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**AHA/ACC/HFSA CLINICAL PRACTICE GUIDELINE**

2022 AHA/ACC/HFSA Guideline for the Management of Heart Failure: A Report of the American College of Cardiology/American Heart Association Joint Committee on Clinical Practice Guidelines

**Writing Committee Members\***

Paul A. Heidenreich, MD, MS, FACC, FAHA, FHFSA, Chair†; Biykem Bozkurt, MD, PhD, FACC, FAHA, FHFSA, Vice Chair†;

David Aguilar, MD, MSc, FAHA†; Larry A. Allen, MD, MHS, FACC, FAHA, FHFSA†; Joni J. Byun†; Monica M. Colvin, MD, MS, FAHA†; Anita Deswal, MD, MPH, FACC, FAHA, FHFSA‡; Mark H. Drazner, MD, MSc, FACC, FAHA, FHFSA†;

Shannon M. Dunlay, MD, MS, FAHA, FHFSA†; Linda R. Evers, JD†; James C. Fang, MD, FACC, FAHA, FHFSA†; Savitri E. Fedson, MD, MA†; Gregg C. Fonarow, MD, FACC, FAHA, FHFSA§; Salim S. Hayek, MD, FACC†; Adrian F. Hernandez, MD, MHS‡; Prateeti Khazanie, MD, MPH, FHFSA†; Michelle M. Kittleson, MD, PhD†;

Christopher S. Lee, PhD, RN, FAHA, FHFSA†; Mark S. Link, MD†; Carmelo A. Milano, MD†; Lorraine C. Nnacheta, DrPH, MPH†; Alexander T. Sandhu, MD, MS†; Lynne Warner Stevenson, MD, FACC, FAHA, FHFSA†; Orly Vardeny, PharmD, MS, FAHA, FHFSA‖; Amanda R. Vest, MBBS, MPH, FHFSA‖; Clyde W. Yancy, MD, MSc, MACC, FAHA, FHFSA†

**AIM:** The “2022 AHA/ACC/HFSA Guideline for the Management of Heart Failure” replaces the “2013 ACCF/AHA Guideline for the Management of Heart Failure” and the “2017 ACC/AHA/HFSA Focused Update of the 2013 ACCF/AHA Guideline for the Management of Heart Failure.” The 2022 guideline is intended to provide patient-centric recommendations for clinicians to prevent, diagnose, and manage patients with heart failure.

**METHODS:** A comprehensive literature search was conducted from May 2020 to December 2020, encompassing studies, reviews, and other evidence conducted on human subjects that were published in English from MEDLINE (PubMed), EMBASE, the Cochrane Collaboration, the Agency for Healthcare Research and Quality, and other relevant databases. Additional relevant clinical trials and research studies, published through September 2021, were also considered. This guideline was harmonized with other American Heart Association/American College of Cardiology guidelines published through December 2021.

**STRUCTURE:** Heart failure remains a leading cause of morbidity and mortality globally. The 2022 heart failure guideline provides recommendations based on contemporary evidence for the treatment of these patients. The recommendations present an evidence-based approach to managing patients with heart failure, with the intent to improve quality of care and align with patients’ interests. Many recommendations from the earlier heart failure guidelines have been updated with new evidence, and new recommendations have been created when supported by published data. Value statements are provided for certain treatments with high-quality published economic analyses.

\*Writing committee members are required to recuse themselves from voting on sections to which their specific relationships with industry may apply; see Appendix 1 for detailed information. †ACC/AHA Representative. ‡ACC/AHA Joint Committee on Clinical Practice Guidelines Liaison. §ACC/AHA Task Force on Performance Measures Representative. ‖HFSA Representative.

ACC/AHA Joint Committee on Clinical Practice Guidelines Members, see page e986.

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**Key Words:** AHA Scientific Statements ■ heart failure ■ heart failure with reduced ejection fraction ■ heart failure with preserved ejection fraction

* heart failure with mildly reduced ejection fraction ■ systolic heart failure ■ heart failure rehabilitation ■ cardiac failure ■ chronic heart failure
  + acute decompensated heart failure ■ cardiogenic shock ■ beta blockers ■ mineralocorticoid receptor antagonists ■ ACE inhibitors
* angiotensin and neprilysin receptor antagonist ■ sacubitril valsartan ■ angiotensin receptor antagonist ■ sodium glucose co-transporter 2
  + SGLT2 inhibitors ■ cardiac amyloidosis ■ atrial fibrillation ■ congestive heart failure ■ guideline-directed medical therapy
* diabetes ■ cardiomyopathy ■ valvular heart disease ■ mitral regurgitation ■ cardiomyopathy in pregnancy ■ reduced ejection fraction
  + right heart pressure ■ palliative care ■ cardio-oncology ■ social determinants of health

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**TOP 10 TAKE-HOME MESSAGES**

1. Guideline-directed medical therapy (GDMT) for heart failure (HF) with reduced ejection fraction (HFrEF) now includes 4 medication classes that include sodium-glucose cotransporter-2 inhibitors (SGLT2i).
2. SGLT2i have a Class of Recommendation 2a in HF with mildly reduced ejection fraction (HFmrEF). Weaker recommendations (Class of Recommendation 2b) are made for ARNi, ACEi, ARB, MRA, and beta blockers in this population.
3. New recommendations for HFpEF are made for SGLT2i (Class of Recommendation 2a), MRAs (Class of Recommendation 2b), and ARNi (Class of Recommendation 2b). Several prior recommen- dations have been renewed including treatment of hypertension (Class of Recommendation 1), treat- ment of atrial fibrillation (Class of Recommendation 2a), use of ARBs (Class of Recommendation 2b), and avoidance of routine use of nitrates or phospho- diesterase-5 inhibitors (Class of Recommendation 3: No Benefit).
4. Improved LVEF is used to refer to those patients with previous HFrEF who now have an LVEF

>40%. These patients should continue their HFrEF treatment.

1. Value statements were created for select rec- ommendations where high-quality, cost-effec- tiveness studies of the intervention have been published.
2. Amyloid heart disease has new recommendations for treatment including screening for serum and urine monoclonal light chains, bone scintigraphy, genetic sequencing, tetramer stabilizer therapy, and anticoagulation.
3. Evidence supporting increased filling pressures is important for the diagnosis of HF if the LVEF is

>40%. Evidence for increased filling pressures can be obtained from noninvasive (eg, natriuretic pep- tide, diastolic function on imaging) or invasive test- ing (eg, hemodynamic measurement).

1. Patients with advanced HF who wish to prolong survival should be referred to a team specializing in HF. A HF specialty team reviews HF manage- ment, assesses suitability for advanced HF thera- pies, and uses palliative care including palliative inotropes where consistent with the patient’s goals of care.
2. Primary prevention is important for those at risk for HF (stage A) or pre-HF (stage B). Stages of

HF were revised to emphasize the new terminolo- gies of “at risk” for HF for stage A and pre-HF for stage B.

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1. Recommendations are provided for select patients with HF and iron deficiency, anemia, hypertension, sleep disorders, type 2 diabetes, atrial fibrillation, coronary artery disease, and malignancy.

# PREAMBLE

Since 1980, the American College of Cardiology (ACC) and American Heart Association (AHA) have translated scientific evidence into clinical practice guidelines with recommendations to improve cardiovascular health. These guidelines, which are based on systematic meth- ods to evaluate and classify evidence, provide a founda- tion for the delivery of quality cardiovascular care. The ACC and AHA sponsor the development and publication of clinical practice guidelines without commercial sup- port, and members volunteer their time to the writing and review efforts. Guidelines are official policy of the ACC and AHA. For some guidelines, the ACC and AHA part- ner with other organizations.

## Intended Use

Clinical practice guidelines provide recommenda- tions applicable to patients with or at risk of developing cardiovascular disease (CVD). The focus is on medical practice in the United States, but these guidelines are rel- evant to patients throughout the world. Although guide- lines may be used to inform regulatory or payer decisions, the intent is to improve quality of care and align with pa- tients’ interests. Guidelines are intended to define prac- tices meeting the needs of patients in most, but not all, circumstances and should not replace clinical judgment.

## Clinical Implementation

Management, in accordance with guideline recommenda- tions, is effective only when followed by both practitioners and patients. Adherence to recommendations can be enhanced by shared decision-making between clinicians and patients, with patient engagement in selecting inter- ventions on the basis of individual values, preferences, and associated conditions and comorbidities.

## Methodology and Modernization

The ACC/AHA Joint Committee on Clinical Practice Guide- lines (Joint Committee) continuously reviews, updates, and modifies guideline methodology on the basis of published standards from organizations, including the National Acad- emy of Medicine (formerly, the Institute of Medicine),1,2 and on the basis of internal reevaluation. Similarly, presentation and delivery of guidelines are reevaluated and modified in

response to evolving technologies and other factors to op- timally facilitate dissemination of information to health care professionals at the point of care.

Numerous modifications to the guidelines have been implemented to make them shorter and enhance “user friendliness.” Guidelines are written and presented in a modular, “knowledge chunk” format in which each chunk includes a table of recommendations, a brief synopsis, rec- ommendation-specific supportive text and, when appro- priate, flow diagrams or additional tables. Hyperlinked references are provided for each modular knowledge chunk to facilitate quick access and review.In recognition of the importance of cost–value considerations, in certain guidelines, when appropriate and feasible, an assessment of value for a drug, device, or intervention may be per- formed in accordance with the ACC/AHA methodology.3

To ensure that guideline recommendations remain cur- rent, new data will be reviewed on an ongoing basis by the writing committee and staff. Going forward, targeted sections/knowledge chunks will be revised dynamically after publication and timely peer review of potentially practice-changing science. The previous designations of “full revision” and “focused update” will be phased out. For additional information and policies on guideline devel- opment, readers may consult the ACC/AHA guideline methodology manual4 and other methodology articles.5–7

## Selection of Writing Committee Members

The Joint Committee strives to ensure that the guide- line writing committee contains requisite content exper- tise and is representative of the broader cardiovascular community by selection of experts across a spectrum of backgrounds, representing different geographic regions, sexes, races, ethnicities, intellectual perspectives/biases, and clinical practice settings. Organizations and profes- sional societies with related interests and expertise are invited to participate as partners or collaborators.

**Relationships With Industry and Other Entities** The ACC and AHA have rigorous policies and methods to ensure that documents are developed without bias or

improper influence. The complete policy on relationships

with industry and other entities (RWI) can be found [online](https://www.acc.org/guidelines/about-guidelines-and-clinical-documents/relationships-with-industry-policy). Appendix 1 of the guideline lists writing committee mem- bers’ relevant RWI; for the purposes of full transparency, their comprehensive disclosure information is available in a [Supplemental Appendix](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063). Comprehensive disclosure in- formation for the Joint Committee is also available [online](https://www.acc.org/guidelines/about-guidelines-and-clinical-documents/guidelines-and-documents-task-forces).

## Evidence Review and Evidence Review Committees

In developing recommendations, the writing committee uses evidence-based methodologies that are based on

all available data.4,5 Literature searches focus on ran- domized controlled trials (RCTs) but also include regis- tries, nonrandomized comparative and descriptive stud- ies, case series, cohort studies, systematic reviews, and expert opinion. Only key references are cited.

An independent evidence review committee is com- missioned when there are 1 questions deemed of utmost clinical importance and merit formal systematic review to determine which patients are most likely to ben- efit from a drug, device, or treatment strategy, and to what degree. Criteria for commissioning an evidence review committee and formal systematic review include absence of a current authoritative systematic review, feasibility of defining the benefit and risk in a time frame consistent with the writing of a guideline, relevance to a substantial number of patients, and likelihood that the findings can be translated into actionable recommendations. Evidence review committee members may include methodologists, epidemiologists, clinicians, and biostatisticians. Recom- mendations developed by the writing committee on the basis of the systematic review are marked “SR.”

## Guideline-Directed Medical Therapy

The term guideline-directed medical therapy (GDMT) encompasses clinical evaluation, diagnostic testing, and both pharmacological and procedural treatments. For these and all recommended drug treatment regimens, the reader should confirm dosage with product insert material and evaluate for contraindications and interactions. Rec- ommendations are limited to drugs, devices, and treat- ments approved for clinical use in the United States.

*Joshua A. Beckman, MD, MS, FAHA, FACC Chair, ACC/AHA Joint Committee on Clinical*

*Practice Guidelines*

*onists; ACE-inhibitors, angiotensin and neprilysin recep- tor antagonist; sacubitril valsartan; angiotensin receptor antagonist; Sodium glucose co-transporter 2 or SGLT2 inhibitors; cardiac amyloidosis; atrial fibrillation; conges- tive heart failure; guideline-directed medical therapy; HFrEF; diabetes mellitus; cardiomyopathy; cardiac amyloidosis; valvular heart disease; mitral regurgitation; cardiomyopathy in pregnancy; reduced ejection fraction; right heart pressure; palliative care*.

Additional relevant studies, published through Sep- tember 2021 during the guideline writing process, were also considered by the writing committee and added to the evidence tables when appropriate. This guideline was harmonized with other ACC/AHA guidelines published through December 2021.The final evidence tables are included in the [Online Data Supplement](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063) and summarize the evidence used by the writing committee to formulate recommendations. References selected and published in the present document are representative and not all- inclusive.

## 1.2. Organization of the Writing Committee

The writing committee consisted of cardiologists, HF specialists, internists, interventionalists, an electro- physiologist, surgeons, a pharmacist, an advanced nurse practitioner, and 2 lay/patient representatives. The writing committee included representatives from the ACC, AHA, and Heart Failure Society of America (HFSA). Appendix 1 of the present document lists writ- ing committee members’ relevant RWI. For the purpos- es of full transparency, the writing committee members’ comprehensive disclosure information is available in a [Supplemental Appendix](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063).

**1.3. Document Review and Approval**

# INTRODUCTION

**CLINICAL STATEMENTS AND GUIDELINES**

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## Methodology and Evidence Review

The recommendations listed in this guideline are, whenever possible, evidence based. An initial exten- sive evidence review, which included literature derived from research involving human subjects, published in English, and indexed in MEDLINE (through PubMed), EMBASE, the Cochrane Collaboration, the Agency for Healthcare Research and Quality, and other selected databases relevant to this guideline, was conducted from May 2020 to December 2020. Key search words included but were not limited to the following: *heart failure; heart failure with reduced ejection fraction; heart failure with preserved ejection fraction; heart failure with mildly reduced ejection fraction; systolic heart failure; heart failure rehabilitation; cardiac failure; chronic heart failure; acute decompensated heart failure; cardiogenic shock; beta blockers; mineralocorticoid receptor antag-*

This document was reviewed by 2 official reviewers nom- inated by the AHA; 1 official reviewer nominated by the ACC; 2 official reviewers from the HFSA; 1 official Joint Committee on Clinical Practice Guidelines reviewer; and 32 individual content reviewers. Reviewers’ RWI informa- tion was distributed to the writing committee and is pub- lished in this document (Appendix 2).This document was approved for publication by the governing bodies of the ACC, AHA, and HFSA.

## Scope of the Guideline

The purpose of the “2022 AHA/ACC/HFSA Guideline for the Management of Heart Failure” (2022 HF guide- line) is to provide an update and to consolidate the “2013 ACCF/AHA Guideline for the Management of Heart Failure”1 for adults and the “2017 ACC/AHA/HFSA Focused Update of the 2013 ACCF/AHA Guideline for the Management of Heart Failure”2 into a new document.

Related ACC/AHA guidelines include recommendations relevant to HF and, in such cases, the HF guideline re- fers to these documents. For example, the 2019 primary prevention of cardiovascular disease guideline3 includes recommendations that will be useful in preventing HF, and the 2021 valvular heart disease guideline4 provides recommendations for mitral valve (MV) clipping in mitral regurgitation (MR).

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**CLINICAL STATEMENTS AND GUIDELINES**

## Areas of focus include:

* + - Prevention of HF.
    - Management strategies in stage C HF, including:
      * New treatment strategies in HF, including sodium-glucose cotransporter-2 inhibitors (SGLT2i) and angiotensin receptor-neprilysin inhibitors (ARNi).
      * Management of HF and atrial fibrillation (AF), including ablation of AF.
      * Management of HF and secondary MR, includ- ing MV transcatheter edge-to-edge repair.
    - Specific management strategies, including:
      * Cardiac amyloidosis.
      * Cardio-oncology.
    - Implantable devices.
    - Left ventricular assist device (LVAD) use in stage D HF.

The intended primary target audience consists of clinicians who are involved in the care of patients with HF. Recommendations are stated in reference to the patients and their condition. The focus is to provide the most up-to-date evidence to inform the clinician during shared decision-making with the patient. Although the present document is not intended to be a procedural- based manual of recommendations that outlines the best practice for HF, there are certain practices that clinicians might use that are associated with improved clinical outcomes.

In developing the 2022 HF guideline, the writing com- mittee reviewed previously published guidelines and related statements. Table 1 contains a list of these guide- lines and statements deemed pertinent to this writing effort and is intended for use as a resource, thus obviating the need to repeat existing guideline recommendations.

## Class of Recommendation and Level of Evidence

The Class of Recommendation (COR) indicates the strength of recommendation, encompassing the estimat- ed magnitude and certainty of benefit in proportion to risk. The Level of Evidence (LOE) rates the quality of sci- entific evidence supporting the intervention on the basis of the type, quantity, and consistency of data from clinical trials and other sources (Table 2).1

## Abbreviations

|  |  |
| --- | --- |
| **Abbreviation** | **Meaning/Phrase** |
| ACEi | angiotensin-converting enzyme inhibitors |
| ACS | acute coronary syndrome |
| ARNi | angiotensin receptor-neprilysin inhibitors |
| ARB | angiotensin (II) receptor blockers |
| AF | atrial fibrillation |
| AL-CM | immunoglobulin light chain amyloid cardiomyopathy |
| ATTR-CM | transthyretin amyloid cardiomyopathy |
| ATTRv | variant transthyretin amyloidosis |
| ATTRwt | wild-type transthyretin amyloidosis |
| BNP | B-type natriuretic peptide |
| CABG | coronary artery bypass graft |
| CAD | coronary artery disease |
| CCM | cardiac contractility modulation |
| CHF | congestive heart failure |
| CKD | chronic kidney disease |
| CMR | cardiovascular magnetic resonance |
| COVID-19 | coronavirus disease 2019 |
| CPET | cardiopulmonary exercise test |
| CRT | cardiac resynchronization therapy |
| CRT-D | cardiac resynchronization therapy with defibrillation |
| CRT-P | cardiac resynchronization therapy with pacemaker |
| CT | computed tomography |
| CVD | cardiovascular disease |
| CVP | central venous pressure |
| DOAC | direct-acting oral anticoagulants |
| DPP-4 | dipeptidyl peptidase-4 |
| ECG | electrocardiogram |
| EF | ejection fraction |
| eGFR | estimated glomerular filtration rate |
| FDA | US Food and Drug Administration |
| FLC | free light chain |
| GDMT | guideline-directed medical therapy |
| HF | heart failure |
| HFimpEF | heart failure with improved ejection fraction |
| HFmrEF | heart failure with mildly reduced ejection fraction |
| HFpEF | heart failure with preserved ejection fraction |
| HFrEF | heart failure with reduced ejection fraction |
| ICD | implantable cardioverter-defibrillator |
| IFE | immunofixation electrophoresis |
| LBBB | left bundle branch block |
| LV | left ventricular |
| LVAD | left ventricular assist device |
| LVEDV | left ventricular end-diastolic volume |
| LVEF | left ventricular ejection fraction |
| LVH | left ventricular hypertrophy |

sociated with reduced survival.3 Therapeutic interven- tions in each stage aim to modify risk factors (stage A), treat risk and structural heart disease to prevent HF (stage B), and reduce symptoms, morbidity, and mortality (stages C and D). To address the evolving role of biomarkers and structural changes for recogni- tion of patients who are at risk of developing HF, who are potential candidates for targeted treatment strat- egies for the prevention of HF, and to enhance the understanding and adoption of these classifications, the writing committee proposed the terminologies listed in Table 3 for the stages of HF. For thresholds of cardiac structural, functional changes, elevated fill- ing pressures, and biomarker elevations, refer to Ap- pendix 3.

|  |  |
| --- | --- |
| **Abbreviation** | **Meaning/Phrase** |
| MCS | mechanical circulatory support |
| MI | myocardial infarction |
| MR | mitral regurgitation |
| MRA | mineralocorticoid receptor antagonist |
| MV | mitral valve |
| NSAID | nonsteroidal anti-inflammatory drug |
| NSVT | nonsustained ventricular tachycardia |
| NT-proBNP | N-terminal prohormone of B-type natriuretic peptide |
| NYHA | New York Heart Association |
| QALY | quality-adjusted life year |
| QOL | quality of life |
| PA | pulmonary artery |
| PCWP | pulmonary capillary wedge pressure |
| PET | positron emission tomography |
| PPAR-γ | peroxisome proliferator-activated receptor gamma |
| PUFA | polyunsaturated fatty acid |
| RA | right atrial |
| RASS | renin-angiotensin-aldosterone system |
| RAASi | renin-angiotensin-aldosterone system inhibitors |
| RCT | randomized controlled trial |
| RV | right ventricular |
| SCD | sudden cardiac death |
| SGLT2i | sodium-glucose cotransporter-2 inhibitors |
| SPECT | single photon emission CT |
| 99mTc-PYP | technetium pyrophosphate |
| TEER | transcatheter mitral edge-to-edge repair |
| TTE | transthoracic echocardiogram |
| VA | ventricular arrhythmia |
| VF | ventricular fibrillation |
| VHD | valvular heart disease |
| VO2 | oxygen consumption/oxygen uptake |
| VT | ventricular tachycardia |

## New York Heart Association (NYHA) Classification

The NYHA classification is used to characterize symp- toms and functional capacity of patients with symp- tomatic (stage C) HF or advanced HF (stage D). It is a subjective assessment by a clinician and can change over time. Although reproducibility and validity can be limited,4,5 the NYHA functional classification is an inde- pendent predictor of mortality,6,7 and it is widely used in clinical practice to determine the eligibility of patients for treatment strategies. Clinicians specify NYHA classifica- tion at baseline after the initial diagnosis and after treat- ment through the continuum of care of a patient with HF. Although a patient with symptomatic HF (stage C) may become asymptomatic with treatment (NYHA class I), that patient will still be categorized as stage C HF. Pa- tients with stage C HF can be classified according to the trajectory of their symptoms (Figure 2).

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# DEFINITION OF HF

## HF Description

HF is a complex clinical syndrome with symptoms and signs that result from any structural or functional im- pairment of ventricular filling or ejection of blood. The writing committee recognizes that asymptomatic stages with structural heart disease or cardiomyopathies are not covered under the above definition as having HF. Such asymptomatic stages are considered at-risk for HF (stage A) or pre-HF (stage B), as explained in Section 2.1, “Stages of HF. ”

## Stages of HF

The ACC/AHA stages of HF (Figure 1, Table 3) emphasize the development and progression of dis- ease,1,2 and advanced stages and progression are as-

## Classification of HF by Left Ventricular Ejection Fraction (LVEF)

LVEF is considered important in the classification of patients with HF because of differing prognosis and response to treatments and because most clinical tri- als select patients based on ejection fraction (EF). RCTs with evidence of survival benefit in patients with HF have mainly enrolled patients with HF with an LVEF 35% or 40%, often labeled HF with reduced ejection fraction (HFrEF).1 In this guideline, HFrEF is defined as LVEF 40% (Table 4). HF with preserved EF (HFpEF) represents at least 50% of the popula- tion with HF, and its prevalence is increasing.2 HFpEF has been variably classified as LVEF >40%, >45%, or

50%. Because some of these patients do not have entirely normal LVEF but also do not have major re- duction in systolic function, the term preserved EF has been used. In this guideline, the threshold for HFpEF is an LVEF 50% (Table 4).

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##### Table 1. Associated Guidelines and Statements

|  |  |  |
| --- | --- | --- |
| **Title** | **Organization** | **Publication Year (Reference)** |
| Guidelines | | |
| 2011 ACCF/AHA Guideline for Coronary Artery Bypass Graft Surgery  *Hillis et al., “2011 ACCF/AHA Guideline for Coronary Artery Bypass Graft Surgery” is now replaced and retired by the “2021 ACC/AHA/SCAI Guideline for Coronary Artery Revascularization”*5 | ACCF/AHA | 20116 |
| 2011 ACCF/AHA/SCAI Guideline for Percutaneous Coronary Intervention  *Levine et al., “2011 ACCF/AHA/SCAI Guideline for Percutaneous Coronary Intervention,” is now replaced and retired by the “2021 ACC/AHA/SCAI Guideline for Coronary Artery Revascularization”*5 | ACCF/AHA/SCAI | 20117 |
| 2015 ACCF/AHA/SCAI Focused Update Guideline for Percutaneous Coronary Intervention | ACCF/AHA/SCAI | 20168 |
| 2020 ACC/AHA Guideline for the Management of Patients With Valvular Heart Disease | ACC/AHA | 20214 |
| 2020 AHA/ACC Guideline for the Diagnosis and Treatment of Patients With Hypertrophic Cardiomyopathy | ACC/AHA | 20209 |
| 2019 ACC/AHA Guideline on the Primary Prevention of Cardiovascular Disease | ACC/AHA | 20193 |
| 2019 AHA/ACC/HRS Focused Update of the 2014 AHA/ACC/HRS Guideline for the Management of Patients With Atrial Fibrillation | AHA/ACC/HRS | 201910 |
| 2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detec- tion, Evaluation, and Management of High Blood Pressure in Adults | ACC/AHA/AAPA/ABC/  ACPM/AGS/AphA/ASH/ ASPC/NMA/PCNA | 201811 |
| 2017 ACC/AHA/HFSA Focused Update of the 2013 ACCF/AHA Guideline for the Management of Heart Failure | ACC/AHA/HFSA | 20172 |
| 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 | ACC/AHA/HFSA | 201612 |
| 2014 ACC/AHA/AATS/PCNA/SCAI/STS Focused Update of the Guideline for the Diagnosis and Management of Patients With Stable Ischemic Heart Disease | ACC/AHA/AATS/PCNA/ SCAI/STS | 201413\* |
| 2013 AHA/ACC Guideline on Lifestyle Management to Reduce Cardiovascular Risk | AHA/ACC | 201414 |
| 2013 AHA/ACC/TOS Guideline for the Management of Overweight and Obesity in Adults | AHA/ACC/TOS | 201415 |
| 2018 AHA/ACC/AACVPR/AAPA/ABC/ACPM/ADA/AGS/AphA/ASPC/NLA/PCNA Guideline on the Manage- ment of Blood Cholesterol | AHA/ACC/AACVPR/AAPA/ ABC/ACPM/ADA/AGS/ AphA/ASPC/NLA/PCNA | 201916 |
| 2013 ACC/AHA Guideline on the Treatment of Blood Cholesterol to Reduce Atherosclerotic Cardiovascular Risk in Adults | ACC/AHA | 201417 |
| 2013 ACC/AHA Guideline on the Assessment of Cardiovascular Risk | ACC/AHA | 201418 |
| 2013 ACCF/AHA Guideline for the Management of Heart Failure | ACCF/AHA | 20131 |
| 2013 ACCF/AHA Guideline for the Management of ST-Elevation Myocardial Infarction | ACCF/AHA | 201319 |
| 2012 ACCF/AHA/HRS Focused Update of the 2008 Guidelines for Device-Based Therapy of Cardiac Rhythm Abnormalities | ACCF/AHA/HRS | 201220 |
| 2012 ACCF/AHA/ACP/AATS/PCNA/SCAI/STS Guideline for the Diagnosis and Management of Patients With Stable Ischemic Heart Disease | ACCF/AHA/ACP/AATS/ PCNA/SCAI/STS | 201221 |
| Effectiveness-Based Guidelines for the Prevention of Cardiovascular Disease in Women—2011 Update | AHA | 201122 |
| AHA/ACCF Secondary Prevention and Risk Reduction Therapy for Patients With Coronary and Other Atheroscle- rotic Vascular Disease: 2011 Update | AHA/ACCF | 201123 |
| 2010 ACCF/AHA Guideline for Assessment of Cardiovascular Risk in Asymptomatic Adults | ACCF/AHA | 201024 |
| Part 9: Post–Cardiac Arrest Care: 2010 American Heart Association Guidelines for Cardiopulmonary Resuscita- tion and Emergency Cardiovascular Care | AHA | 201025 |
| Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure | NHLBI | 200326 |
| Statements | | |
| Cardiac Amyloidosis: Evolving Diagnosis and Management | AHA | 202027 |
| Testing of Low-Risk Patients Presenting to the Emergency Department With Chest Pain | AHA | 201028 |
| Primary Prevention of Cardiovascular Diseases in People With Diabetes Mellitus | AHA/ADA | 200729 |
| Prevention and Control of Influenza | CDC | 200530 |

AATS indicates American Association for Thoracic Surgery; AACVPR, American Association of Cardiovascular and Pulmonary Rehabilitation; AAPA, American Association Academy of Physician Assistants; ABC, Association of Black Cardiologists; ACC, American College of Cardiology; ACCF, American College of Cardiol- ogy Foundation; ACPM, American College of Preventive Medicine; ADA, American Diabetes Association; AGS, American Geriatrics Society; AHA, American Heart Association; AphA, American Pharmacists Association; ASH, American Society of Hypertension; ASPC, American Society for Preventive Cardiology; CDC, Centers for Disease Control and Prevention; ESC, European Society of Cardiology; HFSA, Heart Failure Society of America; HRS, Heart Rhythm Society; NHLBI, National Heart, Lung, and Blood Institute; NICE, National Institute for Health and Care Excellence; NMA, National Medical Association; NLA, National Lipid Association; PCNA, Preventive Cardiovascular Nurses Association; SCAI, Society for Cardiovascular Angiography and Interventions; SIHD, stable ischemic heart disease; STS, Society of Thoracic Surgeons; TOS, The Obesity Society; and WHF, World Heart Federation.

\*The full SIHD guideline is from 2012.21 A focused update was published in 2014.13

**Table 2. Applying American College of Cardiology/American Heart Association Class of Recommendation and Level of Evidence to Clinical Strategies, Interventions, Treatments, or Diagnostic Testing in Patient Care (Updated May 2019)\***

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Patients with HF and an LVEF between the HFrEF and HFpEF range have been termed as “HF with mid- range EF, ”3,4 or “HF with mildly reduced EF. ”4 Because of LVEF being lower than normal, these patients are classified in this document as HF with mildly reduced EF (HFmrEF). Patients with HFmrEF are usually in a dynamic trajectory to improvement from HFrEF or to deterioration to HFrEF (Figure 3). Therefore, for patients whose EF falls into this mildly reduced cat- egory, 1 EF measurement at 1 time point may not be adequate, and the trajectory of LVEF over time and the cause is important to evaluate (Figure 3). Further- more, the diagnosis of HFmrEF and HFpEF can be challenging. Although the classic clinical signs and symptoms of HF, together with EF of 41% to 49%

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or 50%, respectively, are necessary for the diagno- sis of the HFmrEF and HFpEF, the requirements for additional objective measures of cardiac dysfunction can improve the diagnostic specificity. The signs and symptoms of HF are frequently nonspecific and over- lap with other clinical conditions. Elevated natriuretic peptide levels are supportive of the diagnosis, but normal levels do not exclude a diagnosis of HFmrEF or HFpEF. To improve the specificity of diagnosing HFmrEF and HFpEF, the clinical diagnosis of HF in these EF categories should be further supported by objective measures. Therefore, the writing commit- tee proposes the addition of evidence of spontane- ous (at rest) or provokable (eg, during exercise, fluid challenge) increased LV filling pressures (eg, elevated

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##### Figure 1. ACC/AHA Stages of HF.

The ACC/AHA stages of HF are shown. ACC indicates American College of Cardiology; AHA, American Heart Association; CVD, cardiovascular disease; GDMT, guideline-directed medical therapy; and HF, heart failure.

natriuretic peptide, noninvasive/invasive hemody- namic measurement) to the classifications of HFm- rEF and HFpEF (Table 4).

The “2013 ACCF/AHA Guideline for the Man- agement of Heart Failure”1 has used the HFpEF- improved terminology for those whose EF improved from a lower level to EF >40% under the subgroup- ing of patients with HFpEF. Others have proposed a

working definition of HF-recovered EF that included a baseline LVEF 40%, a 10% increase from baseline LVEF, and a second measurement of LVEF >40%.3 Although associated with better outcomes, improve- ment in LVEF does not mean full myocardial recov- ery or normalization of LV function. In most patients, cardiac structural abnormalities, such as LV cham- ber dilatation and ventricular systolic and diastolic

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##### Table 3. Stages of HF

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| **Stages** | **Definition and Criteria** |
| Stage A: At Risk for HF | At risk for HF but without symptoms, structural heart disease, or cardiac biomarkers of stretch or injury (eg, patients with hypertension, atherosclerotic CVD, diabetes, metabolic syndrome and obesity, exposure to cardiotoxic agents, genetic variant for cardiomyopathy, or positive family history of cardiomyopathy). |
| Stage B: Pre-HF | No symptoms or signs of HF and evidence of 1 of the following: |
| *Structural heart disease*\*  Reduced left or right ventricular systolic function Reduced ejection fraction, reduced strain  Ventricular hypertrophy Chamber enlargement Wall motion abnormalities Valvular heart disease |
| *Evidence for increased filling pressures*\*  By invasive hemodynamic measurements  By noninvasive imaging suggesting elevated filling pressures (eg, Doppler echocardiography) |
| *Patients with risk factors and Increased levels of BNPs*\* *or*  *Persistently elevated cardiac troponin*  in the absence of competing diagnoses resulting in such biomarker elevations such as acute coronary syndrome, CKD, pulmonary embolus, or myopericarditis |
| Stage C: Symptomatic HF | Structural heart disease with current or previous symptoms of HF. |
| Stage D: Advanced HF | Marked HF symptoms that interfere with daily life and with recurrent hospitalizations despite attempts to optimize GDMT. |

BNP indicates B-type natriuretic peptide; CKD, chronic kidney disease; CVD, cardiovascular disease; GDMT, guideline-directed medical therapy; and HF, heart failure.

\*For thresholds of cardiac structural, functional changes, elevated filling pressures, and biomarker elevations, refer to Appendix 3.

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##### Figure 2. Trajectory of Stage C HF.

The trajectory of stage C HF is displayed. Patients whose symptoms and signs of HF are resolved are still stage C and should be treated accordingly. If all HF symptoms, signs, and structural abnormalities resolve, the patient is considered to have HF in remission. HF indicates heart failure; and LV, left ventricular. \*Full resolution of structural and functional cardiac abnormalities is uncommon.

dysfunction, may persist. Furthermore, changes in LVEF might not be unidirectional; a patient may have improvement followed by a decrease in EF or vice versa depending on the underlying cause, duration of disease, adherence to the GDMT, or reexposure to cardiotoxicity.5 Therefore, the writing committee elected not to use “recovered EF” or HFpEF, even if subsequent LVEF was >50% but, rather, “HF with improved EF” (HFimpEF) as a subgroup of HFrEF to characterize these patients (Table 4, Figure 3). Impor- tantly, EF can decrease after withdrawal of pharmaco- logical treatment in many patients who had improved EF to normal range with GDMT.5 Trajectory of LVEF can be important, and a significant reduction in LVEF over time is a poor prognostic factor.

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##### Table 4. Classification of HF by LVEF

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| **Type of HF According to LVEF** | **Criteria** |
| HFrEF (HF with reduced EF) | LVEF 40% |
| HFimpEF (HF with improved EF) | Previous LVEF 40% and a follow-up measurement of LVEF >40% |
| HFmrEF (HF with mildly re- duced EF) | LVEF 41%–49%  Evidence of spontaneous or provokable increased LV filling pressures (eg, elevated natriuretic peptide, noninvasive and invasive hemodynamic measurement) |
| HFpEF (HF with preserved EF) | LVEF 50%  Evidence of spontaneous or provokable increased LV filling pressures (eg, elevated natriuretic peptide, noninvasive and invasive hemodynamic measurement) |

Please see Appendix 3 for suggested thresholds for structural heart disease and evidence of increased filling pressures.

HF indicates heart failure; LV, left ventricular; and LVEF, left ventricular ejec- tion fraction.

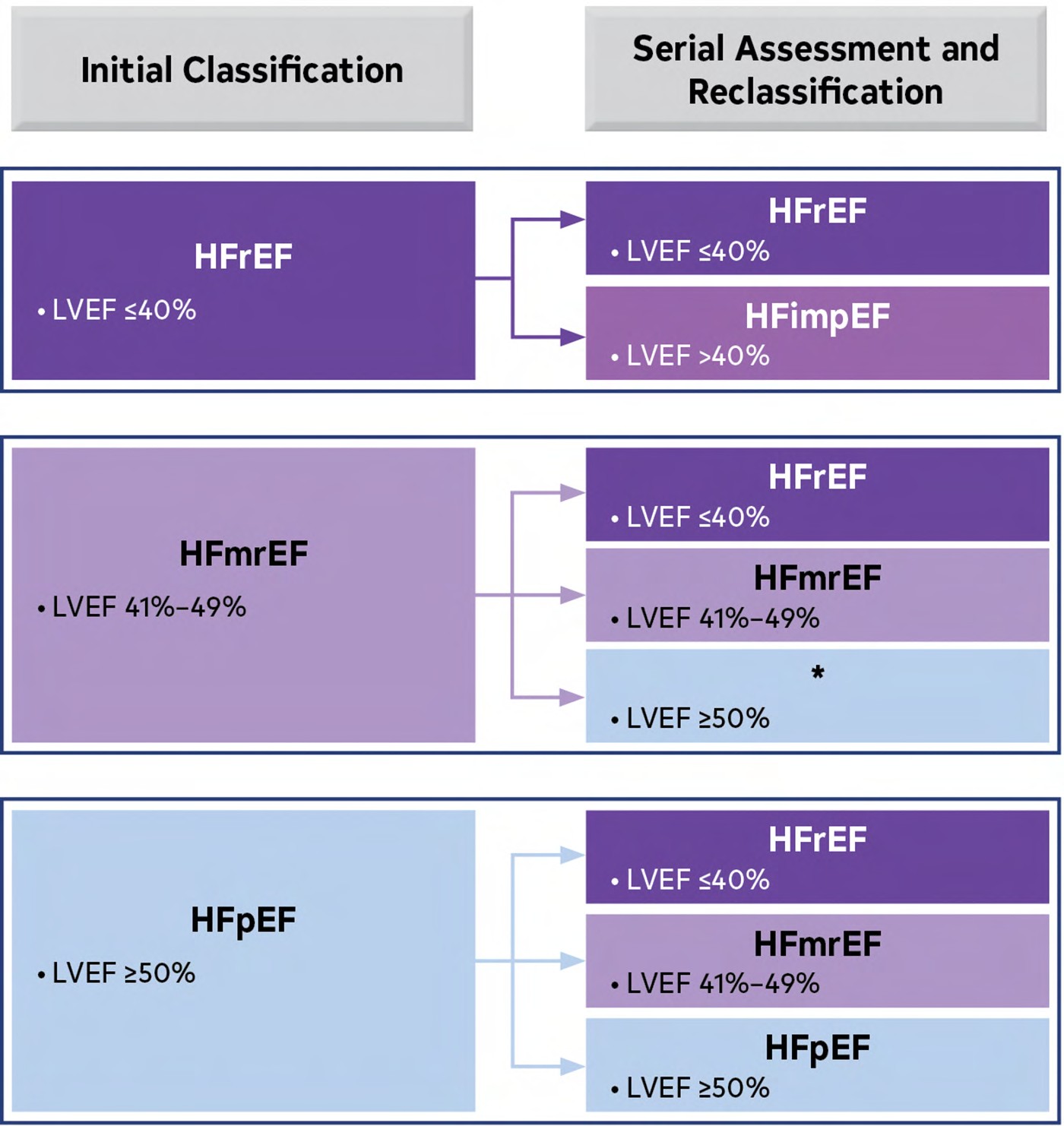
## Diagnostic Algorithm for Classification of HF According to LVEF

Structural and functional alterations of the heart as the underlying cause for the clinical presentation support the diagnosis of HFmrEF and HFpEF1 (Figure 4). The criteria for diagnosis of HFmrEF and HFpEF require evidence of increased LV filling pressures at rest, exer- cise, or other provocations. The criteria can be fulfilled with findings of elevated levels of natriuretic peptides, echocardiographic diastolic parameters such as an E/e′ 15 or other evidence of elevated filling pres- sures, or invasive hemodynamic measurement at rest or exercise. Evidence of structural heart disease (eg, LV structural or functional alterations) may be used to further support the diagnosis of HFpEF. Key structural alterations are an increase in left atrial size and volume (left atrial volume index) and/or an increase in LV mass (LV mass index).

Exercise stress testing with echocardiographic evaluation of diastolic parameters can be helpful if the diagnosis remains uncertain.2,3 Alternatively, or in addition, invasive hemodynamics at rest or with exer- cise, with assessment of filling pressures (pulmonary capillary wedge pressure or LV end diastolic pressures, pulmonary artery [PA] pressures, stroke volumes, and cardiac output) can be performed to help further estab- lish the diagnosis.4

The diagnosis of HFpEF is often challenging. A clin- ical composite score to diagnose HFpEF, the H2FPEF score,5–7 integrates these predictive variables: obesity, atrial fibrillation (AF), age >60 years, treatment with

2 antihypertensive medications, echocardiographic E/e′ ratio >9, and echocardiographic PA systolic pres- sure >35 mm Hg. A weighted score based on these



##### Figure 3. Classification and Trajectories of HF Based on LVEF.

See Appendix 3 for suggested thresholds for laboratory findings. The classification for baseline and subsequent LVEF is shown. Patients with HFrEF who improve their LVEF to >40% are considered to have HFimpEF and should continue HFrEF treatment. HF indicates heart failure; HFimpEF, heart failure with improved ejection fraction; HFmrEF, heart failure with mildly reduced ejection fraction; HFpEF, heart failure with preserved ejection fraction; HFrEF, heart failure with reduced ejection fraction; and LVEF, left ventricular ejection fraction. \*There is limited evidence to guide treatment for patients who improve their LVEF from mildly reduced (41%-49%) to 50%. It is unclear whether to treat these patients as HFpEF or HFmrEF.

6 variables was used to create the composite score ranging from 0 to 9. The odds of HFpEF doubled for each 1-unit score increase (odds ratio, 1.98; 95% CI: 1.74-2.30; *P*<0.0001), with a c-statistic of 0.841. Scores <2 and 6 reflect low and high likelihood, respectively, for HFpEF. A score between 2 and 5 may require further evaluation of hemodynamics with exer- cise echocardiogram or cardiac catheterization to con- firm or negate a diagnosis of HFpEF. The use of this H2FPEF score may help to facilitate discrimination of HFpEF from noncardiac causes of dyspnea and can assist in determination of the need for further diag- nostic testing in the evaluation of patients with unex- plained exertional dyspnea.6,7

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The European Society of Cardiology has developed a diagnostic algorithm.8 This involves a pretest that assesses for HF symptoms and signs, typical clini- cal demographics (obesity, hypertension, diabetes,

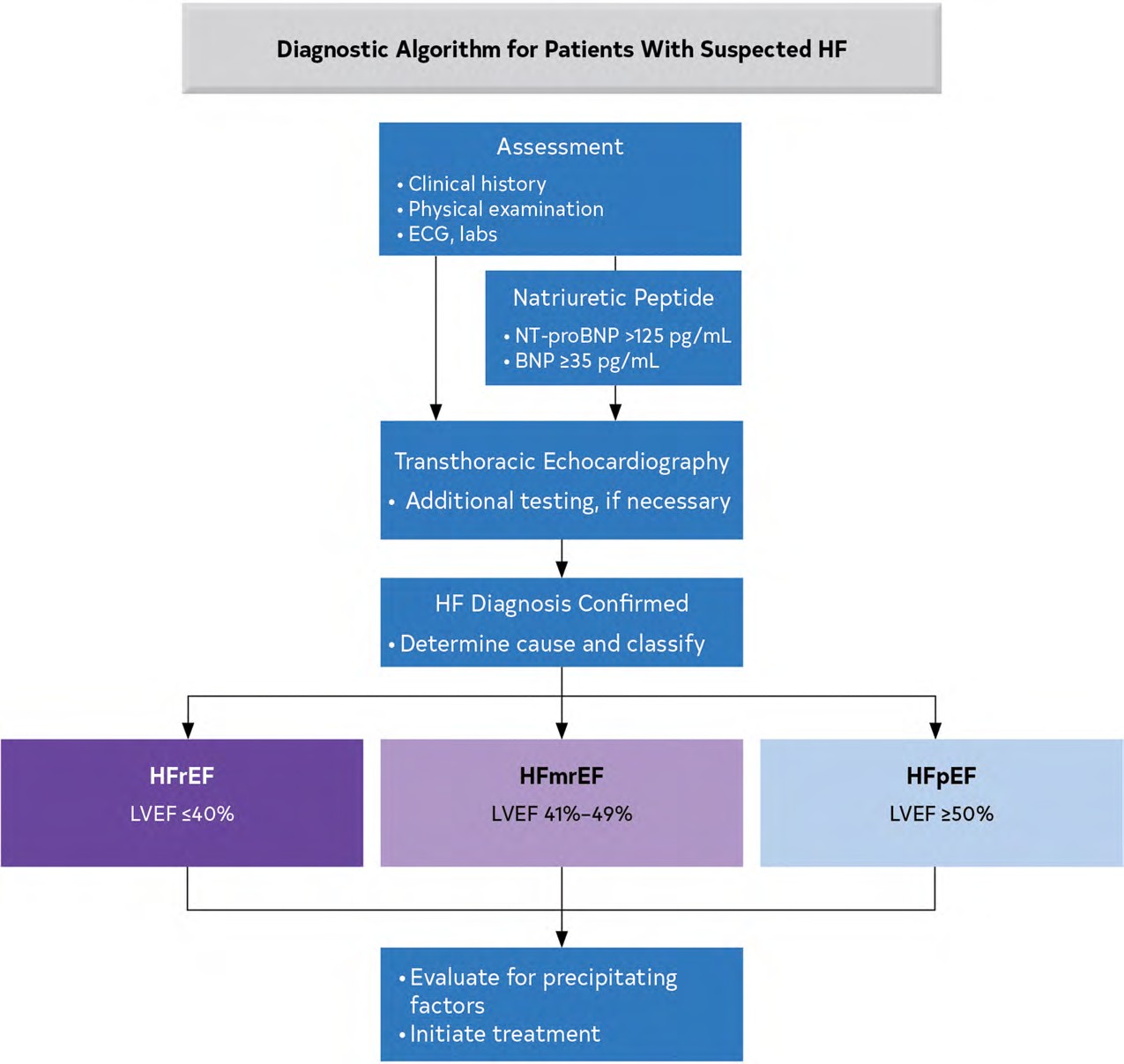
elderly, AF), and diagnostic laboratory tests, ECG, and echocardiography. In the absence of overt noncardiac causes of breathlessness, HFpEF can be suspected if there is a normal LVEF, no significant heart valve disease or cardiac ischemia, and at least 1 typical risk factor. The score used functional, morphological, and biomarker domains. The points score assigns 2 points for a major criterion or 1 point for a minor criterion within each domain, with a maximum of 2 points for each domain.

# EPIDEMIOLOGY AND CAUSES OF HF

## Epidemiology of HF

**Trends in Mortality and Hospitalization for HF** HF is a growing health and economic burden for the United States, in large part because of the aging popula-

##### Figure 4. Diagnostic Algorithm for HF and EF-Based Classification.



The algorithm for a diagnosis of HF and EF-based classification is shown. BNP indicates B-type natriuretic peptide; ECG,

electrocardiogram; EF, ejection fraction; HF, heart failure; HFmrEF, heart failure with mildly reduced ejection fraction; HFpEF, heart failure with preserved ejection fraction; HFrEF, heart failure with reduced ejection fraction; LVEF, left ventricular ejection fraction; LV, left ventricular; and NT-proBNP, N-terminal pro-B type natriuretic peptide.

tion.1,2 Beginning in 2012, the age-adjusted death rate per capita for HF increased for the first time in the Unit- ed States.3 A recent US evaluation found total deaths caused by HF have increased from 275000 in 2009 to 310 000 in 2014.3

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US hospitalizations for HF decreased up until 20124; however, from 2013 to 2017, an increase in HF hospitaliza- tions was observed. In 2017, there were 1.2 million HF hos- pitalizations in the United States among 924 000 patients with HF.4 This represents a 26% increase in HF hospitaliza- tions and number of patients hospitalized with HF.

Although the absolute number of patients with HF has partly grown as a result of the increasing number of older adults, the incidence of HF has decreased.5 Among US Medicare beneficiaries, HF incidence declined from 36 cases per 1000 beneficiaries in 2011 to 27 cases per 1000 beneficiaries in 2014 and remained stable through 2016.5 Divergent trends in the incidence of HF have been observed for those with HFrEF (decreasing incidence) and HFpEF (increasing incidence).6,7 Deaths attributable to cardiomyopathies have been increasing globally because of, in part, increased recognition, diag- nosis, and documentation of specific cardiomyopathies and cardiotoxicity.2

## Racial and Ethnic Disparities in Mortality and Hospitalization for HF

Racial and ethnic disparities in death resulting from HF persist, with non-Hispanic Black patients having the

highest death rate per capita.4 A report examining the US population found age-adjusted mortality rate for HF to be 92 per 100 000 individuals for non-Hispanic Black patients, 87 per 100 000 for non-Hispanic White

patients, and 53 per 100 000 for Hispanic patients.4 Among Medicare beneficiaries, non-Hispanic Black beneficiaries had a slightly greater decrease in HF in- cidence (38 cases per 1000 to 26 cases per 1000, *P*=0.009) than non-Hispanic White beneficiaries (36 cases per 1000 to 28 cases per 1000, *P*=0.003) from 2011 to 2016.4 Among patients with established HF, non-Hispanic Black patients experienced a higher rate of HF hospitalization and a lower rate of death com- pared with non-Hispanic White patients with HF.8–10 Hispanic patients with HF have been found to have similar8 or higher10 HF hospitalization rates and similar10 or lower8 mortality rates compared with non-Hispanic White patients. Asian/Pacific Islander patients with HF have had a similar rate of hospitalization as non-His- panic White patients but a lower rate of death.8,10 These racial and ethnic disparities in outcome, for those with HF, warrant studies and health policy changes to ad- dress health inequity.

## Cause of HF

In the United States, approximately 115 million peo- ple have hypertension, 100 million have obesity, 92 million have prediabetes, 26 million have diabetes, and 125 million have atherosclerotic CVD.1 These are

##### Table 5. Other Potential Nonischemic Causes of HF

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| **Cause** | **Reference** |
| Chemotherapy and other cardiotoxic medications | 23–25 |
| Rheumatologic or autoimmune | 26 |
| Endocrine or metabolic (thyroid, acromegaly, pheochromocy- toma, diabetes, obesity) | 27–31 |
| Familial cardiomyopathy or inherited and genetic heart disease | 32 |
| Heart rhythm–related (eg, tachycardia-mediated, PVCs, RV pacing) | 33 |
| Hypertension | 34 |
| Infiltrative cardiac disease (eg, amyloid, sarcoid, hemochro- matosis) | 21,35,36 |
| Myocarditis (infectious, toxin or medication, immunological, hypersensitivity) | 37,38 |
| Peripartum cardiomyopathy | 39 |
| Stress cardiomyopathy (Takotsubo) | 40,41 |
| Substance abuse (eg, alcohol, cocaine, methamphetamine) | 42–44 |

HF indicates heart failure; PVC, premature ventricular contraction; and RV, right ventricular.

known risk factors with high relative risk and popula- tion attributable risk for development of HF. There- fore, a large proportion of the US population can be categorized as being at-risk for HF or stage A HF. The common causes of HF include ischemic heart disease and myocardial infarction (MI), hypertension, and valvular heart disease (VHD). Other causes can include familial or genetic cardiomyopathies; amyloi- dosis; cardiotoxicity with cancer or other treatments or substance abuse such as alcohol, cocaine, or meth- amphetamine; tachycardia, right ventricular (RV) pac- ing or stress-induced cardiomyopathies; peripartum cardiomyopathy; myocarditis; autoimmune causes, sarcoidosis; iron overload, including hemochromato- sis; and thyroid disease and other endocrine metabol- ic and nutritional causes (Table 5). Furthermore, with cardiac imaging and biomarkers, myocardial injury or cardiac maladaptive structural changes can be de- tected at earlier phases with a higher sensitivity, even in the absence of gross LV dysfunction or symptoms. With the coronavirus disease 2019 (COVID-19) pan- demic, investigators are gaining better insights into infection and inflammation-related myocardial injury and myocarditis. With the increasing ability to detect myocardial injury and with an increasing awareness of cardiotoxicity and injury patterns including inflam- mation, pre-HF or stage B HF will likely continue to increase. Beyond classifications of EF and staging in HF, clinicians should seek the cause of HF be- cause appropriate treatment may be determined by the cause (Table 5).

# INITIAL AND SERIAL EVALUATION

## Clinical Assessment: History and Physical Examination

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| **Recommendations for Clinical Assessment: History and Physical Ex- amination**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. In patients with HF, vital signs and evidence of clinical congestion should be assessed at each encounter to guide overall management, includ- ing adjustment of diuretics and other medica- tions.1–6 |
| **1** | **B-NR** | 2. In patients with symptomatic HF, clinical factors indicating the presence of advanced HF should be sought via the history and physical examina- tion.7–12 |
| **1** | **B-NR** | 3. In patients with cardiomyopathy, a 3-generation family history should be obtained or updated when assessing the cause of the cardiomyopa- thy to identify possible inherited disease.13,14 |
| **1** | **B-NR** | 4. In patients presenting with HF, a thorough history and physical examination should direct diagnos- tic strategies to uncover specific causes that may warrant disease-specific management.15,16 |
| **1** | **C-EO** | 5. In patients presenting with HF, a thorough history and physical examination should be obtained and performed to identify cardiac and noncardiac disorders, lifestyle and behavioral factors, and social determinants of health that might cause or accelerate the development or progression of HF. |

**Synopsis**

The history and physical examination remain a cornerstone in the assessment of patients with HF. The history and physical examination provide information about the cause of an underlying cardiomyopathy, including the possibility of an inherited cardiomyopathy as ascertained by a fam- ily history or a condition requiring disease-specific therapy like amyloid heart disease, as well as reasons why a previ- ously stable patient developed acutely decompensated HF. A critical component of the history and physical examina- tion is to assess for clinical congestion (ie, those signs and symptoms resulting from elevated cardiac filling pressures). Congestion is a target for medication adjustment and is as- sociated with quality of life (QOL) and prognosis. The his- tory and physical examination also allow for the determina- tion of clinical clues that suggest the patient has advanced HF, which may warrant referral to an advanced HF center.

## Recommendation-Specific Supportive Text

* + 1. Clinical congestion can be assessed by various methods, including the presence of jugular venous

distention,17 orthopnea,18 bendopnea,19 a square- wave response to the Valsalva maneuver,20 and leg edema.6 On a practical level, clinicians use extent of clinical congestion to guide titration of pharma- cological treatments, including doses of diuretics. Observational studies have shown that clinical congestion is an important adverse risk factor in patients with HF.1–6,17 Recently, the PARADIGM-HF (The Efficacy and Safety of LCZ696 Compared to Enalapril on Morbidity and Mortality of Patients With Chronic Heart Failure) investigators showed that, in patients with chronic HFrEF, changes in markers of clinical congestion were associated with QOL as assessed by the Kansas City Cardiomyopathy Questionnaire and also provided prognostic infor- mation independently even of natriuretic peptides or the MAGGIC (Meta-analysis Global Group in Chronic Heart Failure) risk score.2 These data highlight the ongoing relevance of clinical con- gestion ascertained by the history and physical examination.

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* + 1. Some patients with HF progress to an advanced state, a condition that can be treated with special- ized interventions such as mechanical circulatory support (MCS) or cardiac transplantation. Such patients should be identified before they progress to a state of extremis, at which point they may suc- cumb to their illness or suffer complications of an intervention as a result of their very advanced state. Several “simple clinical clues” are available to identify advanced HF and should be ascertained via a focused history and physical examination. The recognition that a patient has advanced HF will allow for earlier referral to an advanced HF center, when appropriate, as will be discussed later in this document (see Section 8, “Specialty Referral for Advanced HF”).
    2. Increasingly, familial cardiomyopathy is recognized as a more accurate diagnosis in some patients previously classified as having an idiopathic dilated cardiomyopathy (DCM). A detailed family history may provide the first clue of a genetic basis. A broad array of questions includes whether family members had a weak, enlarged, or thick heart, or HF; muscular dystrophy; a pacemaker or defibril- lator; were on a heart transplant list; or died unex- pectedly. Periodic updating of the family history in patients with a cardiomyopathy of uncertain origin may lead to a diagnosis of familial cardiomyopathy in the event that a relative subsequently develops a cardiomyopathy or a related complication. A 3-gen- eration family pedigree obtained by genetic health

care professionals improved the rate of detection of a familial process as compared with routine care.14 Furthermore, a family history of cardiomy- opathy, as determined by a 3-generation pedigree analysis, was associated with findings of gadolin- ium enhancement on cardiac magnetic resonance imaging (MRI) and increased major adverse car- diac events.13 The possibility of an inherited car- diomyopathy provides the impetus for cascade screening of undiagnosed family members, thereby potentially avoiding preventable adverse events in affected relatives by implementation of GDMT and other management that otherwise would not be initiated.

* + 1. Certain conditions that cause HF require disease- specific therapies. For example, in amyloid heart disease, whether on the basis of transthyretin21 or light chain deposition,22 there are specific treat- ments that otherwise would not be used in patients with HF. Hence, expeditious and accurate diag- nosis of such conditions is important. Currently, important delays have been reported in diagnos- ing amyloid heart disease,16 perhaps not unexpect- edly given the wide spectrum of possible clinical presentations.15 Similarly, HF attributable to sar- coidosis, hemochromatosis, hypothyroidism, hyper- thyroidism, acromegaly, connective tissue disease, tachycardia-induced cardiomyopathy, or high- output HF from an arteriovenous fistula, among others, requires specific therapeutic approaches. Given that the differential diagnosis of HF is broad, the history and physical examination can provide clues to narrow the number of causes to consider and guide the diagnostic approach to identify such conditions (Table 5).
    2. The history and physical examination help to identify the cause of a clinical deterioration. To determine the cause of a clinical deterioration, the clinician assesses for concurrent illness (eg, ongoing myocardial ischemia, pulmonary emboli, or systemic infection), initiation of a medication potentially detrimental in the setting of HF (eg, nonsteroidal anti-inflammatory drugs [NSAIDs]), or the possibility of chronic RV pacing (eg, a newly implanted pacemaker or medications such as amiodarone that leads to bradycardia and resultant chronic RV pacing), nonadherence to a medication or dietary regimen, and ongoing substance abuse. In addition, an assessment of social determinants of health (eg, housing stabil- ity, food security, available transportation) should be made.

* + 1. ***Initial Laboratory and Electrocardiographic Testing***

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| **Recommendations for Initial Laboratory and Electrocardiographic Test- ing**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. For patients presenting with HF, the specific cause of HF should be explored using addi- tional laboratory testing for appropriate man- agement.1–8 |
| **1** | **C-EO** | 2. For patients who are diagnosed with HF, laboratory evaluation should include com- plete blood count, urinalysis, serum electro- lytes, blood urea nitrogen, serum creatinine, glucose, lipid profile, liver function tests, iron studies, and thyroid-stimulating hormone to optimize management. |
| **1** | **C-EO** | 3. For all patients presenting with HF, a 12-lead ECG should be performed at the initial encoun- ter to optimize management. |

## Synopsis

Laboratory evaluation with complete blood count, urinaly- sis, serum electrolytes (including sodium, potassium, cal- cium, and magnesium), blood urea nitrogen, serum cre- atinine, glucose, fasting lipid profile, liver function tests, iron studies (serum iron, ferritin, transferrin saturation), and thyroid-stimulating hormone level and electrocar- diography is part of the standard diagnostic evaluation of a patient with HF. In addition to routine assessment, specific diagnostic testing and evaluation is often neces- sary to identify specific cause and other comorbidities in patients with HF.

## Recommendation-Specific Supportive Text

* + - 1. Identifying the specific cause of HF is impor- tant, because conditions that cause HF may require disease-specific therapies. Depending on the clinical suspicion, additional diagnostic studies are usually required to diagnose specific causes (Table 6) such as ischemic cardiomy- opathy, cardiac amyloidosis, sarcoidosis, hemo- chromatosis, infectious mechanisms (eg, HIV, COVID-19, Chagas), hypothyroidism, hyperthy- roidism, acromegaly, connective tissue disorders, tachycardia-induced cardiomyopathy, Takotsubo, peripartum cardiomyopathy, cardiotoxicity with cancer therapies, or substance abuse would require specific management in addition to or beyond GDMT.1,2,9–15

##### Table 6. Selected Potential Causes of Elevated Natriuretic Peptide Levels50–53

|  |
| --- |
| Cardiac |
| HF, including RV HF syndromes |
| ACS |
| Heart muscle disease, including LVH |
| VHD |
| Pericardial disease |
| AF |
| Myocarditis |
| Cardiac surgery |
| Cardioversion |
| Toxic-metabolic myocardial insults, including cancer chemotherapy |
| Noncardiac |
| Advancing age |
| Anemia |
| Renal failure |
| Pulmonary: Obstructive sleep apnea, severe pneumonia |
| Pulmonary embolism, pulmonary arterial hypertension |
| Critical illness |
| Bacterial sepsis |
| Severe burns |

ACS indicates acute coronary syndromes; AF, atrial fibrillation; HF, heart fail- ure; LVH, left ventricular hypertrophy; RV, right ventricular; and VHD, valvular heart disease.

* + - 1. Laboratory evaluation with complete blood count, urinalysis, serum electrolytes, blood urea nitro- gen, serum creatinine, glucose, fasting lipid pro- file, liver function tests, iron studies (serum iron, ferritin, transferrin saturation), and thyroid-stimu- lating hormone levels provides important informa- tion regarding patients’ comorbidities, suitability for and adverse effects of treatments, potential causes or confounders of HF, severity and prog- nosis of HF, and is usually performed on initial evaluation. Pertinent laboratory tests are repeated with changes in clinical condition or treatments (eg, to monitor renal function or electrolytes with diuretics).
      2. Electrocardiography is part of the routine evalua- tion of a patient with HF and provides important information on rhythm, heart rate, QRS morphol- ogy and duration, cause, and prognosis of HF. It is repeated when there is a clinical indication, such as a suspicion for arrhythmia, ischemia or myocardial injury, conduction, or other cardiac abnormalities.

## Use of Biomarkers for Prevention, Initial Diagnosis, and Risk Stratification

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| **Recommendations for Use of Biomarkers for Prevention, Initial Diagno- sis, and Risk Stratification**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In patients presenting with dyspnea, measure- ment of B-type natriuretic peptide (BNP) or  N-terminal prohormone of B-type natriuretic peptide (NT-proBNP) is useful to support a diagnosis or exclusion of HF.1–12 |
| **1** | **A** | 2. In patients with chronic HF, measurements of BNP or NT-proBNP levels are recommended for risk stratification.11,13–29 |
| **1** | **A** | 3. In patients hospitalized for HF, measurement of BNP or NT-proBNP levels at admission is recommended to establish prognosis.11,13–19 |
| **2a** | **B-R** | 4. In patients at risk of developing HF, BNP or NT-proBNP–based screening followed by team-based care, including a cardiovascular specialist, can be useful to prevent the devel- opment of LV dysfunction or new-onset HF.30,31 |
| **2a** | **B-NR** | 5. In patients hospitalized for HF, a predischarge BNP or NT-proBNP level can be useful to inform the trajectory of the patient and estab- lish a postdischarge prognosis.14,17,20–29 |

**Synopsis**

Assays for BNP and NT-proBNP are frequently used to establish the presence and severity of HF. In general, BNP and NT-proBNP levels are similar, and either can be used in patient care settings as long as their respec- tive absolute values and cut-points are not used inter- changeably.32–34 Obesity is associated with lower levels of BNP and NT-proBNP thereby reducing their diagnostic sensitivity.35,36 A substantial evidence base supports the use of natriuretic peptide biomarkers for excluding HF as a cause of symptoms in ambulatory and emergency department settings. Although a reduction in BNP and NT-proBNP has been associated with better outcomes, the evidence for treatment guidance using serial BNP or NT-proBNP measurements remains insufficient.37–39 Lastly, a widening array of biomarkers including mark- ers of myocardial injury, inflammation, oxidative stress, vascular dysfunction, and matrix remodeling have been shown to provide incremental prognostic information over natriuretic peptides but remain without evidence of an incremental management benefit.13,40–49

## Recommendation-Specific Supportive Text

* + 1. Measurement of BNP and NT-proBNP levels in the ambulatory setting for a suspected cardiac cause of dyspnea provides incremental diagnostic value to clinical judgment when the cause of dyspnea is unclear and the physical examination equivocal.1–9 In the emergency setting, BNP and NT-proBNP

levels have higher sensitivity than specificity and may be more useful for ruling out HF than ruling in HF. Although lower levels of BNP and NT-proBNP may help exclude the presence of HF, and higher levels have high positive predictive value to diag- nose HF, increases in both BNP and NT-proBNP levels have been reported in patients with various cardiac and noncardiac causes (Table 6).50–53

* + 1. and 3. Higher levels of BNP and NT-proBNP are associated with a greater risk for adverse short- and long-term outcomes in patients with HF, includ- ing all-cause and cardiovascular death and major cardiovascular events.11,13–19 Studies have shown incremental prognostic value of these biomark- ers to standard approaches of CVD risk assess- ment.11,16 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 bio- markers in former settings.

1. The STOP-HF (St Vincent’s Screening to Prevent Heart Failure) study is a large single-center trial of patients at risk of HF, defined by the presence of hypertension, diabetes, or known vascular dis- ease but without established LV systolic dys- function or symptomatic HF, who were randomly assigned to screening with BNP testing or usual care.31 Participants in the intervention group with BNP levels 50 pg/mL underwent echocardiog- raphy and referral to a cardiovascular special- ist.31 All patients received coaching by a specialist nurse who provided education on the importance of adherence to medication and healthy lifestyle behaviors.31 BNP-based screening reduced the composite endpoint of incident asymptomatic LV dysfunction with or without newly diagnosed HF. Similarly, accelerated uptitration of renin-angioten- sin-aldosterone system (RAAS) antagonists and beta blockers reduced cardiac events in patients with diabetes and elevated NT-proBNP levels but without cardiac disease at baseline.30 Standardized screening for HF remains challenging as a result of the heterogeneity of risk factors across different patient populations. Studies are needed to assess the cost-effectiveness and risks of such screening, as well as its impact on QOL and mortality.
2. Predischarge BNP and NT-proBNP levels are strong predictors of the risk of death or hospital readmission for HF.14,17,20–29 Although patients in whom levels of BNP or NT-proBNP decreased with treatment had better outcomes than those without any changes or with a biomarker rise,14,23,28,29 tar- geting a certain threshold, value, or relative change in these biomarker levels during hospitalization has not been shown to be consistently effective in improving outcomes.37–39 Patients in which GDMT

##### Table 7. Examples of Factors Implicating Possible Genetic Cardiomyopathy

|  |  |  |
| --- | --- | --- |
| Phenotypic Category | Patient or Family Member Phenotypic Finding\* | Ask Specifically About Family Members\* With |
| Cardiac morphology | Marked LV hypertrophy | Any mention of cardiomyopathy, enlarged or weak heart, HF.  Document even if attributed to other causes, such as alcohol or peri- partum cardiomyopathy |
| LV noncompaction |
| Right ventricular thinning or fatty replacement on imaging or biopsy |
| Findings on 12-lead ECG | Abnormal high or low voltage or conduction, and repolar- ization, altered RV forces | Long QT or Brugada syndrome |
| Dysrhythmias | Frequent NSVT or very frequent PVCs Sustained ventricular tachycardia or fibrillation | ICD  Recurrent syncope  Sudden death attributed to “massive heart attack” without known CAD Unexplained fatal event such as drowning or single-vehicle crash |
| Early onset AF | “Lone” AF before age 65 y |
| Early onset conduction disease | Pacemaker before age 65 y |
| Extracardiac features | Skeletal myopathy Neuropathy Cutaneous stigmata  Other possible manifestations of systemic syndromes | Any known skeletal muscle disease, including mention of Duchenne and Becker’s, Emory-Dreifuss limb-girdle dystrophy  Systemic syndromes: Dysmorphic features Mental retardation Congenital deafness Neurofibromatosis  Renal failure with neuropathy |

AF indicates atrial fibrillation; CAD, coronary artery disease; LV, left ventricular; NSVT, nonsustained ventricular tachycardia; PVC, premature ventricular contraction; and RV, right ventricular.

\*Note that genetic cause is more likely when the person is younger at the onset of events. However, the cardiac morphology and peripheral manifestations of hereditary amyloidosis may present in later life, unlike most other inherited cardiomyopathies.

leads to a reduction in BNP and NT-proBNP lev- els represent a population with improved long-term outcomes compared with those with persistently elevated levels despite appropriate treatment.37–39 BNP and NT-proBNP levels and their change could help guide discussions on prognosis as well as adherence to, and optimization of, GDMT.

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## Genetic Evaluation and Testing

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| **Recommendations for Genetic Evaluation and Testing**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. In first-degree relatives of selected patients with genetic or inherited cardiomyopathies, genetic screening and counseling are recom- mended to detect cardiac disease and prompt consideration of treatments to decrease HF progression and sudden death.1,2 |
| **2a** | **B-NR** | 2. In select patients with nonischemic cardio- myopathy, referral for genetic counseling and testing is reasonable to identify conditions that could guide treatment for patients and family members.3,4 |

**Synopsis**

In patients in whom a genetic or inherited cardiomy- opathy is suspected, a family history should be per- formed, including at least 3 generations and ideally

diagrammed as a family tree pedigree (see Section 4.1, “Clinical Assessment: History and Physical Ex- amination”). Genetic variants have been implicated in 25% to 40% of patients with DCM with a positive fam- ily history but also in 10% to 30% of patients without a recognized family history.3,4 Phenotype and family history are important for identifying patients in whom genetic testing is most likely to yield clinically action- able information (Table 7). Presentation of DCM with conduction disease or ventricular arrhythmias raises concern of sarcoidosis and arrhythmogenic cardio- myopathy, which is of particular concern because of the risk of sudden death in patients and families.5 No controlled studies have shown clinical benefits of ge- netic testing for cardiomyopathy, but genetic testing contributes to risk stratification and has implications for treatment, currently most often for decisions re- garding defibrillators for primary prevention of sudden death5 and regarding exercise limitation for hypertro- phic cardiomyopathy and the desmosomal variants. Consultation with a trained counselor before and af- ter genetic testing helps patients to understand and weigh the implications of possible results for their own lives and those of family members, including pos- sible discrimination on the basis of genetic informa- tion. Unless shown to be free of the genetic variant(s) implicated in the proband, first-degree relatives of af- fected probands should undergo periodic screening with echocardiography and electrocardiography.

## Recommendation-Specific Supportive Text

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* + 1. and 2. Inherited dilated, restrictive, and hypertrophic cardiomyopathies have been identified, although 1 gene variant may cause different phenotypes in the same family. The most common pathogenic variants identified are truncations in the large structural protein titin, which have been implicated in DCM3–5 and also in peripartum or alcoholic cardiomyopathies; however, variants that do not cause disease are also common. Pathogenic variants in lamin A/C can be associated with conduction block and atrial arrhythmias as well as ventricular arrhythmias, which may progress more rapidly than symptoms of HF. Although previously linked with the phenotype of arrhythmogenic RV car- diomyopathy, desmosomal protein variants are now recognized to affect the left ventricle also with or with- out the right ventricle, and the term arrhythmogenic cardiomyopathy is now preferred for the phenotype of arrhythmias combined with DCM. Filamin-C muta- tions have been associated with skeletal myopathies and with isolated cardiomyopathy with ventricular arrhythmias. The identification of pathogenic variants associated with increased risk of sudden death may trigger consideration of primary prevention implant- able cardioverter-defibrillators (ICDs) even in patients who have LVEF >0.35 or <3 months of guideline- recommended therapies.6 Evidence of desmosomal cardiac disease carries the additional implication of advice to avoid strenuous exercise, which may accel- erate ventricular remodeling.7 Genetic confirmation of symptomatic Fabry’s cardiomyopathy is an indication for replacement therapy with the enzyme agalsidase beta, and migalastat was recently approved for this uncommon cardiomyopathy.

## Evaluation With Cardiac Imaging

|  |  |  |
| --- | --- | --- |
| **Recommendations for Evaluation With Cardiac Imaging**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 1. In patients with suspected or new-onset HF, or those presenting with acute decompensated HF, a chest x-ray should be performed to assess heart size and pulmonary congestion and to detect alternative cardiac, pulmonary, and other diseases that may cause or contrib- ute to the patient’s symptoms.1,2 |
| **1** | **C-LD** | 2. In patients with suspected or newly diag- nosed HF, transthoracic echocardiography (TTE) should be performed during initial evaluation to assess cardiac structure and function.3 |

|  |  |  |
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| **Recommendations for Evaluation With Cardiac Imaging (Continued)** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 3. In patients with HF who have had a significant clinical change, or who have received GDMT and are being considered for invasive procedures  or device therapy, repeat measurement of EF, degree of structural remodeling, and valvular function are useful to inform therapeutic interven- tions.4–7 |
| **1** | **C-LD** | 4. In patients for whom echocardiography is inadequate, alternative imaging (eg, cardiac magnetic resonance [CMR], cardiac computed tomography [CT], radionuclide imaging) is rec- ommended for assessment of LVEF.8–15 |
| **2a** | **B-NR** | 5. In patients with HF or cardiomyopathy, CMR can be useful for diagnosis or manage- ment.16–23 |
| **2a** | **B-NR** | 6. In patients with HF, an evaluation for possible ischemic heart disease can be useful to identify the cause and guide management.24–27 |
| **2b** | **B-NR** | 7. In patients with HF and coronary artery disease (CAD) who are candidates for coronary revas- cularization, noninvasive stress imaging (stress echocardiography, single-photon emission CT [SPECT], CMR, or positron emission tomog- raphy [PET]) may be considered for detection of myocardial ischemia to help guide coronary revascularization.28–32 |
| **3: No**  **Benefit** | **C-EO** | 8. In patients with HF in the absence of: 1) clinical status change, 2) treatment interventions that might have had a significant effect on cardiac function, or 3) candidacy for invasive proce- dures or device therapy, routine repeat assess- ment of LV function is not indicated. |

## Synopsis

Cardiac imaging has a key role in the initial evaluation of individuals with suspected HF and, when indicated, in the serial assessment of patients with HF. After a complete history and physical examination, a com- prehensive TTE is the most useful initial diagnostic test given the vast amount of diagnostic and prognos- tic information provided. The determination of LVEF is a fundamental step to classify HF and to guide evidence-based pharmacological and device-based therapy. In certain situations, the echocardiogram is unable to accurately assess cardiac structure and/or function or more information is needed to determine the cause of the cardiac dysfunction. Other imaging modalities, such as CMR, SPECT or radionuclide ven- triculography, PET, or cardiac CT or invasive coronary angiography, can provide additional and complemen- tary information to cardiac ultrasound.11 In general, cardiac imaging tests, including repeat tests, are per- formed only when the results have a meaningful im- pact on clinical care.

## Recommendation-Specific Supportive Text

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* + 1. The chest x-ray is a useful initial diagnostic test for the evaluation of patients presenting with signs and symptoms of HF because it assesses cardiomeg- aly, pulmonary venous congestion, and interstitial or alveolar edema and may reveal alternative causes, cardiopulmonary or otherwise, of the patient’s symptoms.1,2 Apart from congestion, other findings on chest x-ray are associated with HF only in the context of clinical presentation. Importantly, cardio- megaly may be absent in acute HF and, although cephalization, interstitial edema, and alveolar edema are modestly specific for HF, these findings are rel- atively insensitive.2,33 Considering the limited sensi- tivity and specificity, the chest x-ray should not be used as the only determinant of the specific cause or presence of HF.
    2. TTE provides information regarding cardiac struc- ture and function and identifies abnormalities of myocardium, heart valves, and pericardium. Echocardiography reveals structural and functional information that predicts subsequent risk.34–40 Guidelines provide recommendations for quanti- fication of cardiac structure and function, includ- ing LVEF measurements, ventricular dimensions and volumes, evaluation of chamber geometry, and regional wall motion.41 RV size and function, atrial size, and all valves are evaluated for anatomic and flow abnormalities. Guidelines also provide recom- mendations for diastolic function and estimates of LV filling and left atrial pressure.42 The tricus- pid valve regurgitant gradient, coupled with inferior vena cava diameter and its response during respi- ration, provides estimates of systolic PA pressure and central venous pressure. Indices of myocardial deformation, such as global longitudinal strain, may identify subclinical LV systolic dysfunction, which has been associated with greater risk of developing HF or recurrent HF hospitalizations.38,43–46 Given the widespread availability, lack of ionizing radia- tion, and wealth of provided information, echocar- diography is the preferred initial imaging modality for evaluation of patients with suspected HF. Point- of-care cardiac ultrasound is an evolving tool for assessment of cardiac function and assessment of volume status and pulmonary congestion.47–52
    3. Serial echocardiograms to assess changes in EF, structural remodeling, and valvular function, although not recommended routinely in stable patients, are useful in various situations. In patients who have an unexplained, significant change in clinical status, echocardiography can provide important information, such as worsening ven- tricular or valvular function. A subset of patients may also have reverse remodeling, improvement

in LVEF, and valvular function in response to evidence-based medical, revascularization, and device therapies, and repeat assessment of LVEF and remodeling is appropriate in those who have received treatments that might have had a signifi- cant effect on cardiac structure and function.4–7,53–59 Recovery of function appears more common in those with LV systolic dysfunction occurring in the setting of adverse energetic circumstances (eg, chronic tachycardia or thyroid disease), dilated car- diomyopathies associated with immune responses (eg, peripartum cardiomyopathy, acute myocardi- tis, systemic inflammatory responses), or in those who have undergone revascularization or device- based therapies.60 Reevaluation of EF (>40 days after MI, >90 days after revascularization, >90 days after GDMT) is useful to determine candidacy for implantable cardioverter-defibrillator (ICD) or cardiac resynchronization therapy (CRT). Finally, repeat surveillance of LV function is appropriate in patients exposed to treatments that potentially damage the myocardium, such as chemotherapy.

* + 1. If TTE is unable to accurately evaluate cardiac structure and function, additional noninvasive imaging modalities are available to clarify the ini- tial diagnosis and to provide information on car- diac structure and function. The choice between these modalities depends on availability, local expertise, patient characteristics, indication, and goal of limiting radiation exposure. CMR provides an accurate and highly reproducible assessment of cardiac volumes, mass, and EF of the left and right ventricles.8–10 CMR provides high anatomic resolution of all aspects of the heart and surround- ing structures and is not associated with ioniz- ing radiation, leading to its recommended use in known or suspected congenital heart diseases.11,61 Electrocardiographic-gated cardiac CT can also accurately assess ventricular size, EF, and wall motion abnormalities, but it is accompanied with ionizing radiation.13–15 Radionuclide ventriculog- raphy is highly reproducible for measurement of LVEF, although it also exposes the patient to ion- izing radiation.12
    2. CMR provides noninvasive characterization of the myocardium that may provide insights into HF cause.62 Late-gadolinium enhancement, reflect- ing fibrosis and damaged myocardium, can iden- tify acute and chronic MI.63,64 and identify HF caused by CAD65,66 Patterns of late-gadolinium enhancement or specific T-1 and T-2 techniques can suggest specific infiltrative and inflammatory cardiomyopathies, such as myocarditis, sarcoid- osis, Fabry disease, Chagas disease, noncom- paction, iron overload, and amyloidosis.16,20,22,67 T-1 mapping techniques allow for measurement

of interstitial space characteristics and extracel- lular volume fraction and provides diagnostic and prognostic information.19,21–23,68–71 The presence of delayed hyperenhancement has been associ- ated with worse outcomes and can provide risk stratification.72–77 Although registry data show that CMR findings commonly impact patient care management and provide diagnostic information in patients with suspected myocarditis or cardio- myopathy,17,18 a strategy of routine screening with CMR in patients with nonischemic cardiomyopathy was not shown to yield more specific HF causes than a strategy of selective CMR strategy based on echocardiographic and clinical findings in a recent trial.78

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* + 1. HF is often caused by coronary atherosclerosis,79 and evaluation for ischemic heart disease can help in determining the presence of significant coronary artery disease (CAD). Noninvasive stress imag- ing with echocardiography or nuclear scintigraphy can be helpful in identifying patients likely to have obstructive CAD.24,25 Invasive or computed tomog- raphy coronary angiography can detect and char- acterize extent of CAD.26,27
    2. CAD is a leading cause of HF79 and myocardial ischemia may contribute to new or worsening HF symptoms. Noninvasive testing (ie, stress echo- cardiography, SPECT, CMR, or PET) may be con- sidered for detection of myocardial ischemia to help guide coronary revascularization decisions. Multiple nonrandomized, observational studies have reported improved survival with revascular- ization in patients with viable but dysfunctional myocardium.28,30–32 Despite these observational data, RCTs have not shown that viability imag- ing improves guidance of revascularization to a reduction of adverse cardiovascular out- comes.80–82 A prespecified viability substudy of the STICH (Surgical Treatment for Ischemic Heart Failure) trial showed that the presence of myocardial viability did not determine the long- term benefit from surgical revascularization in patients with ischemic cardiomyopathy.81,82 Of note, a relatively small number of individuals enrolled in the STICH substudy did not have viability, which may limit the power of the study. Although these data do not support the concept of routine viability assessment before revascu- larization, myocardial viability is used as one of the tools to inform decisions regarding revas- cularization in patients with high surgical risk or with complex medical problems.
    3. Repeat noninvasive imaging of cardiac struc- ture and function for routine surveillance is rarely appropriate in the absence of a change in clinical status or treatment interventions.11,83

## Invasive Evaluation

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| **Recommendations for Invasive Evaluation**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **2a** | **B-NR** | 1. In patients with HF, endomyocardial biopsy may be useful when a specific diagnosis is sus- pected that would influence therapy.1,2 |
| **2a** | **C-EO** | 2. In selected patients with HF with persistent or worsening symptoms, signs, diagnostic param- eters, and in whom hemodynamics are uncer- tain, invasive hemodynamic monitoring can be useful to guide management. |
| **3: No**  **Benefit** | **B-R** | 3. In patients with HF, routine use of invasive hemodynamic monitoring is not recom- mended.3,4 |
| **3: Harm** | **C-LD** | 4. For patients undergoing routine evaluation of HF, endomyocardial biopsy should not be per- formed because of the risk of complications.5,6 |

**Synopsis**

Invasive evaluation of patients with HF may provide important clinical information to determine the cause of HF and treatment options. Routine right heart cath- eterization does not provide sufficient information to guide treatment decisions.3,4 However, hemodynamic evaluation with right heart catheterization and moni- toring in the setting of acute respiratory distress, sys- temic hypoperfusion including cardiogenic shock, or when hemodynamics are uncertain, may guide treat- ment decisions. Coronary angiography may be useful in patients who are candidates for revascularization7–9 (see Section 4.4, “Evaluation with Cardiac Imaging,” for recommendations). Endomyocardial biopsy may be ad- vantageous in patients with HF in which a histological diagnosis, such as amyloidosis or myocarditis, may in- fluence treatment decisions.1,2

## Recommendation-Specific Supportive Text

* + 1. Endomyocardial biopsy may be useful when seeking a specific diagnosis that would influence treatment, and biopsy should thus be considered in patients with rapidly progressive clinical HF or worsening ventricular dysfunction that persists despite appro- priate medical treatment. Endomyocardial biopsy should also be considered in patients suspected of having acute cardiac rejection status after heart transplantation or having myocardial infiltrative processes. A specific example is to determine treatment for light chain (AL) amyloidosis or trans- thyretin amyloidosis.5 Additional indications for endomyocardial biopsy include patients with rap- idly progressive and unexplained cardiomyopathy and those in whom active myocarditis, especially giant cell myocarditis, is being considered.1

* + 1. Right-heart catherization in patients in acute HF. The ESCAPE (Evaluation Study of Congestive Heart Failure and Pulmonary Artery Catheterization Effectiveness) trial found that routine use of PA catheter monitoring for patients with HF did not provide benefit.3 However, invasive hemodynamic evaluation or monitoring can be useful to guide management in carefully selected patients with acute HF who have persistent symptoms despite treat- ment. This includes patients whose fluid status, perfusion, or systemic or pulmonary vascular resistance is uncertain whose systolic blood pressure (SBP) remains low, or is associated with symptoms, despite initial treatment; whose renal function is worsening with therapy; or who require parenteral vasoactive agents.

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* + 1. There has been no established role for routine or periodic invasive hemodynamic measure- ments in the management of HF. Most drugs used to treat HF are prescribed on the basis of their ability to improve symptoms or survival rather than their effect on hemodynamic vari- ables. The initial and target doses of these drugs are generally selected on the basis of controlled trial experience rather than changes produced in cardiac output or pulmonary capillary wedge pressure.3,4
    2. Patients with HF should not undergo routine endo- myocardial biopsy because of the risk of complica- tions that include perforation, cardiac tamponade, and thrombus formation, as well as limited diag- nostic yield.5,6

## Wearables and Remote Monitoring (Including Telemonitoring and Device Monitoring)

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| --- | --- | --- |
| **Recommendation for Wearables and Remote Monitoring (Including Telemonitoring and Device Monitoring)**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **2b** | **B-R** | 1. In selected adult patients with NYHA class III HF and history of a HF hospitalization in the past year or elevated natriuretic peptide levels, on maximally tolerated stable doses of GDMT with optimal device therapy, the use- fulness of wireless monitoring of PA pressure by an implanted hemodynamic monitor to  reduce the risk of subsequent HF hospitaliza- tions is uncertain.1–4 |
| **Value Statement: Uncertain Value (B-NR)** | | 2. In patients with NYHA class III HF with a HF hospitalization within the previous year, wireless monitoring of the PA pressure by an implanted hemodynamic monitor provides uncertain value.4–7 |

**Synopsis**

HF is a chronic condition punctuated by periods of insta- bility. Despite close longitudinal monitoring via in-person visits, event rates remain high, affording a potential role for remote monitoring strategies to improve clinical outcomes. Strategies tested in randomized trials include an implant- able PA pressure sensor (CardioMEMS), noninvasive tele- monitoring, or monitoring via existing implanted electronic devices (ICDs or CRT-Ds). Results from a single random- ized trial,1–3 and subsequent observational studies,8–10 sup- port consideration of an implantable PA sensor in selected patients with HF to reduce the risk of HF hospitalization. In contrast, a recent trial testing a PA pressure sensor did not meet its primary endpoint.4 Results from previous clini- cal trials do not support the alternative remote monitoring strategies (eg, noninvasive telemonitoring or remote moni- toring of physiological parameters such as patient activity, thoracic impedance, heart rate) for this purpose.11–18

## Recommendation-Specific Supportive Text

* + 1. The CHAMPION (CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in NYHA Class III Heart Failure patients) trial reported a significant 28% reduction of HF-related hospitalizations after 6 months in patients random- ized to an implanted PA pressure monitor compared with a control group.1 Patients had to have a HF hospitalization in the previous year and be on stable doses of a beta blocker and angiotensin-converting enzyme inhibitor (ACEi) (or angiotensin (II) recep- tor blocker [ARB]) if tolerated. The clinical benefit persisted after longer term follow-up and was seen in both subjects with reduced3 and preserved2 LVEF. However, CHAMPION was a nonblinded trial, and there was differential contact of study personnel with patients in the treatment arm, raising methodologi- cal concerns about the opportunity for bias to have influenced its results.19–21 In the recent GUIDE-HF (Haemodynamic-GUIDEed management of Heart Failure) study, hemodynamic-guided management of patients with NYHA class II to IV heart failure did not significantly reduce the composite endpoint rate of mortality and total HF events.4 The usefulness of noninvasive telemonitoring11,12,22,23 or remote moni- toring of physiological parameters13–18 (eg, patient activity, thoracic impedance, heart rate) via implanted electrical devices (ICDs or CRT-Ds) to improve clini- cal outcomes remains uncertain. Further study of these approaches is needed before they can be recommended for routine clinical care.
    2. Three model-based studies5–7 have evaluated the cost-effectiveness of wireless PA pressure monitor- ing using data from the CHAMPION-HF1 study of

the CardioMEMS device. All 3 studies estimated CardioMEMS implantation and monitoring increased survival and quality-adjusted life year (QALY) while increasing costs. Primarily based on differences regarding the expected magnitude of clinical benefit, 2 analyses5,7estimated the device provided high value while the third6 estimated intermediate value. These analyses had several important differences detailed in the evidence tables, including the model duration, QOL data, cost estimates, and assumptions regarding mortality. One analysis6 found the economic value of CardioMEMS implantation was highly dependent on its effect on mortality and duration of treatment ben- efit, both of which remain unclear. Cost-effectiveness studies incorporating data from GUIDE-HF4 have not been published. Additional data regarding clinical outcomes following CardioMEMS implantation will improve estimates of its economic value.

## Exercise and Functional Capacity Testing

their HF. NYHA class II includes patients who are comfortable at rest but have slight symptoms result- ing from HF (dyspnea, fatigue, lightheadedness) with ordinary activity. NYHA class III includes patients who are comfortable at rest but have symptoms of HF with less than ordinary activity. NYHA class IV includes patients who are unable to carry out any physical activity without symptoms and have symp- toms at rest. NYHA functional classification has been widely used in clinical practice, clinical trials, and clinical practice guidelines to determine candi- dacy for drug and device therapy. Limitations include its ability to be inconsistently assessed from 1 clini- cian to another, resulting in poor reproducibility.23

1. Many CPET variables have been associated with prognosis in patients with HF.4,5,12,14,16,24 Peak exer- cise oxygen consumption/oxygen uptake (VO2) is often used to risk stratify patients and make deci- sions about timing of advanced HF therapies, includ- ing heart transplantation and LVAD. In a landmark article,7 investigators divided patients referred for heart transplantation into groups based on their peak VO2.7 Patients with peak VO2 <14 mL/kg/min were listed for transplant, while those with higher peak VO2 values were deferred for being too well. Patients with peak VO2 >14 mL/kg/min who were deferred had 1- and 2-year survival of 94% and 84%, respectively, which was similar to survival after heart transplant.

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| --- | --- | --- |
| **Recommendations for Exercise and Functional Capacity Testing Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 1. In patients with HF, assessment and documen- tation of NYHA functional classification are recommended to determine eligibility for treat- ments.1–3 |
| **1** | **C-LD** | 2. In selected ambulatory patients with HF, cardiopulmonary exercise testing (CPET) is recommended to determine appropriateness of advanced treatments (eg, LVAD, heart transplant).4–8 |
| **2a** | **C-LD** | 3. In ambulatory patients with HF, performing a CPET or 6-minute walk test is reasonable to assess functional capacity.4,5,9–16 |
| **2a** | **C-LD** | 4. In ambulatory patients with unexplained dys- pnea, CPET is reasonable to evaluate the cause of dyspnea.17,18 |

As such, the authors proposed peak VO2 14 mL/

kg/min as a cutoff to distinguish patients who may derive survival benefit from heart transplant.7 Patients tolerating beta blockers may have improved survival

with an equivalent VO2 compared with patients who

do not tolerate beta blockers.25,26 For patients on

beta blockers, a peak VO2 12 mL/kg/min has been

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## Synopsis

Functional impairment and exercise intolerance are com- mon in HF. CPET and the 6-minute walk test are standard- ized, reliable, and reproducible tests to quantify functional capacity.19–22 The NYHA functional classification can be used to grade the severity of functional limitation based on patient report of symptoms experienced with activity1 and is used to define candidates for certain treatments.

## Recommendation-Specific Supportive Text

* 1. NYHA functional classification is an ordinal, categor- ical variable (I-IV) that is used to document functional limitation in patients with cardiac disease, including HF.1 In HF, NYHA functional class I includes patients with no limitations in physical activity resulting from

suggested as a more appropriate cutoff to consider cardiac transplant listing.8

1. Objective assessment of exercise capacity with CPET can be useful in the clinical management of patients with HF. Although CPET remains the gold standard measure of exercise capacity, limita- tions to more widespread use include need for spe- cial equipment and trained personnel, which leads to lack of availability at many hospitals and clinics. Furthermore, it is not well tolerated by some patients. The 6-minute walk test is an alternative way to mea- sure exercise capacity that is widely available and well tolerated by patients. It entails walking for 6 minutes on a measured flat course, and patients are allowed to slow down or stop if needed. A system- atic review of 14 studies found that the 6-minute walk test results correlated moderately with peak VO2 levels and were a reliable and valid indicator of functional capacity in patients with HF who did not walk >490 m.8 Distance walked in the 6-minute walk test has been associated with prognosis in HF

across multiple studies.9–13,15,16,27 A cutoff of <300 m roughly correlates to patients with NYHA class III to IV symptoms and is associated with worse 3-year survival free of heart transplant (62% versus 82% for those walking 300 m).27

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1. Dyspnea is a complex symptom that can reflect abnormalities in a number of different systems and can be influenced by psychological and envi- ronmental factors. CPET involves having patients perform a treadmill (or stationary bicycle) exer- cise test, while also performing ventilatory gas exchange measurements.28 CPET enables the comprehensive assessment of multiple physiologi- cal measures that can impact exercise capacity and contribute to dyspnea. It provides analysis of gas exchange and yields measures of oxygen uptake (VO2), carbon dioxide output, and ventilation. These measures can be integrated with standard exercise testing variables, such as heart rate, blood pres- sure, electrocardiographic findings, and symptoms to provide insights into the physiologic mechanisms underlying a patient’s dyspnea. In particular, CPET can help to distinguish respiratory versus cardiac etiologies of dyspnea. If exercise capacity is dimin- ished but cardiopulmonary responses are normal, other causes of dyspnea, such as metabolic abnor- malities and deconditioning, should be considered.

## Initial and Serial Evaluation: Clinical Assessment: HF Risk Scoring

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| **Recommendation for Initial and Serial Evaluation: Clinical Assessment: HF Risk Scoring**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **2a** | **B-NR** | 1. In ambulatory or hospitalized patients with HF, validated multivariable risk scores can be useful to estimate subsequent risk of mortality.1–14 |

**Synopsis**

Clinicians should routinely assess a patient’s risk for an adverse outcome to guide discussions on prognosis, goals of care, and treatment decisions. Several predictive models of outcomes of patients with HF have been de- veloped and validated using data from clinical trials, regis- tries, and population-based cohorts. The best performing models have focused on predicting short- and long-term mortality, whereas predictive models for hospitalization or readmission for HF have generally had poor or modest discrimination. Predictive models may also assess the risk of incident HF among the general population and should be considered in the prevention of HF. In the course of standard evaluation, clinicians should routinely assess the patient’s potential for adverse outcome, because accurate

##### Table 8. Selected Multivariable Risk Scores to Predict Outcome in HF

|  |  |  |
| --- | --- | --- |
| **Risk Score** | **Reference/Link** | **Year Published** |
| Chronic HF | | |
| All Patients With Chronic HF | | |
| Seattle Heart Failure Model | 2  [https://depts.washing-](https://depts.washington.edu/shfm/?width=1440&height=900) [ton.edu/shfm/?width](https://depts.washington.edu/shfm/?width=1440&height=900)  [=1440&height=900](https://depts.washington.edu/shfm/?width=1440&height=900) 15 | 2006 |
| Heart Failure Survival Score | 1 | 1997 |
| MAGGIC | 3  [http://www.heartfailure-](http://www.heartfailurerisk.org/) [risk.org/](http://www.heartfailurerisk.org/)  16 | 2013 |
| CHARM Risk Score | 4 | 2006 |
| CORONA Risk Score | 5 | 2009 |
| Specific to Chronic HFrEF | | |
| PARADIGM-HF | 6 | 2020 |
| HF-ACTION | 7 | 2012 |
| GUIDE-IT | 8 | 2019 |
| Specific to Chronic HFpEF | | |
| I-PRESERVE Score | 9 | 2011 |
| TOPCAT | 10 | 2020 |
| Acutely Decompensated HF | | |
| ADHERE Classification and Re- gression Tree (CART) Model | 11 | 2005 |
| AHA Get With The Guidelines Score | 12  [https://www.mdcalc.](https://www.mdcalc.com/gwtg-heart-failure-risk-score) [com/gwtg-heart-failure-](https://www.mdcalc.com/gwtg-heart-failure-risk-score) [risk-score](https://www.mdcalc.com/gwtg-heart-failure-risk-score)  17 | 2010, 2021 |
| EFFECT Risk Score | 13  [http://www.ccort.ca/](http://www.ccort.ca/Research/CHFRiskModel.aspx) [Research/CHFRisk-](http://www.ccort.ca/Research/CHFRiskModel.aspx) [Model.aspx](http://www.ccort.ca/Research/CHFRiskModel.aspx)  18 | 2003, 2016 |
| ESCAPE Risk Model and Discharge Score | 14 | 2010 |

ADHERE indicates Acute Decompensated Heart Failure National Registry; AHA, American Heart Association; ARIC, Atherosclerosis Risk in Communi- ties; CHARM, Candesartan in Heart failure-Assessment of Reduction in Mor- tality and morbidity; CORONA, Controlled Rosuvastatin Multinational Trial in Heart Failure; EFFECT, Enhanced Feedback for Effective Cardiac Treatment; ESCAPE, Evaluation Study of Congestive Heart Failure and Pulmonary Artery Catheterization Effectiveness; GUIDE-ID, Guiding Evidence-Based Therapy Using Biomarker Intensified Treatment; HF, heart failure; HFpEF, heart failure with preserved ejection fraction; HF-ACTION, Heart Failure: A Controlled Trial Investigating Outcomes of Exercise Training MAGGIC Meta-analysis Global Group in Chronic Heart Failure; I-PRESERVE, Irbesartan in Heart Failure with Preserved Ejection Fraction Study; PCP-HF, Pooled Cohort Equations to Pre- vent HF; and TOPCAT, Treatment of Preserved Cardiac Function Heart Failure with an Aldosterone Antagonist trial.

risk stratification may help guide therapeutic decision- making, including a more rapid transition to advanced HF therapies. Several methods objectively assess risk (Table 8), including biomarker testing, as well as various multivariable clinical risk scores, and some that include

machine learning.1–14 These risk scores are for use in am- bulatory, hospitalized patients, and the general population.

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## Recommendation-Specific Supportive Text

* + 1. For HF, there are several clinical models to consider that include the spectrum of HF based on EF and clin- ical setting. For chronic HF, the Seattle Heart Failure Model,2 the Heart Failure Survival score,1 and the MAGGIC score3 have commonly been used to provide estimates of survival. The MAGGIC predictive model may be quite useful given its derivation and valida- tion across multiple clinical trials and cohorts, includ- ing more recent studies. For chronic HFrEF, there are additional models that include other clinical variables, including exercise capacity7 and natriuretic peptide levels.8 Likewise, for chronic HFpEF there are more specific predictive models for that population derived from clinical trial data.9,10 In acute HF, several clinical models may be used to predict short-term survival.11–13

# STAGE A (PATIENTS AT RISK FOR HF)

## Patients at Risk for HF (Stage A: Primary Prevention)

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| **Recommendations for Patients at Risk for HF (Stage A: Primary Prevention)**  **Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In patients with hypertension, blood pressure should be controlled in accordance with GDMT for hypertension to prevent symptomatic HF.1–9 |
| **1** | **A** | 2. In patients with type 2 diabetes and either established CVD or at high cardiovascular risk, SGLT2i should be used to prevent hospitaliza- tions for HF.10–12 |
| **1** | **B-NR** | 3. In the general population, healthy lifestyle hab- its such as regular physical activity, maintaining normal weight, healthy dietary patterns, and avoiding smoking are helpful to reduce future risk of HF.13–21 |
| **2a** | **B-R** | 4. For patients at risk of developing HF, natriuretic peptide biomarker–based screening followed by team-based care, including a cardiovascular specialist optimizing GDMT, can be useful to prevent the development of LV dysfunction (systolic or diastolic) or new-onset HF.22,23 |
| **2a** | **B-NR** | 5. In the general population, validated multivari- able risk scores can be useful to estimate sub- sequent risk of incident HF.24–26 |

**Synopsis**

Healthy lifestyle habits such as maintaining regular phys- ical activity; normal weight, blood pressure, and blood glucose levels; healthy dietary patterns, and not smok- ing reduce primordial risk and have been associated with a lower lifetime risk of developing HF.13–21,27 The AHA/

ACC primary prevention guidelines provide recommen- dations for diet, physical activity, and weight control, all of which have been associated with the risk of HF.28 Blood pressure is an important risk factor for HF, and a treatment goal of <130/80 mm Hg is recommended for those with a CVD risk of 10%.29,30 Multiple RCTs have found that patients with diabetes and CVD without HF have improved survival and reduced HF hospitalizations with SGLT2i.31 Patients at risk for HF screened with BNP or NT-proBNP followed by collaborative care, diagnostic evaluation, and treatment in those with elevated levels can reduce combined rates of LV systolic dysfunction, diastolic dysfunction, and HF.22,23 See Figure 5 for COR 1 and 2a for stage A (at risk for HF) and stage B (pre-HF).

## Recommendation-Specific Supportive Text

* + 1. Elevated systolic and diastolic blood pressure are major risk factors for the development of symp- tomatic HF.8,9,32 Many trials have shown that hyper- tension control reduces the risk of HF.1–7 Although the magnitude of benefit varies with the patient population, target blood pressure reduction, and HF criteria, effective hypertension treatment invari- ably reduces HF events. In the SPRINT (Systolic Blood Pressure Intervention Trial) trial, control to an SBP goal <120 mm Hg decreased incident HF by 38% and mortality by 23% compared with an SBP goal of <140 mm Hg6,7 A meta-analysis showed that blood pressure control was associ- ated with an approximately 40% reduction in HF events.5 Therefore, SBP and diastolic blood pres- sure should be controlled in accordance with pub- lished clinical practice guidelines.30
    2. Multiple RCTs in patients with type 2 diabetes and at risk for, or with established CVD or at high risk for CVD, have shown that SGLT2i prevent HF hospital- izations compared with placebo.10–12 The benefit for reducing HF hospitalizations in these trials predomi- nantly reflects primary prevention of symptomatic HF, because only approximately 10% to 14% of partici- pants in these trials had HF at baseline. The mecha- nisms for the improvement in HF events have not been clearly elucidated but seem to be independent of glucose lowering. Proposed mechanisms include reductions in plasma volume, cardiac preload and afterload, alterations in cardiac metabolism, reduced arterial stiffness, and interaction with the Na+/H+ exchanger.33,34 SGLT2i are generally well tolerated, but these agents have not been evaluated in those with severe renal impairment (estimated glomerular filtration rate [eGFR] <25 mL/min/1.73 m2).35
    3. Greater adherence to healthy lifestyle habits such as regular physical activity, avoiding obe- sity, maintaining normal blood pressure and

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**Figure 5. Recommendations (Class 1 and 2a) for Patients at Risk of HF (Stage A) and Those With Pre-HF (Stage B).** Colors correspond to COR in Table 2. COR 1 and COR 2a for patients at risk for HF (stage A) and those with pre-HF (stage B) are shown. Management strategies implemented in patients at risk for HF (stage A) should be continued though stage B. ACEi indicates angiotensin- converting enzyme inhibitor; ARB, angiotensin receptor blocker; BP, blood pressure; COR, Class of Recommendation; CVD, cardiovascular

disease; HF, heart failure; ICD, implantable cardioverter-defibrillator; LVEF, left ventricular ejection fraction; MI, myocardial infarction; and SGLT2i, sodium glucose cotransporter 2 inhibitor.

blood glucose, not smoking, and healthy dietary patterns have been associated with a lower life- time risk of HF and greater preservation of car- diac structure.13–16,27 Healthful eating patterns, particularly those that are based more on con- sumption of foods derived from plants, such as the Mediterranean, whole grain, plant-based diet and the DASH (Dietary Approaches to Stop Hypertension) diet, are inversely associated with incident HF and may offer some protection against HF development.17–21

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* + 1. A large-scale unblinded single-center study (STOP-HF [The St Vincent’s Screening to Prevent Heart Failure])22 of patients at risk of HF (identi- fied by the presence of hypertension, diabetes, or known vascular disease) but without established LV systolic dysfunction or symptomatic HF at base- line found that screening with BNP testing and then intervening on those with levels of 50 pg/ mL (performing echocardiography and referral to a cardiovascular specialist) reduced the composite endpoint of asymptomatic LV dysfunction (systolic

##### Table 9. Selected Multivariable Risk Scores to Predict De- velopment of Incident HF

|  |  |  |
| --- | --- | --- |
| **Recommendations for Management of Stage B: Preventing the Syndrome of Clinical HF in Patients With Pre-HF (Continued)** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 2. In patients with a recent or remote history of MI or ACS, statins should be used to prevent symptomatic HF and adverse cardiovascular events.5–9 |
| **1** | **B-R** | 3. In patients with a recent MI and LVEF 40% who are intolerant to ACEi, ARB should be used to prevent symptomatic HF and reduce mortal- ity.10 |
| **1** | **B-R** | 4. In patients with a recent or remote history of MI or acute coronary syndrome (ACS) and LVEF  40%, evidence-based beta blockers should  be used to reduce mortality.11–13 |
| **1** | **B-R** | 5. In patients who are at least 40 days post-MI with LVEF 30% and NYHA class I symptoms while receiving GDMT and have reasonable expectation of meaningful survival for >1 year, an ICD is recommended for primary prevention of sudden cardiac death (SCD) to reduce total mortality.14 |
| **1** | **C-LD** | 6. In patients with LVEF 40%, beta blockers should be used to prevent symptomatic HF.12,13 |
| **3: Harm** | **B-R** | 7. In patients with LVEF <50%, thiazolidinediones should not be used because they increase the risk of HF, including hospitalizations.15 |
| **3: Harm** | **C-LD** | 8. In patients with LVEF <50%, nondihydropyri- dine calcium channel blockers with negative inotropic effects may be harmful.16,17 |

|  |  |  |
| --- | --- | --- |
| **Risk Score** | **Reference** | **Year Published** |
| Framingham Heart Failure Risk Score | 24 | 1999 |
| Health ABC Heart Failure Score | 25 | 2008 |
| ARIC Risk Score | 26 | 2012 |
| PCP-HF | 29 | 2019 |

ARIC indicates Atherosclerosis Risk in Communities; HF, heart failure; and PCP-HF, Pooled Cohort Equations to Prevent HF.

or diastolic) with or without newly diagnosed HF.22 Similarly, in another small, single-center RCT, accelerated uptitration of RAAS antagonists and beta blockers reduced cardiac events in patients with diabetes and elevated NT-proBNP levels but without cardiac disease at baseline.23

* + 1. Incident HF may be predicted from different mod- els, including those derived from diverse populations (Table 9). The PCP-HF (Pooled Cohort equations to Prevent HF) model provides race- and sex- specific 10-year risk equations from 7 community- based cohorts with at least 12 years of follow-up.29 Predictors of HF included in the race- and sex-spe- cific models were age, blood pressure (treated or untreated), fasting glucose (treated or untreated), body mass index, cholesterol, smoking status, and QRS duration. Models can be applied to the clinical setting of interest, with clinical trial models potentially less generalizable to registry- or population-based models. In addition, predictive models provide the average estimate of risk derived from a population, and individual risk may vary.36 The integration of risk scores into clinical practice have shown improved outcomes. As data generation increases from elec- tronic health records and digital sources, advanced methods with machine learning are expected to pro- liferate the development of risk prediction models. Machine learning models are often not externally validated, and their performance may vary based on the population and clinical setting.37 Patient popula- tions change over time, and models may need to be recalibrated periodically.

## Synopsis

In general, all recommendations for patients with stage A HF also apply to those with stage B HF. Stage B (pre- HF) represents a phase of clinically asymptomatic struc- tural and functional cardiac abnormalities that increases the risk for symptomatic HF.18–21 Identifying individu- als with stage B HF provides an opportunity to initiate lifestyle modification and pharmacological therapy that may prevent or delay the transition to symptomatic HF (stage C/D). Several ACC/AHA clinical practice guide- lines address appropriate management of patients with stage B HF (Table 10). Although multiple studies high- light the increased HF risk associated with asymptomatic LV systolic19,20,22–26 and diastolic dysfunction identified by noninvasive imaging,19,26–30 beneficial pharmacotherapy

for asymptomatic LV systolic dysfunction, such as inhibi-

# STAGE B (PATIENTS WITH PRE-HF)

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## Management of Stage B: Preventing the Syndrome of Clinical HF in Patients With Pre-HF

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| **Recommendations for Management of Stage B: Preventing the Syndrome of Clinical HF in Patients With Pre-HF**  **Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In patients with LVEF 40%, ACEi should be used to prevent symptomatic HF and reduce mortality.1–4 |

tors of the renin-angiotensin system and beta blockers, have been predominantly observed in individuals with depressed LVEF (LVEF <35%–40%).1–4,11–13 Studies of specific treatments to alter the onset of HF in the set- ting of asymptomatic cardiac dysfunction with preserved LVEF (eg, abnormalities of myocardial deformation or diastolic dysfunction) have been limited. Several comor- bid conditions, including diabetes, obesity, and hyperten- sion, have been associated with asymptomatic LV dys- function27,28,30,31 and with progression of asymptomatic LV dysfunction to symptomatic HF.27 Accordingly, these comorbidities are controlled according to current clinical

##### Table 10. Other ACC/AHA Clinical Practice Guidelines Addressing Patients With Stage B HF

|  |  |
| --- | --- |
| **Consideration** | **Reference** |
| Patients with an acute MI who have not developed HF symptoms treated in accordance with GDMT | 2013 ACCF/AHA Guideline for the Management of ST-Elevation Myocardial Infarction51  2014 AHA/ACC Guideline for the Management of Patients With Non–ST-Elevation Acute Coro- nary Syndromes52 |
| Coronary revascularization for patients without symptoms of HF in accordance with GDMT | 2015 ACC/AHA/SCAI Focused Update on Primary Percutaneous Coronary Intervention for Pa- tients With ST-Elevation Myocardial Infarction: An Update of the 2011 ACCF/AHA/SCAI Guideline for Percutaneous Coronary Intervention and the 2013 ACCF/AHA Guideline for the Management of ST-Elevation Myocardial Infarction53 (This guideline has been replaced by Lawton, 2021.54)  2014 ACC/AHA/AATS/PCNA/SCAI/STS Focused Update of the Guideline for the Diagnosis and Management of Patients With Stable Ischemic Heart Disease55  2011 ACCF/AHA Guideline for Coronary Artery Bypass Graft Surgery56 (This guideline has been replaced by Lawton, 2021.54) |
| Valve replacement or repair for patients with hemodynamically significant valvular stenosis or regurgitation and no symptoms of HF in accordance with GDMT | 2020 ACC/AHA Guideline for the Management of Patients With Valvular Heart Disease.57,58 |
| Patients with congenital heart disease that may increase the risk for the development of HF | 2018 AHA/ACC Guideline for the Management of Adults With Congenital Heart Disease59 |

AATS indicates American Association for Thoracic Surgery; ACC, American College of Cardiology; ACCF, American College of Cardiology Foundation; AHA, American Heart Association; GDMT, guideline-directed medical therapy; HF, heart failure; MI, myocardial infarction; PCNA, Preventive Cardiovascular Nurses As- sociation; SCAI, Society for Cardiovascular Angiography and Interventions; and STS, The Society of Thoracic Surgeons.

practice guidelines. The benefits of mineralocorticoid receptor antagonists (MRA) after MI have mostly been shown in patients with symptomatic HFrEF.32–34

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ARNi have not been well studied in stage B HF. The PARADISE-MI (Prospective ARNi vs. ACE inhibi- tor trial to DetermIne Superiority in reducing heart failure Events after Myocardial Infarction) study35 will report the efficacy and safety of sacubitril/valsartan in patients after acute MI, with LVEF 40 and/or pulmo- nary congestion, plus an additional risk-enhancing fac- tor, compared with ramipril.

## Recommendation-Specific Supportive Text

* + - * 1. ACEi have been shown to impede maladaptive remod- eling after acute MI in patients with reduced LVEF.36,37 In survivors of acute MI with asymptomatic LV dys- function (LVEF <35%–40%), RCTs have shown that ACEi reduced mortality, HF hospitalizations, and progression to severe HF compared with placebo.2,4 Similarly, in those individuals with asymptomatic LV dysfunction in the SOLVD (Studies of Left Ventricular Systolic Dysfunction) prevention trial, which included approximately 20% without ischemic heart disease, enalapril was associated with reduced HF hospitaliza- tion and mortality compared with placebo.1,3
        2. In multiple RCTs,42 statins have been shown to pre- vent adverse CAD events in patients with an MI, ACS, and with high cardiovascular risk. These trials have also shown that statin therapy reduces the risk of incident HF.5–9 A meta-analysis of 6 RCTs of >110 000 patients with an ACS showed that intensive statin therapy reduced hospitalizations for HF.5 A subsequent, larger collaborative meta- analysis of up to 17 major primary and secondary prevention RCTs showed that statins reduced HF

hospitalization.42 These data support the use of statins to prevent symptomatic HF and cardiovas- cular events in patients with acute MI or ACS.

* + - * 1. Two major trials have compared ARB with ACEi after MI. The VALIANT (Valsartan in Acute Myocardial Infarction) trial, which included approximately 25% of patients with asymptomatic LV dysfunction, showed that the benefits of valsartan on mortality and other adverse cardiovascular outcomes were compa- rable to captopril.10,38 In the OPTIMAAL (Optimal Trial in Myocardial Infarction with the Angiotensin II Antagonist Losartan) trial, losartan did not meet the noninferiority criteria for mortality compared with captopril.39 It has been hypothesized that the lower dose of losartan (50 mg daily) in the OPTIMAAL trial may have contributed to the greater differ- ence than those seen with valsartan in VALIANT.40 No clinical trials have specifically evaluated ARB in patients with asymptomatic reduced LVEF in the absence of previous MI. Although ARB are alterna- tives for patients with ACEi-induced angioedema, caution is advised because some patients have also developed angioedema with ARB.
        2. Current evidence supports the use of beta block- ers to improve adverse cardiac remodeling and outcomes in patients with asymptomatic reduced LVEF after MI. Among patients with a recent MI and reduced LVEF, carvedilol reduced maladaptive remodeling41 and reduced mortality compared with placebo.11 Among patients with asymptomatic LV systolic dysfunction in the SOLVD prevention trial (which included 80% with previous MI) and the SAVE (Survival and Ventricular Enlargement) trial, secondary analyses showed that the administra- tion of beta blockers in addition to ACEi reduced mortality and hospitalization.12,13

* + - * 1. The Framingham studies have shown a 60% increased risk of death in patients with asymptomatic low LVEF compared with those with normal LVEF, and almost half of these patients remained free of HF before their death.25 MADIT-II (Multicenter Automatic Defibrillator Implantation Trial II) showed a 31% relative risk reduction in all-cause mortality in patients with post-MI with LVEF 30% receiv- ing a prophylactic ICD compared with standard of care.14 These findings provided justification for the broad adoption of ICDs for primary prevention of SCD in the post-MI setting with reduced LVEF, even in the absence of HF symptoms.

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* + - * 1. Although beta blockers have been shown to improve outcomes in patients with symptomatic HFrEF and in patients with reduced LVEF after MI,11 few data exist regarding the use of beta blockers in asymp- tomatic patients with depressed LVEF without a his- tory of MI. There is evidence to support the role of beta blockers to prevent adverse LV remodeling in asymptomatic patients with LV systolic dysfunction, including those with nonischemic cause.43 Also, in a post hoc analysis of the SOLVD prevention trial, which included approximately 20% of participants with nonischemic HF cause, beta blockers were associated with a reduction in the risk of death and in death or hospitalization for symptomatic HF in those patients randomized to enalapril, a finding that was not seen in the placebo group.12 Given the long- term benefits of beta blockers to reduce HF hos- pitalizations in patients with symptomatic HFrEF,44 beta-blocker therapy is recommended to prevent symptomatic HF in patients with reduced LVEF.
        2. Thiazolidinediones have been associated with fluid retention and increased rates of HF in RCTs of patients with type 2 diabetes who were predominantly free of symptomatic HF at baseline.47–49 In a smaller RCT of patients with more severely symptomatic HFrEF, pioglitazone was associated with increased rates of HF hospitalization compared with placebo.50 In patients with more mild symptoms (NYHA class I to II) but with depressed LVEF,15 rosiglitazone was associated with more fluid-related events, includ- ing worsening edema and need for increased HF medications.15 Given the evidence, thiazolidinediones should be avoided in patients with reduced LVEF.
        3. Nondihydropiridine calcium channel blockers diltia- zem and verapamil are myocardial depressants and generally not tolerated in HF. In previous studies of patients with HF or reduced LVEF after acute MI, dil- tiazem was associated with increased risk of HF,16,17 although in a smaller study of patients with non- ischemic cardiomyopathy, diltiazem had no impact on mortality.45 Verapamil had no impact on survival or major cardiovascular events after acute MI.46 Although not specifically tested in asymptomatic

patients with low LVEF, nondihydropyridine calcium channel blockers may be harmful in this population because of their negative inotropic effects.

# STAGE C HF

## Nonpharmacological Interventions

* + 1. ***Self-Care Support in HF***

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| --- | --- | --- |
| **Recommendations for Nonpharmacological Interventions: Self-Care Support in HF**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. Patients with HF should receive care from multidisciplinary teams to facilitate the imple- mentation of GDMT, address potential barriers to self-care, reduce the risk of subsequent rehospitalization for HF, and improve survival.1–4 |
| **1** | **B-R** | 2. Patients with HF should receive specific edu- cation and support to facilitate HF self-care in a multidisciplinary manner.2,5–9 |
| **2a** | **B-NR** | 3. In patients with HF, vaccinating against respi- ratory illnesses is reasonable to reduce mor- tality.10–16 |
| **2a** | **B-NR** | 4. In adults with HF, screening for depression,17,18 social isolation,19–22 frailty,23,24 and low health literacy25,26 as risk factors for poor self-care is reasonable to improve management. |

## Synopsis

Because of the complexity of HF management and coordination of other health and social services re- quired, HF care is ideally provided by multidisciplinary teams27–30 that include cardiologists, nurses, and phar- macists who specialize in HF as well as dieticians, mental health clinicians, social workers, primary care clinicians, and additional specialists.31–33 Self-care in HF comprises treatment adherence and health main- tenance behaviors.34,35 Patients with HF should learn to take medications as prescribed, restrict sodium intake, stay physically active, and get vaccinations.36,37 They also should understand how to monitor for signs and symptoms of worsening HF, and what to do in response to symptoms when they occur.36,37 Knowledge alone is insufficient to improve self-care.38 Patients with HF need time and support to gain skills and overcome bar- riers to effective self-care.37 Measures listed as Class 1 recommendations for patients in stages A and B are recommended where appropriate for patients in stage

C. GDMT, as depicted in Figure 6, should be the main- stay of pharmacological therapy for HFrEF.

## Recommendation-Specific Supportive Text

1. In a meta-analysis of 30 RCTs, multidisciplinary interventions reduced hospital admission and

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##### Figure 6. Treatment of HFrEF Stages C and D.

Colors correspond to COR in Table 2. Treatment recommendations for patients with HFrEF are displayed. Step 1 medications may be started simultaneously at initial (low) doses recommended for HFrEF. Alternatively, these medications may be started sequentially, with sequence guided by clinical or other factors, without need to achieve target dosing before initiating next medication. Medication doses should be increased to target as tolerated. ACEi indicates angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; ARNi, angiotensin receptor-neprilysin inhibitor; COR, Class of Recommendation; CRT, cardiac resynchronization therapy; GDMT, guideline-directed medical therapy; ICD, implantable cardioverter-defibrillator; hydral-nitrates, hydralazine and isosorbide dinitrate; HFrEF, heart failure with reduced ejection fraction; LBBB, left bundle branch block; MCS, mechanical circulatory support; LVEF, left ventricular ejection fraction; MRA, mineralocorticoid receptor antagonist; NSR, normal sinus rhythm; NYHA, New York Heart Association; and SGLT2i, sodium-glucose cotransporter 2 inhibitor. \*Participation in investigational studies is appropriate for stage C, NYHA class II and III HF.

all-cause mortality.1 In a separate meta-analysis of 22 RCTs, specialized multidisciplinary team follow- up was associated with reduced HF hospitalizations and all-cause hospitalizations.2 In a recent meta- analysis of 22 RCTs, multidisciplinary interventions that included a pharmacist reduced HF hospitaliza- tions.3 In a recent Cochrane systematic review and meta-analysis of 43 RCTs, both case management (ie, active management of complex patients by case managers working in integrated care systems) and multidisciplinary interventions (ie, coordinated mul- tidisciplinary health care interventions and commu- nications) were shown to reduce all-cause mortality, all-cause readmission, and readmission for HF.4

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1. Meta-analyses of RCTs have shown that inter- ventions focused on improving HF self-care

significantly reduce the risk of HF-related hos- pitalization,2,5–8 all-cause hospitalization,2,8,9 and all-cause mortality,6,9 as well as improve QOL.5 Interventions that aim to improve self-care knowl- edge and skill,2,5,8 and those that focus on enhanc- ing medication adherence9 or reinforce self-care with structured telephone support,6,7 are effective in patients with HF. There is uncertainty whether mobile health–delivered educational interven- tions improve self-care in patients with HF.39 In a single RCT involving rural patients with HF, an educational intervention was shown to improve knowledge and self-care40 but did not significantly decrease the combined endpoint of cardiac death or HF hospitalization.41 In a recent pragmatic trial, a transitional care services program that included

self-care education improved discharge prepared- ness, quality of transition, and QOL but did not sig- nificantly improve clinical outcomes compared with usual care.42

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1. In propensity-adjusted models, influenza vacci- nation was associated with a significant reduc- tion in all-cause mortality among participants in PARADIGM-HF (Prospective Comparison of ARNi with ACEi to Determine Impact on Global Mortality and Morbidity in Heart Failure).14 In adjusted mod- els, influenza vaccination was associated with significant reductions in all-cause mortality and car- diovascular mortality12 in 1 registry study and was associated with significant reductions in all-cause mortality and the composite of all-cause mortality and cardiovascular hospitalizations in another large cohort study.11 In a self-controlled case series study of patients with HF, influenza vaccination was asso- ciated with a significantly lower risk of cardiovas- cular, respiratory, and all-cause hospitalization.43 In a meta-analysis of 16 studies of patients with CVD, influenza vaccination was associated with a lower risk of all-cause, cardiovascular mortality, and major adverse cardiovascular events compared with con- trol patients.15 In the Cardiovascular Health Study, pneumococcal vaccination was associated with sig- nificant reductions in incident HF, all-cause mortal- ity, and cardiovascular mortality.16 Patients with HF are uniquely susceptible to poor outcomes in the setting of SARS-CoV-2 infection44–47 and should be vaccinated against COVID-19.10
2. Many health and social factors are associated with poor HF self-care.36,37 (Table 11) but have also been linked to poor clinical outcomes and fundamen- tally change how education and support must be delivered. Depression is a risk factor for poor self- care,40 rehospitalization,17 and all-cause mortality18 among patients with HF. Interventions that focus on improving HF self-care have been reported to be effective among patients with moderate/ severe depression with reductions in hospitaliza- tion and mortality risk.5 Nonrandomized studies have provided evidence of a link between social isolation and mortality in patients with HF19,20 In a recent meta-analysis of 29 cohort studies, frailty was associated with an increased risk of all-cause mortality and hospitalization.23 Frailty also has been shown to impair self-care among elderly patients with HF.24 A recent meta-analysis of observational studies revealed social isolation to be common among adults with HF (ie, 37%) and associated with a 55% greater risk of HF-related rehospital- ization.21 Poor social support also has been shown in nonrandomized studies to be associated with lower HF self-care.22 A recent meta-analysis of observational studies showed that inadequate/

marginal health literacy is common among adults with HF (ie, 24%) and associated independently with the risk of mortality and hospitalization.25 Low literacy also is associated with poor HF self-care, as most interventions depend on both literacy and health literacy/numeracy.26

* + 1. ***Dietary Sodium Restriction***

|  |  |  |
| --- | --- | --- |
| **Recommendation for Dietary Sodium Restriction** | | |
| **COR** | **LOE** | **Recommendation** |
| **2a** | **C-LD** | 1. For patients with stage C HF, avoiding exces- sive sodium intake is reasonable to reduce congestive symptoms.1–6 |

## Synopsis

Restricting dietary sodium is a common nonpharmaco- logical treatment for patients with HF symptomatic with congestion, but specific recommendations have been based on low-quality evidence.7 Concerns about the qual- ity of data regarding clinical benefits or harm of sodium restriction in patients with HF include the lack of current pharmacological therapy, small samples without sufficient racial and ethnic diversity, questions about the correct threshold for clinical benefit, uncertainty about which sub- groups benefit most from sodium restriction,7,8 and seri- ous questions about the validity of several RCTs in this area.9–11 However, there are promising pilot trials of so- dium restriction in patients with HF3,5,6 The AHA currently recommends a reduction of sodium intake to <2300 mg/d for general cardiovascular health promotion12; how- ever, there are no trials to support this level of restriction in patients with HF.13 Sodium restriction can result in poor dietary quality with inadequate macronutrient and mi- cronutrient intake.14 Nutritional inadequacies have been associated with clinical instability,15–17 but routine supple- mentation of oral iron,18 thiamine,19 zinc,20 vitamin D,21 or multivitamins has not proven beneficial.22 The DASH diet is rich in antioxidants and potassium, can achieve sodium restriction without compromising nutritional adequacy when accompanied by dietary counseling,5 and may be associated with reduced hospitalizations for HF.23

## Recommendation-Specific Supportive Text

* + - 1. A registered dietitian- or nurse-coached interven- tion with 2 to 3 g/d sodium restriction improved NYHA functional class and leg edema in patients with HFrEF.1 In a nonrandomized study (>2.5 g/d versus <2.5 g/d), lower dietary sodium was associ- ated with worse all-cause mortality in patients with HFrEF.2 In small RCTs, aggressive sodium restriction (0.8 g/d) during hospitalization for acute decom- pensated HF has not reduced weight, congestion, diuretic use, rehospitalization, or all-cause mortality in patients with HFrEF24 or in patients with HFpEF.25

##### Table 11. Potential Barriers to Effective HF Self-Care and Example Interventions

|  |  |  |
| --- | --- | --- |
| **Potential Barrier** | **Example Screening Tools** | **Example Interventions** |
| Medical Barriers | | |
| Cognitive impairment48–50 | Mini-Cog  Mini-Mental State Examination (MMSE) Montreal Cognitive Assessment (MoCA) | Home health aide Home meal deliveries Adult day care  Geriatric psychiatry referral  Memory care support groups |
| Depression51,52 | Hamilton Depression Rating Scale (HAM-D)  Beck Depression Inventory-II (BDI-II) Patient Health Questionnaire-9 (PHQ-9) | Psychotherapy  Selective serotonin reuptake inhibitors Nurse-led support |
| Substance use disorders53 | Tobacco, Alcohol, Prescription medication, and other Sub- stance use (TAPS) | Referral to social work services and community support partners  Referral for addiction psychiatry consultation |
| Frailty54 | Fried frailty phenotype | Cardiac rehabilitation  Registered dietitian nutritionist evaluation for malnutrition |
| Social Barriers | | |
| Financial burden of HF treatments55 | COmprehensive Score for financial Toxicity–Functional As- sessment of Chronic Illness Therapy (COST-FACIT) | PharmD referral to review prescription assistance eligibilities |
| Food insecurity56,57 | Hunger Vital Sign, 2 items  US Household Food Security Survey Module, 6 items | Determine eligibility for the Supplemental Nutrition Assistance Program (SNAP)  Connect patients with community partners such as food pantries/ food banks  Home meal deliveries  Registered dietitian nutritionist evaluation for potential malnutrition |
| Homelessness or housing insecurity58–60 | Homelessness Screening Clinical Reminder (HSCR) | Referral to local housing services  Connect patients with community housing partners |
| Intimate partner violence or elder abuse61,62 | Humiliation, Afraid, Rape, Kick (HARK) questionnaire Partner Violence Screen (PVS)  Woman Abuse Screening Tool (WAST) | Referral to social work services and community support partners |
| Limited English proficiency or other language barriers63 | Routinely inquire in which language the patient is most comfortable conversing | Access to interpreter services covering a wide range of languag- es, ideally in person or, alternatively, via video platform  Printed educational materials in a range of appropriate languages |
| Low health literacy64 | Short Assessment of Health Literacy (SAHL)  Rapid Estimate of Adult Literacy in Medicine–Short Form (REALM-SF)  Brief Health Literacy Screen (BHLS), 3 items | Agency for Healthcare Research and Quality (AHRQ) Health Lit- eracy Universal Precautions Toolkit  Written education tools provided at sixth grade reading level or below Graphic educational documents |
| Social isolation or low social support65 | Patient-Reported Outcomes Measurement Information System (PROMIS) Social Isolation Short Form | Determine eligibility for home care services  Support group referral |
| Transport limitations | No validated tools currently available. | Referral to social work services  Determine eligibility for insurance or state-based transportation, or reduced-cost public transportation  Maximize opportunities for telehealth visits and remote monitoring |

HF indicates heart failure.

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A recent pilot RCT N=27) showed that providing patients with 1.5 g/d sodium meals can reduce uri- nary sodium and improve QOL but not improve clini- cal outcomes.3 Another recent pilot RCT (N=38) of

1.5 versus 2.3 g/d sodium resulted in sodium intake and improvement in BNP levels and QOL in the

1.5 g/d sodium intake arm5; the full trial is due to be completed in 2022. A third pilot RCT (N=66) of home-delivered 1.5 g/d meals showed favorable but nonsignificant trends toward improvement in clini- cal status and readmission rates.6 Moreover, results from RCTs have shown that reducing dietary sodium is difficult to achieve in patients with HF, even with prepared meals3 or home visits.26

#### *Management of Stage C HF: Activity, Exercise* Prescription, and Cardiac Rehabilitation

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| **Recommendations for Management of Stage C HF: Activity, Exercise Prescription, and Cardiac Rehabilitation**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. For patients with HF who are able to par- ticipate, exercise training (or regular physical activity) is recommended to improve functional status, exercise performance, and QOL.1–9 |
| **2a** | **B-NR** | 2. In patients with HF, a cardiac rehabilitation program can be useful to improve functional capacity, exercise tolerance, and health-related QOL.1,2,5,6,8 |

## Synopsis

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Exercise training in patients with HF is safe and has numerous benefits. In a major trial of exercise and HF, exercise training was associated with a reduction in CVD mortality or hospitalizations in the exercise train- ing group after adjustment for risk factors.1 Meta-anal- yses show that cardiac rehabilitation improves func- tional capacity, exercise duration, and health-related QOL. A cardiac rehabilitation program for patients with HF usually includes a medical evaluation, educa- tion regarding the importance of medical adherence, dietary recommendations, psychosocial support, and an exercise training and physical activity counseling program. Patients with HF on optimal GDMT, who are in stable medical condition and are able to participate in an exercise program, are candidates for an exercise rehabilitation program.10,11

## Recommendation-Specific Supportive Text

1. Evidence from RCTs indicates that exercise training improves functional status, exercise performance, and QOL in patients with HFrEF and HFpEF. In HF-ACTION, the largest randomized trial with exer- cise training in patients with HF,1 2331 patients with LVEF 35% (NYHA class II and III) were ran- domized to usual care versus supervised exercise training plus usual care. There were modest reduc- tions in all-cause mortality and hospitalization rates that did not reach significance by primary analysis but, after prespecified adjustment, were associated with reductions in cardiovascular mortality or HF hospitalizations.1 Many RCTs of exercise training in HF have been conducted, but the statistical power of most was low.2–5,9–13 Meta-analyses suggest that exercise training is associated with improvement in functional capacity, exercise duration, health- related QOL, and reduction in HF hospitalizations in patients with HFrEF as well as HFpEF.2–6,8,11,14,15 Most studies and meta-analyses have not shown significant changes in all-cause mortality.2,12,14–22 except for a few showing mortality benefit with longer follow-up6,7 Other benefits of exercise train- ing include improved endothelial function, blunted catecholamine spillover, increased peripheral oxy- gen extraction, and improvement in peak oxygen consumption.2–5,8,10–12,21
2. A formal cardiac rehabilitation program usually includes a medical evaluation, education regard- ing the importance of medical adherence, dietary recommendations, psychosocial support, and an exercise training and physical activity counseling program. Exercise-based cardiac rehabilitation has been associated with an improvement in func- tional capacity, exercise tolerance, the rate of over- all and HF-specific hospitalization, and improved

QOL.3,4,6,7,11,16,17 In a diverse population of older patients who were hospitalized for acute decom- pensated HF, an early, transitional, tailored, pro- gressive rehabilitation intervention that included multiple physical-function domains (strength, bal- ance, mobility, and endurance) initiated during, or early after hospitalization for HF, and continued after discharge, resulted in greater improvement in physical function than usual care.9

## Diuretics and Decongestion Strategies in Patients With HF

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| **Recommendations for Diuretics and Decongestion Strategies in Patients With HF**  **Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. In patients with HF who have fluid retention, diuretics are recommended to relieve conges- tion, improve symptoms, and prevent worsen- ing HF.1–5 |
| **1** | **B-NR** | 2. For patients with HF and congestive symptoms, addition of a thiazide (eg, metolazone) to treat- ment with a loop diuretic should be reserved for patients who do not respond to moderate- or high-dose loop diuretics to minimize electro- lyte abnormalities.6 |

**Synopsis**

Bumetanide, furosemide, and torsemide inhibit reabsorp- tion of sodium or chloride at the loop of Henle, whereas thiazide and thiazide-like diuretics act in the distal con- voluting tubule and potassium-sparing diuretics (eg, spi- ronolactone) in the collecting duct.7,8 Loop diuretics are the preferred diuretic agents for use in most patients with HF. Thiazide diuretics such as chlorthalidone or hydrochlorothiazide may be considered in patients with hypertension and HF and mild fluid retention. Metola- zone or chlorothiazide may be added to loop diuretics in patients with refractory edema unresponsive to loop di- uretics alone. Diuretics should be prescribed to patients who have evidence of congestion or fluid retention. In any patient with a history of congestion, maintenance diuretics should be considered to avoid recurrent symp- toms. The treatment goal of diuretic use is to eliminate clinical evidence of fluid retention, using the lowest dose possible to maintain euvolemia. With the exception of MRAs, the effects of diuretics on morbidity and mortality are uncertain.1–5 As such, diuretics should not be used in isolation but always combined with other GDMT for HF that reduces hospitalizations and prolongs survival. Table 12 lists oral diuretics recommended for use in the treatment of chronic HF. Hyponatremia complicates HF management. If reversing potential causes and free wa- ter restriction do not improve hyponatremia, vasopressin antagonists may be helpful in the acute management of

##### Table 12. Commonly Used Oral Diuretics in Treatment of Congestion for Chronic HF

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| --- | --- | --- | --- |
| **Drug** | **Initial Daily Dose** | **Maximum Total Daily Dose** | **Duration of Action** |
| Loop diuretics | | | |
| Bumetanide | 0.5–1.0 mg once or twice | 10 mg | 4–6 h |
| Furosemide | 20–40 mg once or twice | 600 mg | 6–8 h |
| Torsemide | 10–20 mg once | 200 mg | 12–16 h |
| Thiazide diuretics | | | |
| Chlorthiazide | 250–500 mg once or twice | 1000 mg | 6–12 h |
| Chlorthalidone | 12.5–25 mg once | 100 mg | 24–72 h |
| Hydrochlorothiazide | 25 mg once or twice | 200 mg | 6–12 h |
| Indapamide | 2.5 mg once | 5 mg | 36 h |
| Metolazone | 2.5 mg once | 20 mg | 12–24 h |

HF indicates heart failure.

volume overload to decrease congestion while maintain- ing serum sodium.

## Recommendation-Specific Supportive Text

1. Controlled trials with diuretics showed their effects to increase urinary sodium excretion, decrease physical signs of fluid retention, and improve symptoms, QOL, and exercise tolerance.1–5 Recent data from the nonrandomized OPTIMIZE-HF (Organized Program to Initiate Lifesaving Treatment in Hospitalized Patients with Heart Failure) regis- try revealed reduced 30-day all-cause mortality and hospitalization for HF with diuretic use com- pared with no diuretic use after hospital discharge for HF.9 The most commonly used loop diuretic for the treatment of HF is furosemide, but some patients respond more favorably to other agents in this category (eg, bumetanide, torsemide), poten- tially because of their increased oral bioavailabil- ity.10–12 In outpatients with HF, diuretic therapy is commonly initiated with low doses, and the dose is increased until urine output increases and weight decreases, generally by 0.5 to 1.0 kg daily. Patients may become unresponsive to high doses of diuretic drugs if they consume large amounts of dietary sodium, are taking agents that can block the effects of diuretics (eg, NSAIDs), or have sig- nificant impairment of renal function or perfusion.
2. Diuretic resistance can be overcome in several ways, including escalation of loop diuretic dose, intravenous administration of diuretics (bolus or continuous infusion),6 or combination of different diuretic classes.13–16 The use of a thiazide or thia- zide-like diuretic (eg, metolazone) in combination with a loop diuretic inhibits compensatory distal tubular sodium reabsorption, leading to enhanced

natriuresis. However, in a propensity-score matched analysis in patients with hospitalized HF, the addition of metolazone to loop diuretics was found to increase the risk for hypokalemia, hypo- natremia, worsening renal function, and mortality, whereas use of higher doses of loop diuretics was not found to adversely affect survival.17 Although randomized data comparing the 2 diuretic strate- gies are limited, the DOSE (Diuretic Optimization Strategies Evaluation) trial lends support for the use of high-dose intravenous loop diuretics.18

## Pharmacological Treatment\* for HFrEF

#### *Renin-Angiotensin System Inhibition With* ACEi or ARB or ARNi

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| --- | --- | --- |
| **Recommendations for Renin-Angiotensin System Inhibition With ACEi or ARB or ARNi**  **Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In patients with HFrEF and NYHA class II to III symptoms, the use of ARNi is recommended to reduce morbidity and mortality.1–5 |
| **1** | **A** | 2. In patients with previous or current symptoms of chronic HFrEF, the use of ACEi is beneficial to reduce morbidity and mortality when the use of ARNi is not feasible.6–13 |
| **1** | **A** | 3. In patients with previous or current symptoms of chronic HFrEF who are intolerant to ACEi  because of cough or angioedema and when the use of ARNi is not feasible, the use of ARB is rec- ommended to reduce morbidity and mortality.14–18 |
| **Value Statement: High Value (A)** | | 4. In patients with previous or current symptoms of chronic HFrEF, in whom ARNi is not feasible, treatment with an ACEi or ARB provides high economic value.19–25 |
| **1** | **B-R** | 5. In patients with chronic symptomatic HFrEF NYHA class II or III who tolerate an ACEi or ARB, replacement by an ARNi is recommended to further reduce morbidity and mortality.1–5 |
| **Value Statement: High Value (A)** | | 6. In patients with chronic symptomatic HFrEF, treatment with an ARNi instead of an ACEi pro- vides high economic value.26–29 |
| **3: Harm** | **B-R** | 7. ARNi should not be administered concomi- tantly with ACEi or within 36 hours of the last dose of an ACEi.30,31 |
| **3: Harm** | **C-LD** | 8. ARNi should not be administered to patients with any history of angioedema.32–35 |
| **3: Harm** | **C-LD** | 9. ACEi should not be administered to patients with any history of angioedema.36–39 |

\*See Section 7.2, “Diuretics and Decongestion Strategies in Patients with HF,” for diuretic recommendations.

## Synopsis

Inhibition of the renin-angiotensin system is recommend- ed to reduce morbidity and mortality for patients with HFrEF, and ARNi, ACEi, or ARB are recommended as first-line therapy.1–18 If patients have chronic symptomatic

HFrEF with NYHA class II or III symptoms and they toler- ate an ACEi or ARB, they should be switched to an ARNi because of improvement in morbidity and mortality.1–5 An ARNi is recommended as de novo treatment in hospital- ized patients with acute HF before discharge given im- provement in health status, reduction in the prognostic biomarker NT-proBNP, and improvement of LV remodel- ing parameters compared with ACEi/ARB. Although data are limited, the use of an ARNi may be efficacious as de novo treatment in patients with symptomatic chronic HFrEF to simplify management. ARB may be used as an alternative to ACEi in the setting of intolerable cough, or as alternatives to ACEi and ARNi in patients with a history of angioedema. If patients are switched from an ACEi to an ARNi or vice versa, there should be at least 36 hours between ACEi and ARNi doses.

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## Recommendation-Specific Supportive Text

* + - 1. An ARNi is composed of an ARB and an inhibi- tor of neprilysin, an enzyme that degrades natri- uretic peptides, bradykinin, adrenomedullin, and other vasoactive peptides. In PARADIGM-HF (Prospective Comparison of ARNi with ACEi to Determine Impact on Global Mortality and Morbidity in Heart Failure), an RCT that compared the first approved ARNi, sacubitril-valsartan, with enalapril in symptomatic patients with HFrEF tol- erating an adequate dose of either ACEi or ARB, sacubitril-valsartan significantly reduced the com- posite endpoint of cardiovascular death or HF hospitalization by 20% relative to enalapril.1 The benefit was observed to a similar extent for death and HF hospitalization and was consistent across prespecified subgroups.1 Use of an ARNi is more frequently associated with symptomatic hypoten- sion and a comparable incidence of angioedema when compared with enalapril.1 Sacubitril-valsartan has been approved for patients with symptomatic HF. HF effects and potential off-target effects may be complex with inhibition of the neprily- sin enzyme, which has multiple biological targets. Trial data have included ACEi/ARB-naïve patients before ARNi initiation (53% in the PIONEER-HF [Comparison of Sacubitril-Valsartan versus Enalapril on Effect on NT-proBNP in Patients Stabilized from an Acute Heart Failure Episode] trial and 24% in the TRANSITION [Comparison of Pre- and Post-discharge Initiation of Sacubitril/ Valsartan Therapy in HFrEF Patients After an Acute Decompensation Event] trial) and have shown similar efficacy and safety in treatment- naïve patients.2,3 The PIONEER-HF trial showed that ARNi reduced NT-proBNP levels in patients hospitalized for acute decompensated HF without increased rates of adverse events (worsening renal

function, hyperkalemia, symptomatic hypotension, angioedema) when compared with enalapril.3 Additional outcome analyses suggested reduction in all-cause mortality and rehospitalization for HF but were only hypothesis-generating as exploratory study endpoints. In the open-label TRANSITION trial, patients with HFrEF hospitalized with wors- ening HF were randomized to start ARNi either before or after discharge.2 Safety outcomes were similar for both arms, suggesting that early initia- tion may simplify management (rather than initiat- ing and uptitrating ACEi first and then switching to ARNi).2 ARNi should be initiated de novo in patients hospitalized with acute HFrEF before discharge in the absence of contraindications. ARNi may be ini- tiated de novo in patients with chronic symptom- atic HFrEF to simplify management, although data are limited. The PARADISE-MI (Prospective ARNi vs ACE Inhibitor Trial to DetermIne Superiority in Reducing Heart Failure Events After MI) trial40 will provide information on whether sacubitril-valsartan will significantly reduce the rate of cardiovascular death, HF hospitalization or outpatient HF requir- ing treatment in patients after acute MI, with LVEF

40% and/or pulmonary congestion, and 1 of 8 additional risk-enhancing factors like AF, previous MI, diabetes, compared with the ACEi ramipril; and whether the safety and tolerability of sacubitril-val- sartan was comparable to that of ramipril. Thus, at the present time, the efficacy of ARNi in patients with LV dysfunction, and HF in the early post-MI period, remains uncertain.

* + - 1. ACEi reduce morbidity and mortality in HFrEF. RCTs clearly establish the benefits of ACE inhibition in patients with mild, moderate, or severe symptoms of HF and in patients with or without CAD.6–11 Data suggest that there are no differences among avail- able ACEi in their effects on symptoms or survival.12 ACEi should be started at low doses and titrated upward to doses shown to reduce the risk of car- diovascular events in clinical trials. ACEi can pro- duce angioedema and should be given with caution to patients with low systemic blood pressures, renal insufficiency, or elevated serum potassium (>5.0 mEq/L). If maximal doses are not tolerated, inter- mediate doses should be tried; abrupt withdrawal of ACE inhibition can lead to clinical deterioration and should be avoided. Although the use of an ARNi in lieu of an ACEi for HFrEF has been found to be superior, for those patients for whom ARNi is inap- propriate, continued use of an ACEi for all classes of HFrEF remains strongly advised.
      2. ARB have been shown to reduce mortality and HF hospitalizations in patients with HFrEF in large RCTs.14–16 Long-term treatment with ARB in patients with HFrEF produces hemodynamic,

neurohormonal, and clinical effects consistent with those expected after interference with the renin-angiotensin system.17,18 Unlike ACEi, ARB do not inhibit kininase and are associated with a much lower incidence of cough and angioedema, although kininase inhibition by ACEi may produce beneficial vasodilatory effects. Patients who are intolerant to ACEi because of cough or angio- edema should be started on an ARB. ARB 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. ARB should be given with caution to patients with low systemic blood pressure, renal insufficiency, or elevated serum potassium (>5.0 mEq/L). Although ARB are alternatives for patients with ACEi-induced angio- edema, caution is advised because some patients have also developed angioedema with ARB. For those patients for whom an ACEi or ARNi is inap- propriate, use of an ARB remains advised.

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* + - 1. Several cost-effectiveness analyses consistently found that ACEi therapy provides high value for patients with chronic HF. A model-based analysis, using generic ACEi costs, found ACEi therapy was high value.19 Previous analyses also found ACEi therapy was high value despite previously higher ACEi costs.19,21,22,24,25 This includes a trial-based analysis of SOLVD (Studies of Left Ventricular Dysfunction) that modeled long-term outcomes.21 Previous analyses included a range of clinical sce- narios including asymptomatic LV dysfunction24 and LV dysfunction after MI,25 with ACEi therapy providing high value in each. There are limited data on the cost-effectiveness of ARBs from 2 clinical trials—a within-trial analysis of Val-HeFT (Valsartan Heart Failure Trial)23 and an analysis of the ELITE (Evaluation of Losartan in the Elderly) study20— which both suggested ARB therapy is high value. The high value of ARB therapy is also supported by its similar efficacy as ACEi therapy and the low-cost generic availability for both medication classes.
      2. Patients with chronic stable HFrEF who toler- ate ACEi and ARB should be switched to ARNi. In patients with mild-to-moderate HF who were able to tolerate both a target dose of enalapril (10 mg twice daily) and then subsequently an ARNi (sacubitril-valsartan; 200 mg twice daily, with the ARB component equivalent to valsartan 160 mg), hospitalizations and mortality were significantly decreased with the valsartan-sacubitril compound compared with enalapril.1 Another RCT and meta- analysis showed improvement in LV remodeling parameters with ARNi compared with enalapril.4,5
      3. Multiple model-based analyses evaluated the eco- nomic value of ARNi therapy compared with ACEi

therapy using the results of PARADIGM-HF.26–29,41 Three high-quality analyses26,28,29 consistently found costs per QALY <$60000, which provides high value according to the benchmarks adopted for the current clinical practice guideline. These results were robust to the range of sacubitril-val- sartan costs currently seen in care. These results were sensitive to the estimated mortality reduc- tion and duration of treatment effectiveness. ARNi would need to maintain effectiveness beyond the PARADIGM-HF study period (mean, 27 months) to be considered high value.29 If clinical benefit were limited to 27 months, ARNi would be inter- mediate value. One additional analysis, based on the PIONEER-HF trial, found that inpatient initia- tion of ARNi was also high value compared with delayed initiation postdischarge.27

* + - 1. Oral neprilysin inhibitors, used in combination with ACEi, can lead to angioedema, and concomitant use is contraindicated and should be avoided. A medication that represented a neprilysin inhibitor and an ACEi—omapatrilat—was studied in hyper- tension and HF, but its development was termi- nated because of an unacceptable incidence of angioedema.30,31 and associated significant mor- bidity. This adverse effect was thought to occur because ACEi and neprilysin break down brady- kinin, which can directly or indirectly cause angio- edema31,32 An ARNi should not be administered within 36 hours of switching from or to an ACEi.
      2. Omapatrilat, a neprilysin inhibitor (as well as an ACEi and aminopeptidase P inhibitor), was asso- ciated with a higher frequency of angioedema than that seen with enalapril in an RCT of patients with HFrEF.30 In a very large RCT of hyperten- sive patients, omapatrilat was associated with a 3-fold increased risk of angioedema compared with enalapril.31 Black patients and patients who smoked were particularly at risk. The high inci- dence of angioedema ultimately led to cessation of the clinical development of omapatrilat.33,34 Because of these observations, angioedema was an exclusion criterion in the first large trial assess- ing ARNi therapy in patients with hypertension35 and then in the large trial that showed clinical ben- efit of ARNi therapy in HFrEF.1 The rates of angio- edema were numerically higher in patients treated with ARNi than in patients treated with ACEi in PARADIGM-HF, although this difference did not reach significance.1 ARNi therapy should not be administered in patients with a history of angio- edema because of the concern that it will increase the risk of a recurrence of angioedema.
      3. Angioedema attributable to ACEi is thought to result from defective degradation of the vasoactive peptides bradykinin, des-Arg9-BK (a metabolite

of bradykinin), and substance P.36,37 ACEi should not be administered to patients with any history of angioedema, but ARB do not interfere as directly with bradykinin metabolism and have been associ- ated with low rates of angioedema.38,39

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* + 1. ***Beta Blockers***

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| **Recommendation for Beta Blockers**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **1** | **A** | 1. In patients with HFrEF, with current or previ- ous symptoms, use of 1 of the 3 beta blockers proven to reduce mortality (eg, bisoprolol, carvedilol, sustained-release metoprolol succi- nate) is recommended to reduce mortality and hospitalizations.1–3 |
| **Value Statement: High Value (A)** | | 2. In patients with HFrEF, with current or previous symptoms, beta-blocker therapy provides high economic value.4–8 |

## Synopsis

Treatment with beta blockers reduces the risk of death and the combined risk of death or hospitalization in pa- tients with HFrEF.1–3 In addition, this treatment can im- prove LVEF, lessen the symptoms of HF, and improve clinical status.1–3,9–11 Clinical trials have shown that beta blockers should be prescribed to all patients when HFrEF is diagnosed, including in-hospital, unless contra- indicated or not tolerated.1–3,9–11 These benefits of beta blockers were observed in patients with or without CAD, and in patients with or without diabetes, older patients, as well as in women and across racial and ethnic groups but not in patients with AF.1–3,10–12 Even if symptoms do not improve, long-term treatment should be maintained to reduce the risk of major cardiovascular events. Beta blockers should be initiated at low doses, and every ef- fort should be made to achieve the target doses of the beta blockers shown to be effective in major clinical tri- als, as tolerated1–3,9,10 (see Section 7.3.8, “GDMT Dosing, Sequencing and Uptitration”).

## Recommendation-Specific Supportive Text

* + - 1. Three beta blockers have been shown to be effec- tive in reducing the risk of death in patients with HFrEF: bisoprolol, sustained-release metoprolol (succinate), and carvedilol.1–3 The favorable find- ings with these 3 agents, however, should not be considered a beta-blocker class effect in HFrEF. Other beta blockers are not included in this rec- ommendation for use.13–15 Even when asymptom- atic, or when symptoms are mild or improve with other therapies, beta-blocker therapy is important and should not be delayed until symptoms return or disease progression is documented.16 Data show that beta blockers can be safely initiated before

hospital discharge, provided patients are clinically stabilized and do not require intravenous inotropic therapy for HF.17 If a contraindication or intoler- ance are noted, they should be documented, and the patient restarted on beta-blocker therapy in the future, so long as an absolute contraindication is not present. Even if symptoms or LVEF improve, long-term treatment with beta blockers and use of target doses should be maintained to reduce the risk of progression in LV dysfunction or major car- diovascular events.18,19 Abrupt withdrawal of beta- blocker therapy can lead to clinical deterioration and should be avoided unless indicated.18

* + - 1. Multiple analyses have shown the high value of beta-blocker therapy among HF patients. A model- based analysis, using generic beta-blocker costs, found beta-blocker therapy was high value.4 These results were consistent with earlier model-based cost-effectiveness analyses5–7 and a trial-based economic analysis of the US Carvedilol Heart Failure (CHF) Trials Program.8 Each of these stud- ies also found treatment with a beta blocker was high value despite using previously higher beta- blocker costs.
    1. ***Mineralocorticoid Receptor Antagonists (MRAs)***

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| **Recommendations for Mineralocorticoid Receptor Antagonists (MRAs) Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In patients with HFrEF and NYHA class II to IV symptoms, an MRA (spironolactone or eplere- none) is recommended to reduce morbidity and mortality, if eGFR is >30 mL/min/1.73 m2 and serum potassium is <5.0 mEq/L. Careful moni- toring of potassium, renal function, and diuretic dosing should be performed at initiation and closely monitored thereafter to minimize risk of hyperkalemia and renal insufficiency.1–3 |
| **Value Statement: High Value (A)** | | 2. In patients with HFrEF and NYHA class II to IV symptoms, MRA therapy provides high eco- nomic value.4–7 |
| **3: Harm** | **B-NR** | 3. In patients taking MRA whose serum potas- sium cannot be maintained at <5.5 mEq/L, MRA should be discontinued to avoid life- threatening hyperkalemia.8,9 |

## Synopsis

MRA (also known as aldosterone antagonists or anti-min- eralocorticoids) show consistent improvements in all-cause mortality, HF hospitalizations, and SCD across a wide range of patients with HFrEF.1–3 Patients at risk for renal dysfunc- tion or hyperkalemia require close monitoring, and eGFR

30 mL/min/1.73 m2 or serum potassium 5.0 mEq/L are contraindications to MRA initiation.10,11 Because of the higher selectivity of eplerenone for the aldosterone receptor, adverse effects such as gynecomastia and vaginal bleeding

are observed less often in patients who take eplerenone than in those who take spironolactone.

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## Recommendation-Specific Supportive Text

* + - 1. Clinical trials taken on MRA together—RALES (Randomized Aldactone Evaluation Study)1 ran- domized highly symptomatic patients with LVEF

35%; EPHESUS (Eplerenone Post–Acute Myocardial Infarction Heart Failure Efficacy and Survival Study)2 randomized patients post-MI with LVEF 40%; and EMPHASIS-HF (Eplerenone in Mild Patients Hospitalization and Survival Study in Heart Failure)3 randomized patients with mild symptoms and LVEF 30%—suggest a benefit of MRA across the spectrum of HFrEF, inclusive of a wide range of etiologies and disease severities. Initiation in the ambulatory or hospital setting is appropriate.12 The starting dose of spironolactone and eplerenone is 25 mg orally daily, increased to 50 mg daily orally after a month; for eGFR 31 to 49 mL/min/1.73 m2, dosing should be reduced by half. Regular checks of serum potassium levels and renal function should be performed accord- ing to clinical status, approximately 1 week, then 4 weeks, then every 6 months after initiating or inten- sifying MRA, with more frequent testing for clinical instability. We elected to remove the 2013 recom- mendation “Aldosterone receptor antagonists are recommended to reduce morbidity and mortality following an acute MI in patients who have LVEF of 40% or less who develop symptoms of HF or who have a history of diabetes mellitus, unless contrain- dicated” because the new recommendation covers the spectrum of symptomatic patients with HF.

* + - 1. The economic value of MRA therapy was assessed by both RCTs (RALES5 and EPHESUS6,7) and a model-based analysis.4 The model-based analysis used generic MRA costs and found therapy was high value with a cost per QALY of under $1000.4 The earlier trial-based economic analyses of MRAs from RALES and EPHESUS also found MRA ther- apy was high value despite using previously higher MRA costs.5–7
      2. Spironolactone and eplerenone are partially excreted through the kidneys, raising concerns about safety when eGFR is 30 mL/min/1.73 m2.10,11 Spironolactone and eplerenone decrease renal potassium excretion, raising the risk of hyperkalemia, particularly when MRA is initiated at serum potassium 5.0 mEq/L and continued

5.5 mEq/L. The incidence of clinically significant hyperkalemia events was <1% in EPHESUS and EMPHASIS-HF, without a significant difference between eplerenone and placebo.2,3 however, in the closely monitored setting of a RCT with enrollment

of younger patients with fewer multiple chronic conditions than seen in the general HFrEF popula- tion, safety may be overstated. Observational data have raised concerns about less favorable out- comes of MRA use for HFrEF during usual care.8,9 Coadministration of MRA with ACEi or ARB mildly increases the risk of hyperkalemia. Hyperkalemia risk was lower with ARNi in patients with chronic HF in the PARADIGM-HF trial13 but not different in patients with HF who were decompensated in the PIONEER-HF trial14 when compared with ACEi. Diarrhea causing dehydration or loop diuretic ther- apy interruption, because of worsening renal func- tion or hyperkalemia, should be a consideration for temporarily holding the MRA. The development of worsening renal function or hyperkalemia is often a reflection of acute clinical change or progressive disease, prompting careful evaluation of the entire medical regimen and other causes of hyperkalemia, in addition to holding the MRA. The efficacy of the use of potassium binders (eg, patiromer, sodium zir- conium cyclosilicate) to improve outcomes by facili- tating continuation of MRA is uncertain15,16 and is addressed in Section 7.3.6, “Other Drug Treatment.”

* + 1. ***Sodium-Glucose Cotransporter 2 Inhibitors***

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| **Recommendation for SGLT2i**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **1** | **A** | 1. In patients with symptomatic chronic HFrEF, SGLT2i are recommended to reduce hospital- ization for HF and cardiovascular mortality, irre- spective of the presence of type 2 diabetes.1,2 |
| **Value Statement: Intermediate Value (A)** | | 2. In patients with symptomatic chronic HFrEF, SGLT2i therapy provides intermediate eco- nomic value.3,4 |

## Synopsis

Several RCTs in patients with type 2 diabetes and ei- ther established CVD or high risk for CVD have shown that SGLT2i prevent HF hospitalizations compared with placebo.5–7 The overall 31% reduction in HF hospitaliza- tions was noted irrespective of the presence or absence of preexisting HF, although only 10% to 14% of par- ticipants had HF at baseline. The benefit appears in- dependent of the glucose-lowering effects.8 Therefore, several trials were launched to examine the efficacy of SGLT2i on outcomes in patients with HF, irrespective of the presence of type 2 diabetes. The DAPA-HF (Dapa- gliflozin and Prevention of Adverse Outcomes in Heart Failure) trial and EMPEROR-Reduced (EMPagliflozin outcomE tRial in Patients With chrOnic heaRt Failure With Reduced Ejection Fraction) showed the benefit of SGLT2i (dapagliflozin and empagliflozin, respectively) versus placebo on outcomes (median follow-up, 16–18

months).1,2 Patients enrolled had symptomatic chronic HFrEF (LVEF 40%, NYHA class II to IV, and elevated natriuretic peptides) and were already on GDMT. Impor- tant exclusions were eGFR <20 (EMPEROR-Reduced) or <30 mL/min/1.73 m2 (DAPA-HF), type 1 diabetes, or lower SBP <95 to 100 mm Hg.

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* + - 1. In the DAPA-HF and EMPEROR-Reduced tri- als, SGLT2i compared with placebo reduced the composite of cardiovascular death or HF hospi- talization by approximately 25%.1,2,9 The benefit in reduction of HF hospitalization was greater (30%) in both trials.9 Risk of cardiovascular death was significantly lowered (18%) with dapagliflozin, as was risk of all-cause mortality (17%). Although no significant cardiovascular mortality benefit was observed with empagliflozin in a meta-analysis of DAPA-HF and EMPEROR-Reduced trials, SGLT2i therapy was associated with a reduction in all-cause mortality and cardiovascular death.9 The benefits in both trials were seen irrespective of baseline diabetes status. Furthermore, serious renal outcomes were less frequent, and the rate of decline in eGFR was slower in patients treated with SGLT2i.1,2,9 In the SOLOIST-WHF (Effect of Sotagliflozin on Cardiovascular Events in Patients With Type 2 Diabetes And Worsening Heart Failure) trial, patients with diabetes and HF hospitalization (79%: LVEF, <50%) were enrolled before dis- charge or within 3 days of discharge. Sotagliflozin, a dual inhibitor of sodium-glucose co-transporters 1 and 2, reduced the combined endpoint of car- diovascular death, HF hospitalization, or urgent HF visits by 33%10 but has not been approved by the US Food and Drug Administration (FDA) as of 2021. Although SGLT2i increased risk for genital infections, they were otherwise well tolerated in the trials. As the use of SGLT2i is translated into clini- cal practice, caution is warranted for euglycemic ketoacidosis, genital and soft tissue infections, and adjustment of diuretics, if needed, to prevent vol- ume depletion.11
      2. Two model-based analyses evaluated the economic value of dapagliflozin therapy compared with usual care based on the results of the DAPA-HF trial.3,4 Both analyses found costs per QALY between

$60000 and $90000, which is consistent with intermediate value according to the benchmarks adopted for the current guideline. The results were most sensitive to the magnitude of cardiovascular mortality reduction, with a 8% reduction in cardio- vascular mortality necessary for a cost per QALY below $150000 in 1 study.3 There are a wide range of costs currently seen with dapagliflozin.

These 2 analyses estimated a cost per QALY below

$50 000 with annual dapagliflozin costs of $3240 (43% reduction from main analysis) and $2500 (40% reduction from main analysis), respectively.3,4 A smaller reduction in drug cost would lead to a cost per QALY of under $60 000, the threshold for high value in this guideline.

* + 1. ***Hydralazine and Isosorbide Dinitrate***

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| **Recommendations for Hydralazine and Isosorbide Dinitrate Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. For patients self-identified as African American with NYHA class III-IV HFrEF who are receiv- ing optimal medical therapy, the combination of hydralazine and isosorbide dinitrate is rec- ommended to improve symptoms and reduce morbidity and mortality.1,2 |
| **Value Statement: High Value (B-NR)** | | 2. For patients self-identified as African Ameri- can with NYHA class III to IV HFrEF who are receiving optimal medical therapy with ACEi or ARB, beta blockers, and MRA, the combination of hydralazine and isosorbide dinitrate provides high economic value.3 |
| **2b** | **C-LD** | 3. In patients with current or previous symp- tomatic HFrEF who cannot be given first-line agents, such as ARNi, ACEi, or ARB, because of drug intolerance or renal insufficiency, a combination of hydralazine and isosorbide dini- trate might be considered to reduce morbidity and mortality.4,5 |

## Synopsis

Two RCTs, V-HeFT I (Vasodilator Heart Failure Trial) and A-HeFT (African-American Heart Failure Trial), es- tablished benefit of the combination of hydralazine-iso- sorbide dinitrate in self-identified African Americans.2,4 A-HeFT was terminated early because of evidence of remarkable benefit, but the result is vulnerable to a small number of events and the exigencies of early cessation of RCTs.2 The benefit in both trials was seen only at doses achieved in those trials that are higher than doses typi- cally used in clinical practice and with short-acting nitrate therapy.2,4 Uptake of this regimen has been modest as a result of the complexity of the medical regimen and the array of drug-related adverse effects.5 Even when pre- scribed, there is marked underusage based on very low prescription refill rates. Race-based medical therapy re- mains a challenging issue, as well, with ongoing research now focused on biological hypotheses, particularly ab- sence of European ancestry, which may be associated with responsiveness to this combination. There are insuf- ficient data to guide the use of hydralazine-isosorbide dinitrate with ARNi. In patients with HFrEF who cannot receive first-line agents such as ARNi, ACEi, or ARB, re- ferral to a HF specialist can provide guidance for further management because the use of hydralazine and isosor- bide dinitrate in these patients is uncertain.

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* + - 1. In a large-scale trial that compared the vasodila- tor combination with placebo, the use of hydrala- zine and isosorbide dinitrate reduced mortality in patients with HF treated with digoxin and diuretics but not an ACEi or beta blocker.4 However, in 2 other trials that compared the vasodilator combina- tion with an ACEi, the ACEi produced more favor- able effects on survival.6,7 A post hoc retrospective analysis of these vasodilator trials showed particu- lar efficacy of isosorbide dinitrate and hydralazine in the African American cohort.1 In a subsequent trial, which was limited to patients self-identified as African American, the addition of a fixed-dose combination of hydralazine and isosorbide dini- trate to standard therapy with an ACEi or ARB, a beta blocker, and MRA offered significant benefit.2 Thus, the combination of hydralazine and isosor- bide dinitrate is appropriate for African Americans with HFrEF who remain symptomatic despite con- comitant use of ACEi (or ARB), beta blockers, and MRA. There are insufficient data for concomitant use with ARNi.
      2. The economic value of hydralazine and isosorbide nitrate therapy was assessed by the A-HeFT trial.3 This analysis found hydralazine and isosorbide dinitrate increased survival and reduced health care costs over the 12.8-month trial. Extrapolating beyond the trial, the analysis found hydralazine and isosorbide dinitrate remained high value over a life- time with a cost per life-year <$60 000 despite conservative assumptions regarding the durabil- ity of therapy effectiveness and previously higher hydralazine and isosorbide dinitrate costs.
      3. It is unclear if a benefit of hydralazine-isosorbide dinitrate (suggested in a trial before the use of ACEi)4 exists for non–African Americans with HFrEF. Despite the lack of data with the vaso- dilator combination in patients who are intoler- ant of ACEi or ARB, especially those with renal insufficiency, the combined use of hydralazine and isosorbide dinitrate might be considered as a therapeutic option in such patients. Although the potential benefit is unknown and has not been shown in recent observational datasets,5 in V-HeFT I, the use of hydralazine and isosorbide dinitrate reduced mortality in patients with HF treated with digoxin and diuretics, compared with placebo.4 If patients are unable to tolerate first- line agents, such as ARNi, ACEi, or ARB, because of drug intolerance, hypotension, or renal insuf- ficiency, referral to a HF specialist can provide guidance for further management, and the use of hydralazine and isosorbide dinitrate in these patients might be considered.
    1. ***Other Drug Treatment***

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| **Recommendations for Other Drug Treatment**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **2b** | **B-R** | 1. In patients with HF class II to IV symptoms, omega-3 polyunsaturated fatty acid (PUFA) supplementation may be reasonable to use as adjunctive therapy to reduce mortality and car- diovascular hospitalizations.1–4 |
| **2b** | **B-R** | 2. In patients with HF who experience hyperkale- mia (serum potassium level 5.5 mEq/L) while taking a renin-angiotensin-aldosterone system inhibitor (RAASi), the effectiveness of potas- sium binders (patiromer, sodium zirconium cyclosilicate) to improve outcomes by facilitat- ing continuation of RAASi therapy is uncer- tain.5,6 |
| **3: No**  **Benefit** | **B-R** | 3. In patients with chronic HFrEF without a spe- cific indication (eg, venous thromboembolism [VTE], AF, a previous thromboembolic event, or a cardioembolic source), anticoagulation is not recommended.7–9 |

## Synopsis

Trials in prevention of CVD, including HF, showed that omega-3 PUFA supplementation results in a 10% to 20% risk reduction in fatal and nonfatal cardiovascular events when used with other evidence-based thera- pies.2,3,10 Hyperkalemia is common in HF and can lead to arrhythmias and underuse of GDMT.11,12 Two newer gastrointestinal potassium-binding agents—patiromer and sodium zirconium cyclosilicate—have been shown to lower potassium levels and enable treatment with a RAASi in patients with HF.5,6,13

## Recommendation-Specific Supportive Text

* + - 1. Supplementation with omega-3 PUFA has been evaluated as an adjunctive therapy for CVD and HF.14 The GISSI-HF (Effect of n-3 polyunsaturated fatty acids in patients with chronic heart failure) trial showed a reduction in death among post-MI patients taking 1 g of omega-3 PUFA (850–882 mg of eicosapentaenoic acid [EPA] and docosa- hexaenoic acid [DHA] as ethyl esters in the ratio of 1:1.2).10 A post hoc subgroup analysis revealed that this reduction in mortality and SCD was con- centrated in the approximately 2000 patients with reduced LVEF.10 The GISSI-HF investigators ran- domized symptomatic patients with HF to 1 g daily of omega-3 PUFA (850–882 mg of EPA-DHA) or placebo. Death from any cause was reduced from 29% with placebo to 27% in those treated with omega-3 PUFA.2 The outcome of death or admission to hospital for a cardiovascular event was also significantly reduced. The REDUCE-IT trial randomized patients with established CVD

or diabetes with risk factors to 2 g of icosapent ethyl (a highly purified EPA) twice daily or placebo and showed a reduced risk for the composite of cardiovascular death, nonfatal MI, nonfatal stroke, coronary revascularization, or unstable angina.3 In reported studies, omega-3 PUFA therapy has been well tolerated. Recent studies have reported that in patients with cardiovascular risk treated with omega-3 fatty acid, there may be a dose-related risk of AF.3,15,16

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* + - 1. Hyperkalemia is common in HF as a result of the syndrome itself, comorbidities (diabetes, CKD), and use of RAASi, and can increase the risk for ventricular arrhythmias and mortality.11 Hyperkalemia results in dose reductions or dis- continuation of RAASi, compromising their cardio- renal benefit in HF.12 Two newer gastrointestinal potassium binders—patiromer (RLY5016) and sodium zirconium cyclosilicate (SZC)—remove potassium by exchanging cations (calcium for pat- iromer, and sodium and hydrogen for SZC), lead- ing to increased fecal excretion. Both agents have been FDA approved for treatment of hyperkalemia for patients receiving RAASi. In the PEARL-HF (Evaluation of the efficacy and safety of RLY5016, a polymeric potassium binder in patients with chronic heart failure) trial, patiromer led to lower potassium levels, less hyperkalemia, and a higher proportion of patients able to increase spironolac- tone dose to 50 mg daily compared with placebo.5 The HARMONIZE (Hyperkalemia Randomized Intervention Multidose ZS-9 Maintenance) trial included 94 patients (out of 258 total) with HF (87 of whom entered the double-blind phase).6,13 The SZC groups achieved lower potassium lev- els overall compared with placebo, and a higher proportion maintained normokalemia (potassium levels, <5.1 mEq/L). Whether patiromer or SZC improve clinical outcomes is under investigation. Adverse effects for the newer potassium bind- ers include hypomagnesemia (for patiromer) and edema (for SZC).
      2. In several retrospective analyses, the risk of throm- boembolic events was not lower in patients with HF taking warfarin than in patients not treated with antithrombotic drugs.17–19 The use of warfarin was associated with a reduction in major cardiovascular events and death in patients with HF in some stud- ies but not in others.20–22 An RCT that compared the outcome of patients with HFrEF assigned to aspirin, warfarin, or clopidogrel found that no therapy was superior.7 Another trial that compared aspirin with warfarin in patients with reduced LVEF, sinus rhythm, and no cardioembolic source showed no difference in either the primary outcome of death, stroke, or intracerebral hemorrhage, and no

difference in the combined outcome of death, isch- emic stroke, intracerebral hemorrhage, MI, or HF hospitalization.8 There was a significant increase in major bleeding with warfarin. A trial of rivaroxa- ban in patients with HFrEF, CAD, and normal sinus rhythm showed no difference in mortality, MI, and stroke compared with placebo.9 Therefore, there is no evidence of benefit for anticoagulation in HF patients without a specific indication (eg, VTE, AF, a previous thromboembolic event, or a cardioem- bolic source).

* + 1. ***Drugs of Unproven Value or That May Worsen HF***

|  |  |  |
| --- | --- | --- |
| **Recommendations for Drugs of Unproven Value or Drugs That May Worsen HF**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **3: No**  **Benefit** | **A** | 1. In patients with HFrEF, dihydropyridine calcium channel-blocking drugs are not recommended treatment for HF.1,2 |
| **3: No**  **Benefit** | **B-R** | 2. In patients with HFrEF, vitamins, nutritional supplements, and hormonal therapy are not recommended other than to correct specific deficiencies.3–9 |
| **3: Harm** | **A** | 3. In patients with HFrEF, nondihydropyridine calcium channel-blocking drugs are not recom- mended.10–13 |
| **3: Harm** | **A** | 4. In patients with HFrEF, class IC antiarrhythmic medications and dronedarone may increase the risk of mortality.14–16 |
| **3: Harm** | **A** | 5. In patients with HFrEF, thiazolidinediones increase the risk of worsening HF symptoms and hospitalizations.17–21 |
| **3: Harm** | **B-R** | 6. In patients with type 2 diabetes and high cardiovascular risk, the dipeptidyl pepti- dase-4 (DPP-4) inhibitors saxagliptin and alogliptin increase the risk of HF hospitaliza- tion and should be avoided in patients with HF.22–24 |
| **3: Harm** | **B-NR** | 7. In patients with HFrEF, NSAIDs worsen HF symptoms and should be avoided or withdrawn whenever possible.25–28 |

## Synopsis

Although there is strong evidence for benefit with se- lected medications for HFrEF as outlined in Section 7.3, “Pharmacological Treatment for HF With Reduced Ejec- tion Fraction (HFrEF),” there remain several classes of medications that have either unproven value or poten- tial for harm (Table 13). These recommendations are not exhaustive but focus on the most relevant and com- monly encountered medications in the management of patients with HFrEF: calcium channel blockers; antiar- rhythmic agents; NSAIDs; medications for treatment of type 2 diabetes including thiazolidinediones and DPP-4 inhibitors; and vitamins, hormones, and nutritional sup- plements.

##### Table 13. Selected Prescription Medications That May Cause or Exacerbate HF

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Drug or Therapeutic Class** | **Associated With HF** | | **Magnitude of HF Induction or Precipitation** | **LOE for HF Induction or Precipitation** | **Possible Mechanism(s)** | **Onset** |
| **Causes Direct Myocardial Toxicity** | **Exacerbates Underlying Myocardial Dysfunction** |
| COX, nonselective inhibi- tors (NSAIDs) |  | X | Major | B | Prostaglandin inhibition leading to sodium and water retention, increased systemic vascular resistance, and blunted response to diuretics | Immediate |
| COX, selective inhibitors (COX-2 inhibitors) |  | X | Major | B |
| Thiazolidinediones |  | X | Major | A | Possible calcium channel blockade | Intermediate |
| Saxagliptin |  | X | Major | A | Unknown | Intermediate to delayed |
| Alogliptin |  | X | Major | A |
| Flecainide |  | X | Major | A | Negative inotrope, proarrhythmic effects | Immediate to intermediate |
| Disopyramide |  | X | Major | B |  |  |
| Sotalol |  | X | Major | A | Proarrhythmic properties, beta blockade | Immediate to intermediate |
| Dronedarone |  | X | Major | A | Negative inotrope |  |
| **Alpha-1 blockers** | | | | | | |
| Doxazosin |  | X | Moderate | B | Beta-1-receptor stimulation with increas- es in renin and aldosterone | Intermediate to delayed |
| Diltiazem |  | X | Major | B | Negative inotrope | Immediate to intermediate |
| Verapamil |  | X | Major | B |
| Nifedipine |  | X | Moderate | C |

COX indicates cyclo-oxygenase; HF, heart failure; LOE, Level of Evidence; and NSAID, nonsteroidal anti-inflammatory drug. Adapted from Page RL 2nd et al.57 Copyright 2016 American Heart Association Inc.

## Recommendation-Specific Supportive Text

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1. Second-generation dihydropyridine calcium channel blockers, including amlodipine and felodipine, have greater selectivity for calcium channels in vascular smooth muscle cells and less myocardial depressant activity. By reduc- ing peripheral vasoconstriction and LV afterload, calcium channel blockers were thought to have a potential role in the management of chronic HF. The PRAISE-1 (Prospective Randomized Amlodipine Survival Evaluation-1) study showed a reduction in mortality in the subgroup of patients with nonischemic cardiomyopathy who received amlodipine.1 However, in the PRAISE-2 (Prospective Randomized Amlodipine Survival Evaluation 2) trial, which enrolled only patients with nonischemic cardiomyopathy, no survival benefit was observed, indicating the limitations of conclusions derived from subgroup analyses.29 However, dihydropyridine calcium channel block- ers may be used for treatment of hypertension in patients who have elevated blood pressure despite optimization of GDMT.
2. Many nutritional supplements and hormonal therapies have been proposed for the treatment of HF.3–9,30,31 Ultimately, most studies are limited by small sample sizes, surrogate endpoints, or

nonrandomized design.32,33 In addition, adverse effects and drug-nutraceutical interactions remain unresolved. There is a lack of evidence of benefit from vitamin D,3–5 thiamine,34–36 carnitine,37 and taurine38,39 and potential harm from vitamin E.6,7 The largest RCT of coenzyme Q10—Q-SYMBIO (Coenzyme Q10 as adjunctive treatment of chronic heart failure with focus on SYMptoms, BIomarker status [Brain-Natriuretic Peptide], and long-term Outcome [hospitalisations/mortality])—showed no changes in NYHA functional classification at 16 weeks, although the incidence of major adverse cardiovascular events at 2 years was significantly reduced (hazard ratio, 0.50; 95% CI, 0.32-0.80; *P*=0.003).8 Despite these findings, concerns about slow recruitment in this trial have tempered enthusi- asm for coenzyme Q10 supplementation in clinical practice.9,31 Hormonal therapies have been proposed for the treatment of HF, but trials have shown a neu- tral effect of testosterone,40,41 growth hormone,30,42 and thyroid hormone43–45 in HF outcomes.

1. Nondihydropyridine calcium channel blockers—dil-

tiazem and verapamil—are myocardial depressants and generally not well tolerated in HF. Verapamil had no impact of survival or major cardiac events post-MI, including in those patients with HFrEF after acute MI.10 In patients with nonischemic

cardiomyopathy, diltiazem had no impact on mor- tality13 but, in HFrEF after acute MI, diltiazem was associated with a higher risk of recurrent HF.11,12

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1. In the CAST (Cardiac Arrhythmia Suppression) trial, patients with asymptomatic ventricular arrhythmias post-MI on the class IC antiarrhyth- mics encainide or flecainide had increased mortal- ity.14 The applicability of CAST to patients without recent MI or to other class I antiarrhythmic drugs is uncertain, but class IC antiarrhythmic agents are generally avoided in patients with structural heart disease. In ANDROMEDA (Antiarrhythmic Trial with Dronedarone in Moderate to Severe CHF Evaluating Morbidity Decrease Study), for the class III antiarrhythmic dronedarone, patients with HFrEF who were hospitalized had increased mortality.16 In the SWORD (Survival With ORal D-sotalol) trial of the class III antiarrhythmic sotalol, patients with HF post-MI had increased mortality.15 However, SWORD was published in 1996, and whether sotalol would be harmful in the current era of GDMT and ICDs is uncertain; sotalol may be used for refractory atrial-ventricular arrhyth- mias with close monitoring for decompensation. Amiodarone46,47 and dofetilide48,49 are the only anti- arrhythmic agents with neutral effects on mortal- ity in clinical trials of patients with HFrEF. Class IA antiarrhythmic agents such as quinidine and class IB agents such as mexiletine have not been stud- ied and may be indicated for the management of refractory ventricular arrhythmias in the context of the individual patient’s risk benefit calculus and in conjunction with electrophysiology consultation.
2. Thiazolidinediones increase insulin sensitivity by activating nuclear peroxisome proliferator-acti- vated receptor gamma (PPAR-γ). Expressed in virtually all tissues, PPAR-γ also regulates sodium reabsorption in the collecting ducts of the kidney. In observational cohort studies,17 meta-analysis,18 and clinical trials,19–21 thiazolidinediones have been associated with increased incidence of fluid reten- tion and HF events in those patients with19,21or without18,20a previous history of HF.
3. DPP-4 is a cell-surface enzyme that deactivates several peptides include glucose-dependent insulinotropic polypeptide and glucagon-like peptide 1. DPP-4 inhibitors affect glucose reg- ulation through multiple mechanisms, includ- ing enhancement of glucose-dependent insulin secretion, slowed gastric emptying, and reduction of postprandial glucagon and of food intake. The impact of DPP-4 inhibitors on cardiovascular out- comes in patients with diabetes and high cardio- vascular risk has been assessed in multiple RCTs. Saxagliptin increased the risk of hospitalization for HF,22 as did alogliptin in a post hoc analysis

including only patients with no HF history,23,50 but sitagliptin51,52and linagliptin53–55 did not; these findings may have been a result of baseline dif- ferences in the use of metformin, thiazolidinedio- nes, and insulin, which also affect HF risk. The FDA recommends discontinuation specifically of saxagliptin and alogliptin in patients who develop HF,56 and whether the risk of worsening HF is a class effect of DPP-4 inhibitors is unclear.

1. NSAIDs inhibit the synthesis of renal prosta- glandins, which mediate vasodilation in the kid- neys and directly inhibit sodium resorption in the thick ascending loop of Henle and collecting tubule. Hence, NSAIDs can cause sodium and water retention and blunt the effects of diuret- ics. Several observational cohort studies have revealed increased morbidity and mortality in patients with HF using either nonselective or selective NSAIDs.25–28
   * 1. ***GDMT Dosing: Sequencing and Uptitration***

|  |  |  |
| --- | --- | --- |
| **Recommendations for GDMT Dosing: Sequencing and Uptitration Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In patients with HFrEF, titration of guideline- directed medication dosing to achieve target doses showed to be efficacious in RCTs is rec- ommended, to reduce cardiovascular mortality and HF hospitalizations, unless not well toler- ated.1–10 |
| **2a** | **C-EO** | 2. In patients with HFrEF, titration and optimiza- tion of guideline-directed medications as frequently as every 1 to 2 weeks depending on the patient’s symptoms, vital signs, and labora- tory findings can be useful to optimize manage- ment. |

## Synopsis

Clinical trials of ACEi, ARB, ARNi, beta blockers, and most other HFrEF medications had therapy initiated at low dose by trial protocol.1–9,11–14 If the initial dose was tolerated, the protocol would then direct the uptitration of medication dose over time to a specified target dose (Table 14), unless not well tolerated. Even if symptoms improved or other indicators of response were shown at lower doses, the medication dose would still be in- creased to the trial-defined target doses. Because these target doses were the ones that established the efficacy and safety of these medications in HFrEF and serve as the basis of the guideline recommendations (Table 15), use of these target doses is recommended, if tolerated.1–9,11–14 Use of all 4 drug classes has been estimated to reduce all-cause mortality by 73% com- pared with no treatment.15

If the target dose cannot be achieved or is not well tol- erated, then the highest tolerated dose is recommended.

##### Table 14. Drugs Commonly Used for HFrEF (Stage C HF)

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Drug** | **Initial Daily Dose(s)** | **Target Doses(s)** | **Mean Doses Achieved in Clinical Trials** | **References** |
| ACEi | | | | |
| Captopril | 6.25 mg 3 times daily | 50 mg 3 times daily | 122.7 mg total daily | 19 |
| Enalapril | 2.5 mg twice daily | 10–20 mg twice daily | 16.6 mg total daily | 3 |
| Fosinopril | 5–10 mg once daily | 40 mg once daily | NA | … |
| Lisinopril | 2.5–5 mg once daily | 20–40 mg once daily | 32.5–35.0 mg total daily | 17 |
| Perindopril | 2 mg once daily | 8–16 mg once daily | NA | … |
| Quinapril | 5 mg twice daily | 20 mg twice daily | NA | … |
| Ramipril | 1.25–2.5 mg once daily | 10 mg once daily | NA | … |
| Trandolapril | 1 mg once daily | 4 mg once daily | NA | … |
| ARB | | | | |
| Candesartan | 4–8 mg once daily | 32 mg once daily | 24 mg total daily | 20 |
| Losartan | 25–50 mg once daily | 50–150 mg once daily | 129 mg total daily | 18 |
| Valsartan | 20–40 mg once daily | 160 mg twice daily | 254 mg total daily | 21 |
| ARNi | | | | |
| Sacubitril-valsartan | 49 mg sacubitril and 51 mg val- sartan twice daily (therapy may be initiated at 24 mg sacubitril and 26 mg valsartan twice daily) | 97 mg sacubitril and 103 mg val- sartan twice daily | 182 mg sacubitril and 193 mg valsartan total daily | 22 |
| Beta blockers | | | | |
| Bisoprolol | 1.25 mg once daily | 10 mg once daily | 8.6 mg total daily | 1 |
| Carvedilol | 3.125 mg twice daily | 25–50 mg twice daily | 37 mg total daily | 23 |
| Carvedilol CR | 10 mg once daily | 80 mg once daily | NA | … |
| Metoprolol succinate extended release (metoprolol CR/XL) | 12.5–25 mg once daily | 200 mg once daily | 159 mg total daily | 11 |
| Mineralocorticoid receptor antagonists | | | | |
| Spironolactone | 12.5–25 mg once daily | 25–50 mg once daily | 26 mg total daily | 6 |
| Eplerenone | 25 mg once daily | 50 mg once daily | 42.6 mg total daily | 13 |
| SGLT2i | | | | |
| Dapagliflozin | 10 mg once daily | 10 mg once daily | 9.8 mg total daily | 8 |
| Empagliflozin | 10 mg once daily | 10 mg once daily | NR | 9 |
| Isosorbide dinitrate and hydralazine | | | | |
| Fixed dose combination | 20 mg isosorbide dinitrate and  37.5 mg hydralazine 3 times daily | 40 mg isosorbide dinitrate and 75 mg hydralazine 3 times daily | 90 mg isosorbide dinitrate and  ∼175 mg hydralazine total daily | 10 |
| Isosorbide dinitrate and hydrala- zine | 20–30 mg isosorbide dinitrate and 25–50 mg hydralazine 3–4 times daily | 120 mg isosorbide dinitrate total daily in divided doses and 300 mg hydralazine total daily in di- vided doses | NA | 24 |
| If Channel inhibitor | | | | |
| Ivabradine | 5 mg twice daily | 7.5 mg twice daily | 12.8 total daily | 25–27 |
| Soluble guanylate cyclase stimulator | | | | |
| Vericiguat | 2.5 mg once daily | 10 mg once daily | 9.2 mg total daily | 28 |
| Digoxin | 0.125–0.25 mg daily (modified according to monogram) | Individualized variable dose to achieve serum digoxin concentra- tion 0.5–<0.9 ng/mL | NA | 29,30 |

ACE indicates angiotensin-converting enzyme; ARB, angiotensin receptor blocker; CR, controlled release; CR/XL, controlled release/extended release; HF, heart failure; HFrEF, heart failure with reduced ejection fraction; NA, not applicable; NR, not reported; and SGLT2i, sodium glucose cotransporter 2 inhibitor.

There are no direct data showing that use of lower doses of HFrEF medications among patients, where higher target doses could be tolerated, would produce the same or simi-

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lar degree of clinical benefit. In trials that have evaluated dose response for outcomes, composite event rates were lower with target doses compared with lower dose.16–18

**Table 15. Benefits of Evidence-Based Therapies for Patients With HFrEF3–6,8,10–14,23,31–42**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Evidence-Based Therapy** | **Relative Risk Reduction in All-Cause Mortality in Pivotal RCTs, %** | **NNT to Prevent All-Cause Mortality Over Time\*** | **NNT for All-Cause Mortality (Standardized to 12 mo)** | **NNT for All- Cause Mortality (Standardized to 36 mo)** |
| ACEi or ARB | 17 | 22 over 42 mo | 77 | 26 |
| ARNi† | 16 | 36 over 27 mo | 80 | 27 |
| Beta blocker | 34 | 28 over 12 mo | 28 | 9 |
| Mineralocorticoid receptor antagonist | 30 | 9 over 24 mo | 18 | 6 |
| SGLT2i | 17 | 43 over 18 mo | 63 | 22 |
| Hydralazine or nitrate‡ | 43 | 25 over 10 mo | 21 | 7 |
| CRT | 36 | 12 over 24 mo | 24 | 8 |
| ICD | 23 | 14 over 60 mo | 70 | 23 |

ACEi indicates angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; ARNi, angiotensin receptor neprilysin inhibitor; CRT, cardiac resyn- chronization therapy; HFrEF, heart failure with reduced ejection fraction; ICD, implantable cardioverter-defibrillator; NNT, number needed to treat; RCT, randomized controlled trial; and SGLT2i, sodium-glucose cotransporter-2 inhibitor.

\*Median duration follow-up in the respective clinical trial.

†Benefit of ARNi therapy incremental to that achieved with ACEi therapy. For the other medications shown, the benefits are based on comparisons to placebo control.

‡Benefit of hydralazine-nitrate therapy was limited to African American patients in this trial.

## Recommendation-Specific Supportive Text

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1. The use of these specific medications for HFrEF should involve initiation at low-starting doses, uptitration at specified intervals as tolerated, and achieving-maintaining the target doses shown to be effective in major clinical trials. Every effort should be made by clinicians to achieve and main- tain the clinical trial–defined target doses (Table 13) of guideline-directed medications, as long as they are well tolerated by the patient. Patients should be monitored for changes in heart rate, blood pressure, electrolytes, renal function, and symptoms during this uptitration period. Planned uptitration of a HF medication should be delayed until any adverse effects observed with lower doses have resolved. When such a strategy is used for dose titration, most patients (approxi- mately 70%–85%) enrolled in clinical trials who received these medications were able to tolerate short-, intermediate-, and long-term treatment with these agents and achieve and maintain the trial defined target dose.1–9,11–14 Repeated attempts at uptitration can result in optimization, even if initial attempts may fail. In patients with HFrEF, beta blockers provide dose-dependent improvements in LVEF, reduction in HF hospi- talizations, and reduction in all-cause mortality.17 Trials of lower versus higher dose of ACEi and ARB have shown lower risk of cardiovascular death or HF hospitalization with higher doses, with similar safety and tolerability.17,18
2. Initiation and titration should be individualized and optimized without delay according to patient’s symptoms, vital signs, functional status, tolerance, renal function, electrolytes, comorbidities, specific

cause of HF, and ability of follow-up. In patients with HFrEF, simultaneous initiation or sequencing, and order of guideline-directed medications are usually individualized according to patient’s symp- toms, vital signs, functional status, tolerance, renal function, electrolytes, comorbidities, specific cause of HF, and ability of follow-up, and does not neces- sarily need to be done according to the sequence of trial publications and should not be delayed.

#### *Additional Medical Therapies*

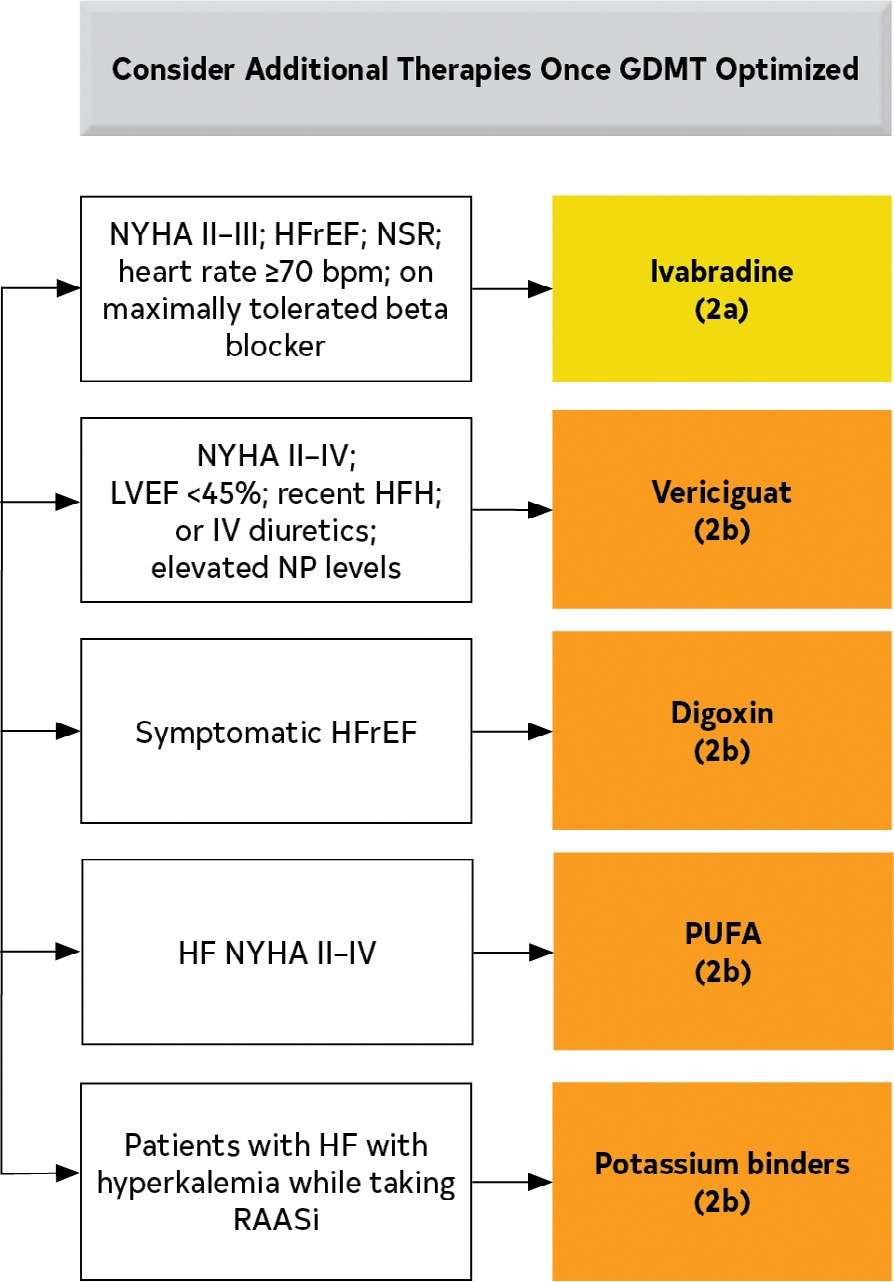
* + - 1. *Management of Stage C HF: Ivabradine*

|  |  |  |
| --- | --- | --- |
| **Recommendation for the Management of Stage C HF: Ivabradine Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **2a** | **B-R** | 1. For patients with symptomatic (NYHA class II to III) stable chronic HFrEF (LVEF 35%) who are receiving GDMT, including a beta blocker at maximum tolerated dose, and who are in sinus rhythm with a heart rate of 70 bpm at rest, ivabradine can be beneficial to reduce HF hospitalizations and cardiovascular death.1,2 |

## Synopsis

Heart rate is a strong predictor of cardiovascular out- comes in the general population and in patients with CVD, including HF. The SHIFT (Ivabradine and Outcomes in Chronic Heart Failure) trial tested the hypothesis that reducing heart rate in patients with HF improves cardio- vascular outcomes.1 SHIFT demonstrated the efficacy of ivabradine, a sinoatrial node modulator that selectively inhibits the If current, in reducing the composite endpoint of cardiovascular death or HF hospitalization in patients with HF. See Figure 7 for a summary of additional medi- cal therapy recommendations.

* + - 1. *Pharmacological Treatment for Stage C HFrEF: Digoxin*



|  |  |  |
| --- | --- | --- |
| **Recommendation for the Pharmacological Treatment for Stage C HFrEF: Digoxin**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **2b** | **B-R** | 1. In patients with symptomatic HFrEF despite GDMT (or who are unable to tolerate GDMT), digoxin might be considered to decrease hospi- talizations for HF.1,2 |

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**Figure 7. Additional Medical Therapies for Patients With HFrEF.** Colors correspond to COR in Table 2. Recommendations for additional medical therapies that may be considered for patients with HF are shown. GDMT indicates guideline-directed medical therapy; HF,

heart failure; HFH, heart failure hospitalization; HFrEF, heart failure with reduced ejection fraction; IV, intravenous; LVEF, left ventricular ejection fraction; LVESD, left ventricular end systolic dimension; MV, mitral valve; MR, mitral regurgitation; NP, natriuretic peptide; NSR, normal sinus rhythm; NYHA, New York Heart Association; and RAASi, renin-angiotensin-aldosterone system inhibitors.

## Recommendation-Specific Supportive Text

1. Although the primary outcome in SHIFT was a com- posite of hospitalization and cardiovascular death, the greatest benefit was a reduction in HF hospi- talization. SHIFT included patients with HFrEF and LVEF 35% who were in sinus rhythm with a resting heart rate of 70 bpm. Participants were predomi- nantly NYHA class II and III. Participants had been hospitalized for HF in the preceding 12 months and were on stable GDMT for 4 weeks before initiation of ivabradine therapy.1–4 The target of ivabradine is heart rate, and the benefit of ivabradine results from a reduction in heart rate. However, only 25% of patients studied in SHIFT were on optimal doses of beta-blocker therapy. Given the well-proven mor- tality benefits of beta-blocker therapy, these agents should be initiated and uptitrated to target doses, as tolerated, before assessing the resting heart rate for consideration of ivabradine initiation.5,6

## Synopsis

To date, there has been only 1 large-scale, RCT of digox- in in patients with HF.1 This trial, which predated current GDMT, primarily enrolled patients with NYHA class II to III HF and showed that treatment with digoxin for 2 to 5 years had no effect on mortality but modestly reduced the combined risk of death and hospitalization. The trial also found no significant effect on health-related QOL in a subset of the trial patients.3 The effect of digoxin on hospitalizations has been supported by retrospective analyses and meta-analyses.2,4–6 Additionally, observa- tional studies and retrospective analyses have shown improvement in symptoms and exercise tolerance in mild to moderate HF; however, they have mostly shown either lack of mortality benefit or increased mortality associated with digoxin.7 The benefit in patients on current GDMT is unclear because most trials preceded current GDMT. Thus, use of digoxin requires caution in patients with HF and is reserved for those who remain symptomatic de- spite optimization of GDMT.

## Recommendation-Specific Supportive Text

1. Digoxin is usually initiated at a low dose because higher doses are rarely required in the management of HF and are potentially detrimental. Two retro- spective analyses of large-scale clinical trials have shown a linear relationship between mortality and digoxin serum concentration in patients with AF and at risk for stroke, including those with HF, and in patients with HF. The risk of death was indepen- dently associated with serum digoxin concentration, with a significantly higher risk observed in those with concentrations 1.2 ng/mL and 1.6 ng/mL.8,9 The benefit of digoxin in patients with HF remains con- troversial. GDMT is expected to be optimized before considering the addition of digoxin. Clinical worsen- ing after withdrawal of digoxin has been shown.10 Therapy with digoxin may either be continued in the absence of a contraindication or discontinued with caution.11 Therapy with digoxin is commonly initi- ated and maintained at a dose of 0.125 to 0.25 mg daily. Low doses (0.125 mg daily or every other day) should be used initially if the patient is >70 years of

age, has impaired renal function, or has a low lean body mass. Higher doses (eg, digoxin 0.375 to 0.50 mg daily) are rarely used or needed in the manage- ment of patients with HF.

* + - 1. *Pharmacological Treatment for Stage C HFrEF: Soluble Guanylyl Cyclase Stimulators*

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| **Recommendation for Pharmacological Treatment for Stage C HFrEF: Soluble Guanylyl Cyclase Stimulators**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **2b** | **B-R** | 1. In selected high-risk patients with HFrEF and recent worsening of HF already on GDMT, an oral soluble guanylate cyclase stimulator (vericiguat) may be considered to reduce HF hospitalization and cardiovascular death.1 |

## Synopsis

In patients with progression of HFrEF despite GDMT, there may be a role for novel therapeutic agents. Oral soluble guanylyl cyclase stimulator (eg, vericiguat) di- rectly binds and stimulates sGC and increases cGMP production. cGMP has several potentially beneficial effects in patients with HF, including vasodilation, im- provement in endothelial function, as well as decrease in fibrosis and remodeling of the heart.2–7 The VICTORIA (Vericiguat Global Study in Subjects with Heart Failure with Reduced Ejection Fraction) trial randomized 5050 higher-risk patients with worsening HFrEF to vericiguat versus placebo.1

## Recommendation-Specific Supportive Text

* + - * 1. Patients with HFrEF in the VICTORIA trial had LVEF

<45%, NYHA class II to IV, were on GDMT, with elevated natriuretic peptides (BNP 300 pg/mL or NT-proBNP 1000 pg/mL if in sinus rhythm; higher cutoffs with AF), and recent HF worsening (hospitalized within 6 months or recently received intravenous diuretic therapy without hospitaliza- tion). Patients on long-acting nitrates, with SBP

<100 mm Hg, or eGFR <15 mL/min/1.73 m2 were

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| **Recommendations for ICDs and CRTs**  **Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In patients with nonischemic DCM or ischemic heart disease at least 40 days post-MI with LVEF 35% and NYHA class II or III symp- toms on chronic GDMT, who have reasonable expectation of meaningful survival for >1 year, ICD therapy is recommended for primary pre- vention of SCD to reduce total mortality.1–9 |
| **Value Statement: High Value (A)** | | 2. A transvenous ICD provides high economic value in the primary prevention of SCD particularly when the patient’s risk of death caused by ventricular arrythmia is deemed high and the risk of nonarrhythmic death (either cardiac or noncardiac) is deemed low based on the patient’s burden of comorbidi- ties and functional status.10–15 |
| **1** | **B-R** | 3. In patients at least 40 days post-MI with LVEF  30% and NYHA class I symptoms while receiving GDMT, who have reasonable expec- tation of meaningful survival for >1 year, ICD therapy is recommended for primary prevention of SCD to reduce total mortality.6 |
| **1** | **B-R** | 4. For patients who have LVEF 35%, sinus rhythm, left bundle branch block (LBBB) with a QRS duration 150 ms, and NYHA class II, III, or ambulatory IV symptoms on GDMT, CRT is indicated to reduce total mortality, reduce hospi- talizations, and improve symptoms and QOL.16–21 |
| **Value Statement: High Value (B-NR)** | | 5. For patients who have LVEF 35%, sinus rhythm, LBBB with a QRS duration of 150 ms, and NYHA class II, III, or ambulatory IV symptoms on GDMT, CRT implantation pro- vides high economic value.22–27 |
| **2a** | **B-R** | 6. For patients who have LVEF 35%, sinus rhythm, a non-LBBB pattern with a QRS duration 150 ms, and NYHA class II, III, or ambulatory class IV symptoms on GDMT, CRT can be useful to reduce total mortality, reduce hospitalizations, and improve symp- toms and QOL.16–21,28–33 |
| **2a** | **B-R** | 7. In patients with high-degree or complete heart block and LVEF of 36% to 50%, CRT is reasonable to reduce total mortality, reduce hospitalizations, and improve symptoms and QOL.34,35 |

a higher risk population. Although not statistically significant, symptomatic hypotension (9.1% versus 7.9%; *P*=0.12) and syncope (4.0% versus 3.5%; *P*=0.30) were numerically higher in the vericiguat group versus placebo. There was heterogeneity by subgroup analysis, and patients in the highest quartile of NT-proBNP subgroup (NT proBNP level

>5314 pg/mL) did not have benefit from vericiguat when compared with placebo.

## Device and Interventional Therapies for HFrEF

#### *ICDs and CRTs*

excluded. Over a median follow-up of 10.8 months,

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the primary outcome, cardiovascular death or HF hospitalization, occurred in 35.5% with vericiguat compared with 38.5% with placebo (HR, 0.90; *P*=0.019). All-cause mortality occurred in 20.3% in the vericiguat group and 21.2% in the placebo group (HR, 0.95; 95% CI, 0.84-1.07; *P*=0.38) and

composite of any-cause death or HF hospitaliza- tion was also lower in the vericiguat group versus placebo group (HR, 0.90; 95% CI, 0.83–0.98; *P*=0.02). The relative risk reduction of 10% in the primary outcome was lower than expected, even in

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| **Recommendations for ICDs and CRTs (Continued)** | | |
| **COR** | **LOE** | **Recommendations** |
| **2a** | **B-NR** | 8. For patients who have LVEF 35%, sinus rhythm, LBBB with a QRS duration of 120 to 149 ms, and NYHA class II, III, or ambu- latory IV symptoms on GDMT, CRT can  be useful to reduce total mortality, reduce hospitalizations, and improve symptoms and QOL.16–21,28–33 |
| **2a** | **B-NR** | 9. In patients with AF and LVEF 35% on GDMT, CRT can be useful to reduce total mortality, improve symptoms and QOL, and increase LVEF, if: a) the patient requires ventricular pac- ing or otherwise meets CRT criteria and b) atrioventricular nodal ablation or pharmacologi- cal rate control will allow near 100% ventricu- lar pacing with CRT.16–21,28–33 |
| **2a** | **B-NR** | 10. For patients on GDMT who have LVEF  35% and are undergoing placement of  a new or replacement device implantation with anticipated requirement for significant (>40%) ventricular pacing, CRT can be useful to reduce total mortality, reduce hospitalizations, and improve symptoms and QOL.16–21,28–33 |
| **2a** | **B-NR** | 11. In patients with genetic arrhythmogenic car- diomyopathy with high-risk features of sudden death, with EF 45%, implantation of ICD is reasonable to decrease sudden death.36,37 |
| **2b** | **B-NR** | 12. For patients who have LVEF 35%, sinus rhythm, a non-LBBB pattern with QRS dura- tion of 120 to 149 ms, and NYHA class III or ambulatory class IV on GDMT, CRT may be considered to reduce total mortality, reduce hospitalizations, and improve symptoms and QOL.16–21,28–33 |
| **2b** | **B-NR** | 13. For patients who have LVEF 30%, ischemic cause of HF, sinus rhythm, LBBB with a QRS duration 150 ms, and NYHA class I symp- toms on GDMT, CRT may be considered to reduce hospitalizations and improve symptoms and QOL.16–21,28–33 |
| **3: No**  **Benefit** | **B-R** | 14. In patients with QRS duration <120 ms, CRT is not recommended.36–41 |
| **3: No**  **Benefit** | **B-NR** | 15. For patients with NYHA class I or II symptoms and non-LBBB pattern with QRS duration  <150 ms, CRT is not recommended.16–21,28–33 |
| **3: No**  **Benefit** | **C-LD** | 16. For patients whose comorbidities or frailty limit survival with good functional capacity to  <1 year, ICD and cardiac resynchronization therapy with defibrillation (CRT-D) are not indi- cated.1–9,16–21 |

## Synopsis

RCTs have informed the decisions regarding cardiac implantable devices (ICDs and CRTs) over the past 20 years. In fact, the seminal RCTs for ICDs and CRTs are unlikely to be repeated. Subgroup analyses of these tri- als have also informed decisions, but these were not the primary endpoints of these studies and thus should be interpreted with caution. GDMT is optimized before ICD and CRT implantation to assess whether the LVEF im- proves. Figures 8 and 9 summarize device and interven- tional therapy recommendations.

## Recommendation-Specific Supportive Text

* + - 1. ICDs were first assessed in patients who had been resuscitated from a cardiac arrest. In AVID (Antiarrhythmics versus Implantable Defibrillators trial), CASH (Cardiac Arrest Study Hamburg), and CIDS (Canadian Implantable Defibrillator StudyS), benefit was observed in those who were randomized to ICDs.1–3 Extension of benefit was then shown in other patient populations that were at perceived risk of SCD. In the first MADIT (Multicenter Automated Defibrillator Implantation Trial) trial, patients with previous MI, LVEF 35% with nonsustained VT had a mortality benefit with ICD.4 Similar populations in MUSTT (Multicenter UnSustained Tachycardia Trial) also showed bene- fit.5 In MADIT-II (Multicenter Automatic Defibrillator Implantation Trial II), patients with no arrhyth- mia qualifier but with previous MIs and LVEF

30% derived benefit from ICD.6 The DEFINITE

(Defibrillators in Non-Ischemic Cardiomyopathy Treatment Evaluation) study included only non- ischemic patients with LVEF 35% and frequent premature ventricular contractions (PVCs) or non- sustained ventricular tachycardia (VT).7 There was a trend to mortality benefit, but it ultimately did not achieve significance. In SCD-HEFT (Sudden Cardiac Death in Heart Failure Trial), patients with ischemic and nonischemic cardiomyopathy, LVEF

35%, and HF class II to III showed benefit with

an ICD compared with either amiodarone or pla- cebo.8 More recently, the DANISH (Defibrillator Implantation in Patients with Nonischemic Systolic Heart Failure) trial enrolled patients with nonisch- emic cardiomyopathy and LVEF 35% to ICD or standard care.9 There was no reduction in the pri- mary endpoint of total mortality, but there was a reduction in SCD risk. In the DANISH trial, 58% of patients in each limb received CRT, possibly miti- gating the benefit of an ICD.

* + - 1. Economic outcomes of ICD implantation for pri- mary prevention of SCD were assessed in 3 RCTs (MADIT-I,13 MADIT-II,15 and SCD-HeFT,12 1 obser- vational study,10 and 3 simulation models,11,14,42 all of which had generally consistent results. All stud- ies reported increased survival and life expectancy and higher lifetime costs of medical care with an ICD than without an ICD. The incremental cost- effectiveness ratios were generally <$60 000 per year of life added by an ICD, which provides high value according to the benchmarks adopted for the current guideline. The value provided by an ICD was consistently high when life expectancy was projected to increase by >1.4 years.14 In contrast, when survival was not increased by ICD implanta- tion, as in the coronary artery bypass graft (CABG) Patch trial,43 the ICD did not provide value, because

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##### Figure 8. Algorithm for CRT Indications in Patients With Cardiomyopathy or HFrEF.

Colors correspond to COR in Table 2. Recommendations for cardiac resynchronization therapy (CRT) are displayed. AF indicates atrial fibrillation; Amb, ambulatory; CM, cardiomyopathy; GDMT, guideline-directed medical therapy; HB, heart block; HF, heart failure; HFrEF, heart failure with reduced ejection fraction; LBBB, left bundle branch block; LV, left ventricular; LVEF, left ventricular ejection fraction; NSR, normal sinus rhythm; NYHA, New York Heart Association; and RV, right ventricular.

the higher costs were unaccompanied by a gain in life expectancy.14

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* + - 1. The MADIT-II trial randomized patients with previ- ous MI and LVEF <30%, without any limitation of HF class, to ICDs or not.6 Thirty-seven percent of the patients were in class I congestive heart failure (CHF). Mortality was reduced with an ICD.
      2. Most of the relevant data for the guidelines of CRT in HF come from seminal trials published from 2002 to 2010. The first of these was the MIRACLE (Multicenter InSync Randomized Clinical Evaluation) trial, which took patients with LVEF

35%, moderate to severe HF, and QRS duration

130 ms.16 There was a benefit in the 6-minute walk test, QOL, functional HF classification, and LVEF. The COMPANION (Comparison of Medical Therapy, Pacing and Defibrillation in Heart Failure) trial, which enrolled NYHA class III to IV patients with QRS 120 ms, included 3 arms: GDMT, CRT-D, and CRT pacemaker (CRT-P).17 The primary end- point of death or hospitalization was decreased

with CRT-P and CRT-D. The CARE-HF (Cardiac Resynchronization Heart Failure) trial included a similar group with NYHA class III to IV, LVEF

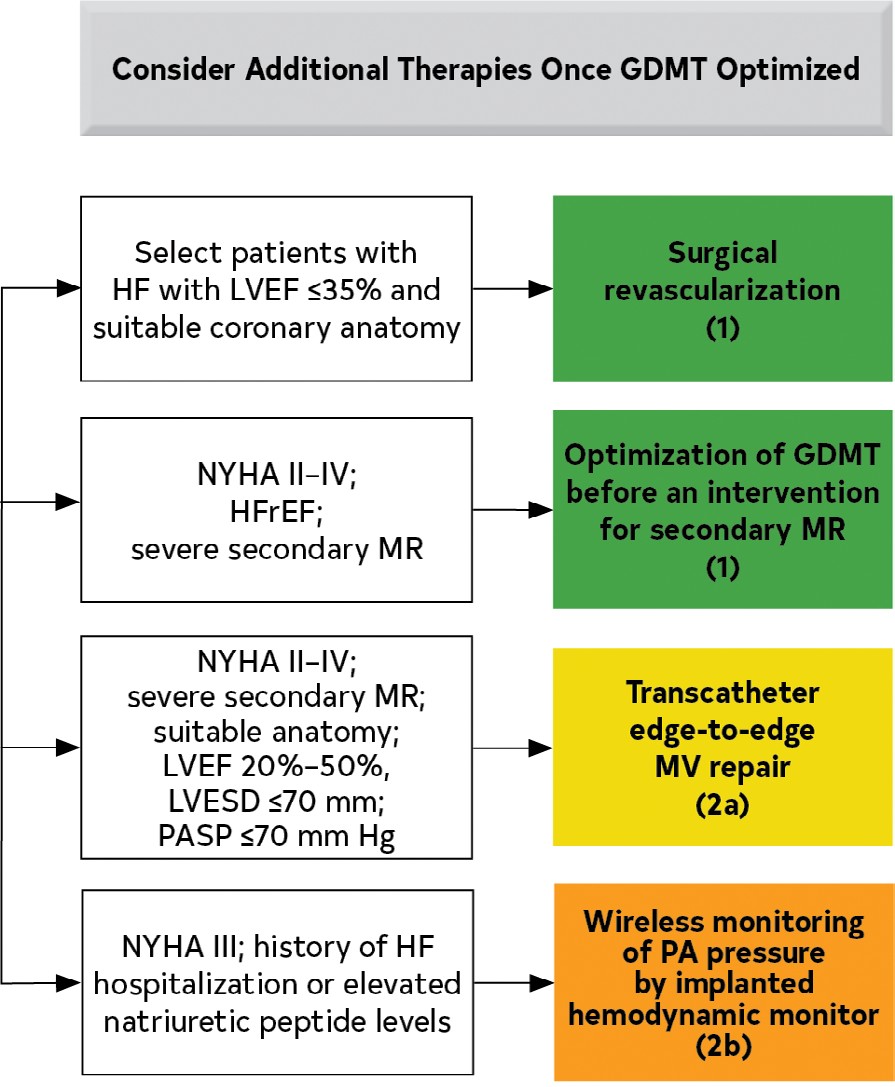
35%, QRS >120 ms, and showed a significant reduction in primary and endpoint of death or hos- pitalization.18 In the REVERSE (Resynchronization Reverses Remodeling in Systolic Left Ventricular Dysfunction) trial, patients with NYHA class I to II and LVEF 40% were randomized to CRT-D on for 1 year and CRT-D off for 1 year or vice versa.19 A HF composite endpoint was less common when CRT was activated. MADIT-CRT enrolled NYHA class I and II HF with LVEF 30% and QRS 130 ms and compared CRT-D with ICD.20 The primary endpoint of death or HF was reduced by CRT-D. The RAFT (Resynchronization-Defibrillation for Ambulatory Heart Failure) trial randomized patients with NYHA class II to III HF, LVEF 30%, QRS

>120 ms, or paced QRS 200 ms and compared

CRT-D with ICD.21 Again, there was a reduction in the primary endpoint of death or HF hospitalization.

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##### Figure 9. Additional Device Therapies.

Colors correspond to COR in Table 2. Recommendations for additional nonpharmaceutical interventions that may be considered for patients with HF are shown. GDMT indicates guideline-directed medical therapy; HF, heart failure; HFH, heart failure hospitalization; HFrEF, heart failure with reduced ejection fraction; IV, intravenous; LVEF, left ventricular ejection fraction; LVESD, left ventricular end systolic dimension; MV, mitral valve; MR, mitral regurgitation; NP, natriuretic peptide; NSR, normal sinus rhythm; NYHA, New York Heart Association; and PASP, pulmonary artery systolic pressure.

* + - 1. The economic value of CRT has been evaluated by 3 RCTs (COMPANION,22 MADIT-CRT,26 and REVERSE23), 2 model-based analyses,25,27 and 1 observational study.24 These analyses consistently found CRT increased survival and QOL in addition to increasing health care costs. However, the eco- nomic value of CRT likely varies as a result of the shown variation in treatment effect.26 Among popu- lations with larger expected mortality reduction and improvement in QOL, such as patients with a LBBB with QRS duration >150 ms, the cost per QALY is

<$60 000.22,26,27 Among other populations expected to have smaller treatment benefit, the economic value is more uncertain. However, a model-based analysis of patients with NYHA class I to II found the incremental cost-effectiveness ratio remained

<$150 000 per QALY with even small reductions in all-cause mortality.27 Therefore, CRT likely provides at least intermediate value for patients with other guideline-indicated recommendations in which CRT is expected to reduce mortality.

* + - 1. Subgroup analysis of the previously mentioned tri- als has informed us of the predictors of benefit,

including longer QRS duration, and LBBB versus non-LBBB.28 The most benefit was gained with wider QRS durations and with LBBB. This was true in COMPANION, CARE-HF, MADIT-CRT, REVERSE, and RAFT.17,29–32 A QRS duration >150

ms was also a predictor of response, and in those with non-LBBB, a prolonged PR predicted benefit in MADIT-CRT but not in REVERSE.33

* + - 1. Extension of benefit to those with LVEF between 35% and 50% has been seen. In the BLOCK-HF (Biventricular versus Right Ventricular Pacing in Heart Failure) trial, patients with NYHA class I to III HF, LVEF 50%, and atrioventricular block randomized to RV pacing or CRT, there was ben- efit to CRT in reduction in the primary outcome of death, urgent HF visit, or 15% increase in LV end systolic volume.34
      2. In the previously mentioned CRT trials, there was some benefit for those with LBBB and QRS dura- tions between 120 and 149, but not as much ben- efit as those with LBBB 150 ms.17,28–32
      3. Several trials have included patients with AF. In the MUSTIC AF (Multisite Stimulation in Cardiomyopathies),44 RAFT,45 and the SPARE (Spanish Atrial Fibrillation and Resynchronization)46 trials, there were benefits in patients with AF, while in COMPANION,47 AF attenuated the ben- efit of CRT. In the PAVE (Post AV Nodal Ablation Evaluation) study, patients with NYHA class II to III, mean LVEF of 46%, and AF undergoing atrioven- tricular node ablation, CRT improved the 6-minute walk test and LVEF compared with those who were RV paced.35
      4. In patients in whom there is an expected high bur- den of ventricular pacing, especially if >40%, CRT may be used to reduce mortality, reduce hospital- izations, and improve symptoms and QOL.35,48
      5. Identification of specific arrhythmogenic genetic variants such as *LMNA/C*, desmosomal proteins, phospholamban, and Filamin-C carry implications for implantation of ICDs for primary prevention of sudden death even in patients who have LVEF

>35%, or <3 months of GDMT. Most patients with *LMNA/C* cardiomyopathy will progress to cardiac transplantation, sometimes precipitated by refrac- tory arrhythmias more than by pump failure.36–38,49

* + - 1. Subgroup analysis of the CRT RCTs has shown that patients with LVEFs 35%, non-LBBB, and QRS duration of 120 to 149 ms and NYHA class III to ambulatory class IV did not derive as much benefit as those with LBBB 120 ms.17,28–32
      2. The MADIT-CRT trial included NYHA class I (and class II) patients with ischemic heart disease, LVEF

30%, and QRS >130 ms.39 Patients with non- ischemic cardiomyopathy were enrolled if they had NYHA class II HF.

* + - 1. Extension of benefit to patients with narrow QRS has been attempted but has generally failed. In the RETHINQ (Cardiac Resynchronization Therapy in Patients with Heart Failure and Narrow QRS) trial, patients with QRS duration <130 ms were ran- domized to CRT or not.40 There was no benefit from CRT, but subgroup analysis showed there was a benefit with QRS durations between 120 and 130 ms. In the ECHO-CRT (Echocardiography Guided Cardiac Resynchronization Therapy) trial, patients with NYHA class III to IV HF, LVEF 35% and a QRS duration 130 ms, and mechanical dysyn- chrony on echocardiography underwent random- ization to CRT.50 There was no benefit to CRT in this trial. And in the LESSER-EARTH (Evaluation of Resynchronization Therapy for Heart Failure) trial, patients with severe LV dysfunction and QRS

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<120 ms derived no benefit from CRT.51 The

NARROW-CRT (Narrow QRS Ischemic Patients Treated With Cardiac Resynchronization Therapy) was the only trial that showed a benefit in a clinical composite score in patients with an indication for an ICD and QRS <120 ms.52

* + - 1. Subgroup analysis of the CRT trials has shown no benefit for those with LVEF 35%, non-LBBB 120 to 149, and NYHA class I-II HF.17,28–32
      2. The 1-year survival is a standard inclusion for ICD and CRT trials.1–9,16–21
    1. ***Other Implantable Electrical Interventions*** Autonomic nervous system modulation is intriguing as a treatment for HFrEF because of the heightened sym- pathetic response and decreased parasympathetic re- sponse in HF.1 Trials of device stimulation of the vagus nerve, spinal cord, and baroreceptors have had mixed responses.2 An implantable device that electrically stimu- lates the baroreceptors of the carotid artery has been approved by the FDA for the improvement of symptoms in patients with advanced HF who are unsuited for treat- ment with other HF devices including CRT. In a prospec- tive, multicenter, RCT with a total of 408 patients with current or recent NYHA class III HF, LVEF 35%, baro- receptor stimulation was associated with improvements in QOL, exercise capacity, and NT-proBNP levels.3 To date, there are no mortality or hospitalization rates results available with this device. Although early trials of vagus nerve stimulation were positive, the largest and latest trial did not show a reduction in mortality and HF hospital- izations.4 Multisite LV pacing studies initially were prom- ising.5,6 However, more recent data have not confirmed benefit, and the larger phase 2 trial was terminated early for low probability of benefit.7 His bundle and left bundle pacing are attractive because they use the intrinsic con- duction system. In observational data, there does appear to be a benefit over RV pacing8; however, comparisons to CRT are limited.9,10 Cardiac contractility modulation

(CCM), a device-based therapy that involves applying relatively high-voltage, long-duration electric signals to the RV septal wall during the absolute myocardial refrac- tory period, has been associated with augmentation of LV contractile performance. CCM is FDA-approved for patients with NYHA class III with LVEF of 25% to 45% who are not candidates for CRT. Four RCTs have shown benefits in exercise capacity and QOL but, as of yet, no benefits in death or hospitalizations.11–14 Most patients in these trials were class III CHF.3

* + 1. ***Revascularization for CAD***

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| **Recommendation for Revascularization for CAD**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| COR | **LOE** | **Recommendation** |
| **1** | **B-R** | 1. In selected patients with HF, reduced EF (EF  35%), and suitable coronary anatomy, surgical revascularization plus GDMT is beneficial to improve symptoms, cardiovascular hospitaliza- tions, and long-term all-cause mortality.1–8 |

## Synopsis

CAD is commonly associated with HF, necessitating re- vascularization in selected patients with angina or HF symptoms. Data from the STICH Trial showed that, com- pared with optimal medical management alone, CABG surgery plus GDMT did not reduce the primary endpoint of all-cause mortality at a median of 56 months; how- ever, at 10 years’ follow-up, CABG+GDMT resulted in significant reductions in all-cause mortality, cardiovas- cular mortality, and death from any cause or cardiovas- cular hospitalization in patients with LVEF 35% and ischemic cardiomyopathy.7,8 Furthermore, a retrospective analysis showed significant reductions in first and recur- rent all-cause, cardiovascular, and HF hospitalizations at 10 years in patients receiving CABG+ optimal medical therapy compared with optimal medical therapy alone.2 Similar benefits from percutaneous coronary intervention revascularization, in this cohort, have not yet been shown in an RCT, although the REVIVED-BCIS2 (Study of Ef- ficacy and Safety of Percutaneous Coronary Intervention to Improve Survival in Heart Failure) trial, which com- pares percutaneous coronary intervention with medical therapy in a similar population, is ongoing.9 Recent data continue to show a benefit of CABG over percutaneous coronary intervention in patients with diabetes, CAD, and LV dysfunction and in patients with left main CAD and moderate or severe LV dysfunction.4,6,10 Figure 9 sum- marizes revascularization and additional device therapy recommendations.

## Recommendation-Specific Supportive Text

* + - 1. CABG has been shown to improve outcomes in patients with left main or left main equivalent

disease and HF.1,4,10–14 Long-term follow-up shows a reduction in all-cause, cardiovascular, and HF hospitalizations and in all-cause and cardiovascu- lar mortality in patients with LV dysfunction who receive CABG and GDMT compared with GDMT alone.2,7 The long-term survival benefit is greater in those with more advanced ischemic cardiomyopa- thy (lower EF or 3-vessel disease) and diminishes with increasing age.5,7 CABG also improves QOL compared with GDMT alone.3 An RCT of CABG combined with surgical ventricular remodeling com- pared with CABG alone did not show a reduction in death or hospitalization, or improvement in symp- toms with surgical ventricular remodeling.15 Surgical ventricular remodeling performed at the time of CABG may be useful in patients with intractable HF, large thrombus, or persistent arrhythmias resulting from well-defined aneurysm or scar, if other thera- pies are ineffective or contraindicated.15,16

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## Valvular Heart Disease

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| **Recommendations for Valvular Heart Disease**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-R** | 1. In patients with HF, VHD should be managed in a multidisciplinary manner in accordance with clini- cal practice guidelines for VHD to prevent wors- ening of HF and adverse clinical outcomes.1–11 |
| **1** | **C-LD** | 2. In patients with chronic severe secondary MR and HFrEF, optimization of GDMT is recom- mended before any intervention for secondary MR related to LV dysfunction.3–5,12–14 |

**Synopsis**

GDMT applies to all patients with HFrEF, irrespective of the presence of VHD. Significant valve disease warrants evaluation by a multidisciplinary team with expertise in VHD, and management should proceed in accordance with the VHD guidelines.15

#### *Mitral Regurgitation*

Optimization of GDMT can improve secondary MR as- sociated with LV dysfunction and obviate the need for intervention.14,16,17 Therefore, optimizing GDMT and re- assessing MR before MV interventions are important. Patients with persistent severe secondary MR despite GDMT may benefit from either surgical or transcatheter repair, depending on clinical scenario. Thus, patient- centric conversation with a multidisciplinary cardiovas- cular team that includes a cardiologist with expertise in HF is essential when considering MV intervention.15 Two RCTs of transcatheter mitral valve edge-to-edge repair (TEER) in patients with HFrEF and severe sec- ondary MR have been performed. The COAPT trial showed significant reduction in HF and all-cause mor-

tality in patients treated with TEER and GDMT com- pared with GDMT alone, while MITRA-FR (Multicentre Study of Percutaneous Mitral Valve Repair MitraClip Device in Patients With Severe Secondary Mitral Re- gurgitation) showed no benefit of TEER over GDMT in reducing death or hospitalization.6 Specifically, trans- catheter edge-to-edge MV repair has been shown to be beneficial in patients with persistent symptoms despite GDMT, appropriate anatomy on transesophageal echo- cardiography and with LVEF between 20% and 50%, LVESD 70 mm, and pulmonary artery systolic pressure

70 mm Hg6 (Figure 10). Optimal management of sec-

ondary MR may depend on the degree of MR relative to LV remodeling.4,5,14,18–22 Disproportionate MR (MR out of proportion to LV remodeling) may respond better to procedural interventions that reduce MR, such as CRT, TEER, and MV surgery. Proportionate MR may respond to measures that reverse LV remodeling and reduce LV volumes, such as GDMT and CRT.

#### *Aortic Stenosis*

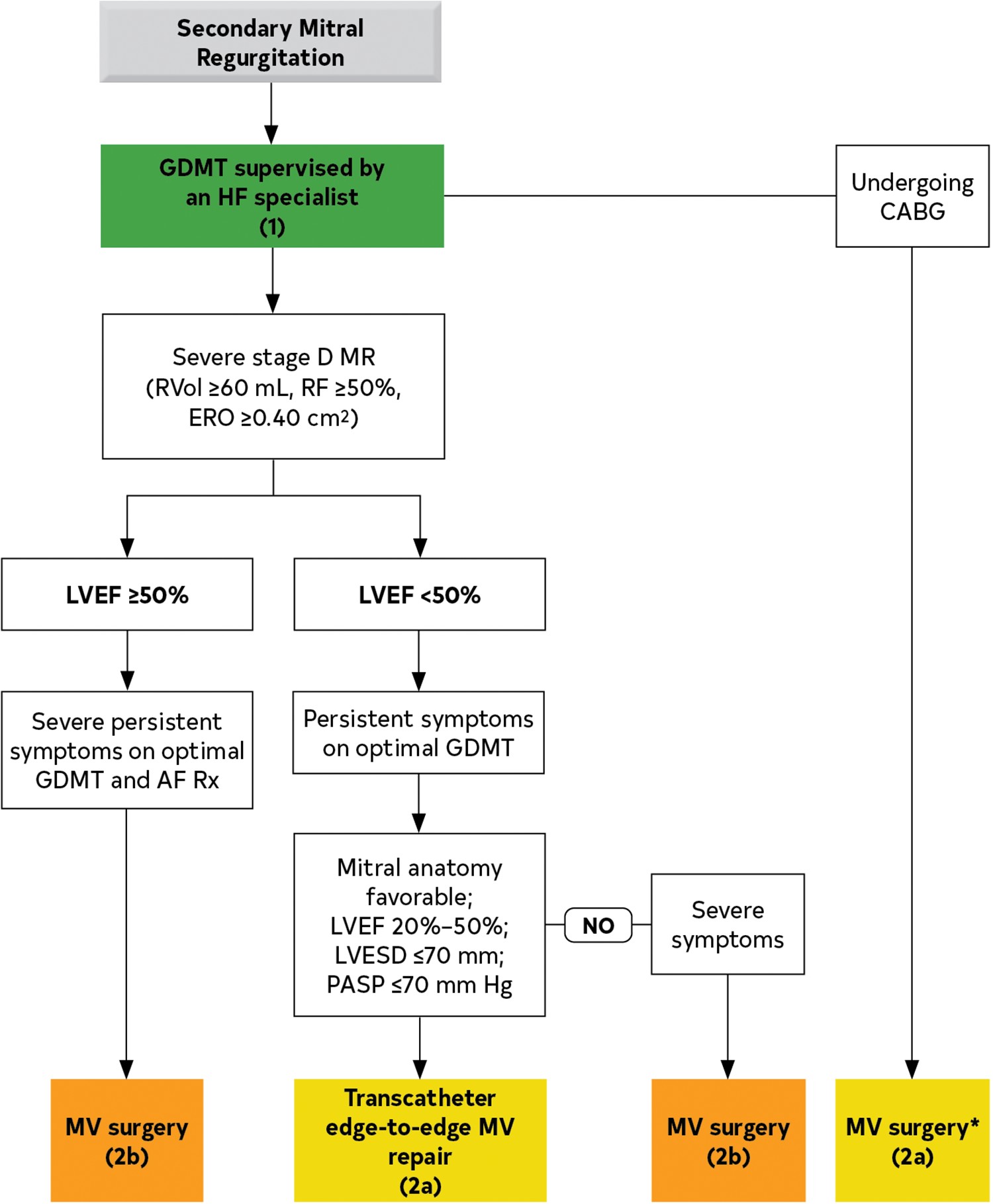
In patients with symptomatic aortic stenosis, transcath- eter and surgical aortic valve repair can improve survival, symptoms, and LV function.15 However, the choice of transcatheter aortic valve implantation versus surgical aortic valve replacement is based on shared decision- making, indications, and assessment of the risk-benefit profile.23,24 The benefit of GDMT in nonsevere aortic ste- nosis and HFrEF is being evaluated in the TAVR UNLOAD (Transcatheter Aortic Valve Replacement to Unload the Left Ventricle in Patients With Advanced Heart Failure) trial.25 GDMT is usually continued in conjunction with clinical surveillance and imaging in patients with nonse- vere aortic stenosis and reduced EF.

#### *Tricuspid Regurgitation*

The severity of secondary tricuspid regurgitation may be dynamic, depending on RV function and pulmonary hypertension, and management entails focusing on underlying causes, such as pulmonary hypertension, RV failure, and HFrEF. Referral to the multidisciplinary team for consideration of intervention might be helpful in patients with refractory tricuspid regurgitation.

## Recommendation-Specific Supportive Text

1. VHD is a significant cause of HF. In patients with HF, management of VHD should be performed by a multidisciplinary team with expertise in HF and VHD, in accordance with the VHD guidelines.15 Cardiologists with expertise in the management of HF are integral to the multidisciplinary team and to guiding the optimization of GDMT in patients with HF and coexisting valve disease. Severe aortic ste- nosis, aortic regurgitation, MR, and tricuspid regur- gitation are associated with adverse outcomes and require timely assessment, optimization of medical



##### Figure 10. Treatment Approach in Secondary Mitral Regurgitation.

Colors correspond to Table 2. AF indicates atrial fibrillation; CABG, coronary artery bypass graft; ERO, effective regurgitant orifice; GDMT, guideline-directed medical therapy; HF, heart failure; LVEF, left ventricular ejection fraction; LVESD, left ventricular end-systolic diameter; MR, mitral regurgitation; MV, mitral valve; PASP, pulmonary artery systolic pressure; RF, regurgitant fraction; RVol, regurgitant volume; and Rx,

medication. \*Chordal-sparing MV replacement may be reasonable to choose over downsized annuloplasty repair. Adapted from Otto CM, et al.15 Copyright 2021 American College of Cardiology Foundation and American Heart Association, Inc.

therapies, and consideration of surgical or transcath- eter interventions accordingly to prevent worsening of HF and other adverse outcomes.1–10,12–20,22–35

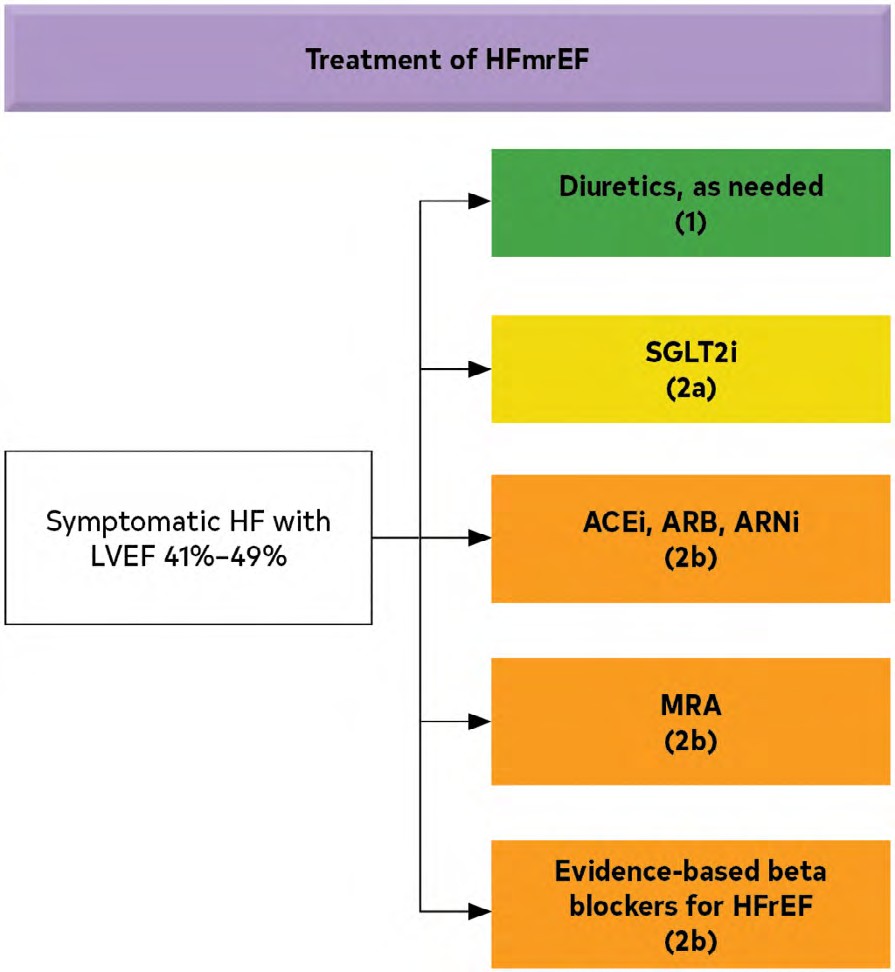
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1. GDMT, including RAAS inhibition, beta blockers, and biventricular pacing, improves MR and LV dimensions in patients with HFrEF and second- ary MR, particularly MR that is proportionate to LV dilatation.1–5,12,13,17 In a small RCT, sacubitril-valsar- tan resulted in a significant reduction in effective regurgitant area and in regurgitant volume when compared with valsartan. The COAPT trial showed a mortality benefit with TEER in patients with

severe secondary MR, LVEF between 20% and 50%, LV end-systolic diameter 70 mm, PA systolic pressure 70 mm Hg, and persistent symptoms (NYHA class II to IV) while on optimal GDMT,28 and these criteria apply when considering TEER. A cardiologist with expertise in the management of HF is integral to shared decision-making for valve intervention and should guide optimization of GDMT to ensure that medical options for HF and secondary MR have been effectively applied for an appropriate time period and exhausted before considering intervention.

pear to respond to medical therapies similarly to patients with HFrEF. Thus, it may be reasonable to treat these pa- tients with GDMT used for treatment of HFrEF. Patients with HFmrEF should have repeat evaluation of LVEF to determine the trajectory of their disease process. Future prospective studies are needed to further clarify treat- ment recommendations for patients with HFmrEF. Figure 11 summarizes COR 1, 2a, and 2b for HFmrEF.



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##### Figure 11. Recommendations for Patients With Mildly Reduced LVEF (41%–49%).

Colors correspond to COR in Table 2. Medication recommendations for HFmrEF are displayed. ACEi indicates angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; ARNi, angiotensin receptor-neprilysin inhibitor; HFmrEF, heart failure with mildly reduced ejection fraction; HFrEF, heart failure with reduced ejection fraction; LVEF, left ventricular ejection fraction; MRA, mineralocorticoid receptor antagonist; and SGLT2i, sodium-glucose cotransporter 2 inhibitor.

## Heart Failure With Mildly Reduced EF (HFmrEF) and Improved EF (HFimpHF)

* + 1. ***HF With Mildly Reduced Ejection Fraction***

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| **Recommendations for HF With Mildly Reduced Ejection Fraction Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **2a** | **B-R** | 1. In patients with HFmrEF, SGLT2i can be ben- eficial in decreasing HF hospitalizations and cardiovascular mortality.1 |
| **2b** | **B-NR** | 2. Among patients with current or previous symp- tomatic HFmrEF (LVEF, 41%–49%), use of evidence-based beta blockers for HFrEF, ARNi, ACEi, or ARB, and MRAs may be considered to reduce the risk of HF hospitalization and cardiovascular mortality, particularly among patients with LVEF on the lower end of this spectrum.2–9 |

## Synopsis

There are no prospective RCTs for patients specifically with HFmrEF (LVEF, 41%–49%). All data for HFmrEF are from post hoc or subsets of analyses from previous HF trials with patients now classified as HFmrEF. LVEF is a spectrum, and among patients with LVEF 41% to 49%, patients with LVEF on the lower end of this spectrum ap-

## Recommendation-Specific Supportive Text

* + - 1. EMPEROR-Preserved (Empagliflozin Outcome Trial in Patients with Chronic Heart Failure with Preserved Ejection Fraction) showed a significant benefit of the SGLT2i, empagliflozin, in patients with symptom- atic HF, with LVEF >40% and elevated natriuretic peptides.1 The 21% reduction in the primary com- posite endpoint of time to HF hospitalization or car- diovascular death was driven mostly by a significant 29% reduction in time to HF hospitalization (nonsig- nificant lower cardiovascular death [HR, 0.91; 95% CI, 0.76–1.0]), with no benefit on all-cause mortality. Empagliflozin also resulted in a significant reduction in total HF hospitalizations, decrease in the slope of the eGFR decline, and a modest improvement in QOL at 52 weeks. Of note, the benefit was simi- lar irrespective of the presence or absence of dia- betes at baseline. In a subgroup of 1983 patients with LVEF 41% to 49% in EMPEROR-Preserved, empagliflozin, a SGLT2i, reduced the risk of the pri- mary composite endpoint of cardiovascular death or hospitalization for HF.1 Although the benefit in the primary endpoint did not have a significant interac- tion by LVEF subgroups (41%–49%, 50%–<60%, and >60%),1 in a subgroup analysis by EF, there was a signal for lower benefit on the primary com- posite endpoint, first and recurrent hospitalizations for HF at higher LVEFs >62.5%.10
      2. Post hoc and subsets of analyses of HFrEF trials that included HFmrEF (LVEF 41%–49%) have suggested benefit from use of GDMT for HFrEF (ie, beta blockers, ARNi, ACEi or ARB, and spironolac- tone).2,3,5–8 The BBmeta-HF (Beta-blockers in Heart Failure Collaborative Group) performed a meta-anal- ysis of 11 HF trials; in a subgroup of 575 patients with LVEF 40% to 49% in sinus rhythm, beta blockers reduced the primary outcome of all-cause and cardiovascular mortality.2 A subgroup analysis of the PARAGON-HF (Prospective Comparison of ARNi with ARB Global Outcomes in HF with Preserved Ejection Fraction) trial for patients with LVEF 45% to 57% (lower range of EFs in the trial) suggested benefit of sacubitril-valsartan versus val- sartan alone (rate ratio, 0.78; 95% CI, 0.64-0.95).3 In a subgroup of 1322 patients with LVEF 41% to 49% in a post hoc analysis of pooled data from the

CHARM (Candesartan in Heart failure-Assessment of Reduction in Mortality and morbidity) trials, can- desartan reduced risk of cardiovascular death and HF hospitalization, the risk of first HF hospitaliza- tion, and the risk of recurrent HF hospitalization.5 In a subgroup of 520 patients with LVEF 44% to 49% in a post hoc analysis of TOPCAT (Treatment of Preserved Cardiac Function Heart Failure With an Aldosterone Antagonist), spironolactone reduced the risk of the primary composite endpoint of car- diovascular death, HF hospitalization, or resusci- tated sudden death, which was mostly caused by a reduction in cardiovascular mortality with spirono- lactone and among patients enrolled in North and South America.6 Spironolactone is preferred among HFmrEF patients with poorly controlled hyperten- sion given previous evidence supporting its use for blood pressure management.1 Continuation of GDMT for patients with improved HFrEF and HFmrEF is important to reduce risk of recrudescent HF.4 Meta-analyses report diverse findings with neu- rohormonal antagonism in patients with HFmrEF, specifying benefit in certain subgroups, underlining the heterogeneity of this phenotype.2,9 Patients with HFmrEF should have repeat evaluation of LVEF to determine the trajectory of their disease process and should undergo testing as clinically indicated to diagnose conditions warranting disease-specific therapy (eg, CAD, sarcoidosis, amyloidosis).

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* + 1. ***HF With Improved Ejection Fraction***

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| **Recommendation for HF With Improved Ejection Fraction**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **1** | **B-R** | 1. In patients with HFimpEF after treatment, GDMT should be continued to prevent relapse of HF and LV dysfunction, even in patients who may become asymptomatic.1 |

## Synopsis

Although GDMT can result in improvement in symptoms, functional capacity, LVEF, and reverse remodeling in patients with HFrEF,2 in most patients, LV function and structural abnormalities do not fully normalize, and symp- toms and biomarker abnormalities may persist or reoc- cur. Many patients deemed to have recovered from HF with resolution of symptoms and improvement of LVEF and natriuretic peptide levels will relapse after withdrawal of GDMT.1 Resolution of symptoms and improvement in cardiac function and biomarkers after treatment does not reflect full and sustained recovery but, rather, remission, which requires treatment to be maintained.3 Stage C HF patients are defined as patients with structural heart dis- ease with previous or current symptoms of HF. In those patients who do not improve (ie, patients who remain

symptomatic or with LV dysfunction), GDMT should not only be continued but also optimized.

## Recommendation-Specific Supportive Text

* + - 1. In an open-label RCT,1 phased withdrawal of HF medications in patients with previous DCM—who were now asymptomatic, whose LVEF had improved from <40% to 50%, whose left ventricular end- diastolic volume (LVEDV) had normalized, and who had an NT-proBNP concentration <250 ng/L— resulted in relapse of cardiomyopathy and HF in 40% of the patients within 6 months. Relapse was defined by at least 1 of these: 1) a reduction in LVEF by >10% and <50%; 2) an increase in LVEDV by >10% and to higher than the normal range; 3) a 2-fold rise in NT-proBNP concentration and to >400 ng/L; or 4) clinical evidence of HF. Treatment was withdrawn successfully in only 50% of patients.1 Secondary analyses showed worsening Kansas City Cardiomyopathy Questionnaire scores, a substantial reduction in LVEF, and nonsignificant increases in NT-proBNP and LV volumes with with- drawal of HF medications.

## Preserved EF (HFpEF)

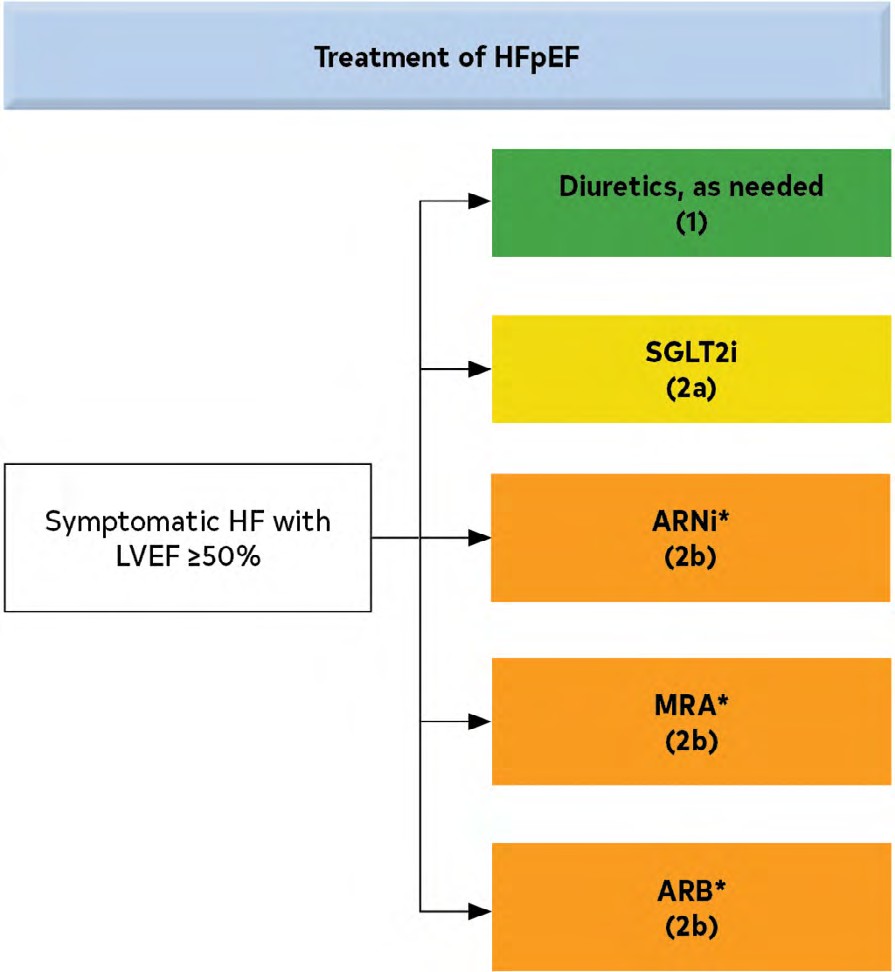
#### *HF With Preserved Ejection Fraction*

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| **Recommendations for HF With Preserved Ejection Fraction\* Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 1. Patients with HFpEF and hypertension should have medication titrated to attain blood pres- sure targets in accordance with published clini- cal practice guidelines to prevent morbidity.1–3 |
| **2a** | **B-R** | 2. In patients with HFpEF, SGLT2i can be ben- eficial in decreasing HF hospitalizations and cardiovascular mortality.4 |
| **2a** | **C-EO** | 3. In patients with HFpEF, management of AF can be useful to improve symptoms. |
| **2b** | **B-R** | 4. In selected patients with HFpEF, MRAs may be considered to decrease hospitalizations, par- ticularly among patients with LVEF on the lower end of this spectrum.5–7 |
| **2b** | **B-R** | 5. In selected patients with HFpEF, the use of ARB may be considered to decrease hospital- izations, particularly among patients with LVEF on the lower end of this spectrum.8,9 |
| **2b** | **B-R** | 6. In selected patients with HFpEF, ARNi may be considered to decrease hospitalizations, par- ticularly among patients with LVEF on the lower end of this spectrum.10,11 |
| **3: No-**  **Benefit** | **B-R** | 7. In patients with HFpEF, routine use of nitrates or phosphodiesterase-5 inhibitors to increase activity or QOL is ineffective.12,13 |

\*See Section 7.2, “Diuretics and Decongestion Strategies in Patients with HF,” and Section 10.2, “Management of Atrial Fibrillation (AF) in HF” for recom- mendations for use of diuretics and management of AF in HF.

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##### Figure 12. Recommendations for Patients With Preserved LVEF (50%).

Colors correspond to COR in Table 2. Medication recommendations for HFpEF are displayed. ARB indicates angiotensin receptor blocker; ARNi, angiotensin receptor-neprilysin inhibitor; HF, heart failure; HFpEF, heart failure with preserved ejection fraction; LVEF, left ventricular ejection fraction; MRA, mineralocorticoid receptor antagonist; and SGLT2i, sodium-glucose cotransporter-2 inhibitor.

\*Greater benefit in patients with LVEF closer to 50%.

## Synopsis

HFpEF (LVEF 50%) is highly prevalent, accounting for up to 50% of all patients with HF, and is associ- ated with significant morbidity and mortality.14 HFpEF is a heterogenous disorder, contributed to by comor- bidities that include hypertension, diabetes, obesity, CAD, CKD, and specific causes such as cardiac amy- loidosis.15–17 Clinical trials have used variable defini- tions of HFpEF (eg, LVEF 40%, 45%, or 50%, and the varying need for accompanying evidence of struc- tural heart disease or elevated levels of natriuretic peptides).18 Until recently, clinical trials had been generally disappointing, with no benefit on mortality and marginal benefits on HF hospitalizations.5,8,11,19,20 Currently, recommended management is that used for HF in general with use of diuretics to reduce congestion and improve symptoms (see Section

* + 1. for recommendations for nonpharmacological management and Section 7.2 for recommendations for diuretics), identification and treatment of spe- cific causes such as amyloidosis, and management of contributing comorbidities such as hypertension, CAD, and AF (see Section 10.2 for recommendations on management of AF). Figure 12 summarizes COR 1, 2a, and 2b for HFpEF.

## Recommendation-Specific Supportive Text

* + - 1. The role of blood pressure control is well established for the prevention of HF, as well as for reduction of other cardiovascular events and HF mortality in patients without prevalent baseline HF.1–3,21–24 The SPRINT (Systolic Blood Pressure Intervention) trial and meta-analyses established that more intensive blood pressure control in patients with high cardiovascular risk significantly reduces HF and other cardiovascular outcomes.2,3,25 In recent clinical practice guidelines for hypertension, blood pressure targets in HFpEF are extrapolated from those for treatment of patients with hypertension in general.26 However, the optimal blood pressure goal and antihypertensive regimens are not known for patients with HFpEF. RAAS antagonists includ- ing ACEi, ARB, MRA, and possibly ARNi, could be first-line agents given experience with their use in HFpEF trials.8,10,16,20,27,28 Beta blockers may be used to treat hypertension in patients with a history of MI,27 symptomatic CAD, or AF with rapid ventricular response. These effects need to be balanced with the potential contribution of chronotropic incompe- tence to exercise intolerance in some patients.29
      2. EMPEROR-Preserved (Empagliflozin Outcome Trial in Patients with Chronic Heart Failure with Preserved Ejection Fraction) showed a significant benefit of the SGLT2i, empagliflozin, in symptom- atic patients with HF with LVEF >40% and ele- vated natriuretic peptides.30 The 21% reduction in the primary composite endpoint of time to HF hospitalization or cardiovascular death was driven mostly by a significant 29% reduction in time to HF hospitalization (nonsignificant lower cardiovascular death [HR, 0.91; 95% CI, 0.76-1.0]), with no benefit on all-cause mortality. Empagliflozin also resulted in a significant reduction in total HF hospitalizations, decrease in the slope of the eGFR decline, and a modest improvement in QOL at 52 weeks. Of note, the benefit was similar irrespective of the pres- ence or absence of diabetes at baseline. Although the benefit in the primary endpoint did not have a significant interaction by LVEF subgroups (<50%, 50%–<60%, and >60%),30 in a subgroup analysis by EF, there was a signal for lower benefit on the primary composite endpoint, first and recurrent HF hospitalizations at higher LVEFs >62.5%.31
      3. Large, randomized clinical trial data are unavail- able to specifically guide therapy in patients with HFpEF and AF. Currently, the comprehensive care of AF can be extrapolated from the clinical practice guidelines for AF, with individualization of strategies for rate or rhythm control in patients with HFpEF (see also Section 10.2, “Management of Atrial

Fibrillation (AF) in HF,” for HF specific recommen- dations for AF). Although beta blockers and nondi- hydropyridine calcium channel blockers are often considered as first-line agents for heart rate con- trol in patients with HFpEF, a recent smaller open- label trial, RATE-AF in elderly patients with AF and symptoms of HF (most with preserved LVEF), compared the use of the beta blocker, bisoprolol, to digoxin.32 At 6 months, the primary endpoint of QOL was similar between the 2 groups. However, sev- eral secondary QOL endpoints, functional capacity, and reduction in NT-proBNP favored digoxin at 12 months. There was a similar heart rate reduction in both groups. Of note, more adverse events such as higher rates of dizziness, lethargy, and hypoten- sion occurred with beta blockers than digoxin. The comprehensive care of AF is beyond the scope of these guidelines. AF-specific care recommenda- tions can be found in separate ACC/AHA clinical practice guidelines.33,34

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* + - 1. MRAs improve diastolic function in patients with HFpEF.35 The TOPCAT trial investigated the effects of spironolactone in patients with HFpEF. The small reduction (HR, 0.89) in the composite of death, aborted cardiac death, and HF hospitaliza- tion was not statistically significant, although HF hospitalization was reduced (HR, 0.83); adverse effects of hyperkalemia and increasing creati- nine levels were more common in the treatment group.5 A post hoc analysis6 showed efficacy in the Americas (HR 0.83) but not in Russia-Georgia (HR 1.10). A sample of the Russia-Georgia popu- lation in the active treatment arm had nondetect- able levels of a spironolactone metabolite. Post hoc analyses have limitations, but they suggest a pos- sibility of benefit in appropriately selected patients with symptomatic HFpEF (LVEF 45%, elevated BNP level or HF admission within 1 year, eGFR

>30 mL/min/1.73 m2, creatinine <2.5 mg/dL, and potassium <5.0 mEq/L). Furthermore, another post hoc analysis suggested that the potential effi- cacy of spironolactone was greatest at the lower end of the LVEF spectrum.7 Careful monitoring of potassium, renal function, and diuretic dosing at initiation and follow-up are key to minimizing the risk of hyperkalemia and worsening renal function.

* + - 1. Although RAAS inhibition strategies have been successful in the treatment of HFrEF, and RAAS activation is suggested in HFpEF,36,37 clinical tri- als with RAAS inhibition have not showed much benefit in patients HFpEF. In the CHARM- Preserved (Candesartan in patients with chronic HF and preserved left-ventricular ejection frac- tion) trial, patients with LVEF >40% were ran- domized to an ARB, candesartan, or to placebo.38 The primary endpoint (cardiovascular death or

HF hospitalization) was not significantly differ- ent between the 2 groups (HR, 0.89; 95% CI,

0.77–1.03, *P*=0.118; covariate-adjusted HR, 0.86; *P*=0.051). Cardiovascular mortality was identical in the 2 groups; HF hospitalizations were lower in the candesartan arm, with borderline statistical signifi- cance on the covariate-adjusted analysis only (HR, 0.84; 95% CI, 0.70–1.00; *P*=0.047; unadjusted

*P*=0.072). The number of individuals hospitalized for HF (reported by the investigator) was lower in the candesartan group than placebo (230 versus 279; *P*=0.017). A post hoc analysis of the CHARM trials showed that improvement in outcomes with candesartan was greater at the lower end the LVEF spectrum.39 In a meta-analysis of 7694 patients with HFpEF in 4 trials evaluating ARB, there was no signal for benefit on cardiovascular mortality (HR, 1.02), all-cause mortality (HR, 1.02), or HF hospitalization (HR, 0.92; 95% CI, 0.83–1.02).40,41

* + - 1. In the PARAMOUNT-HF (Prospective Comparison of ARNi With ARB on Management of Heart Failure With Preserved Ejection Fraction) trial, a phase II RCT in patients with HFpEF (LVEF 45%), sacubitril-valsartan resulted in a lower level of NT-proBNP after 12 weeks of treatment compared with the ARB, valsartan.42 In the PARAGON-HF (Prospective Comparison of Angiotensin Receptor Neprilysin Inhibitor With Angiotensin Receptor Blocker Global Outcomes in Heart Failure and Preserved Left Ventricular Ejection Fraction) trial, in 4822 patients with HFpEF (LVEF 45%, HF admission within 9 months or elevated natri- uretic peptide levels, and eGFR 30 mL/min/m2), sacubitril-valsartan compared with valsartan did not achieve a significant reduction in the primary composite endpoint of cardiovascular death or total (first and recurrent) HF hospitalizations (rate ratio, 0.87; 95% CI, 0.75-1.01; *P*=0.06).10 Given the primary outcome was not met, other analyses are exploratory. There was no benefit of sacubitril- valsartan on cardiovascular death (HR, 0.95) or total mortality (HR, 0.97). There was a signal of benefit for the ARNi for HF hospitalizations (rate ratio, 0.85; 95% CI, 0.72–1.00; *P*=0.056). The occurrence of hyperkalemia and the composite outcome of decline in renal function favored sacu- bitril-valsartan, but it was associated with a higher incidence of hypotension and angioedema. In pre- specified subgroup analyses, a differential effect by LVEF and sex was noted. A benefit of sacubitril- valsartan compared with valsartan was observed in patients with LVEF below the median (45%–57%; rate ratio, 0.78; 95% CI, 0.64–0.95), and in women (rate ratio, 0.73; 95% CI, 0.59–0.90).10,43,44
      2. Nitrate therapy can reduce pulmonary conges-

tion and improve exercise tolerance in patients

with HFrEF. However, the NEAT-HFpEF (Nitrate’s Effect on Activity Tolerance in Heart Failure With Preserved Ejection Fraction) trial45 randomized 110 patients with EF 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. Although the routine use of nitrates in patients with HFpEF does not appear beneficial, patients with HFpEF and symptom- atic CAD may still receive symptomatic relief with nitrates. Phosphodiesterase-5 inhibition augments the nitric oxide system by upregulating cGMP activ- ity. The RELAX (Phosphodiesterase-5 Inhibition to Improve Clinical Status and Exercise Capacity in Heart Failure with Preserved Ejection Fraction) trial13 randomized 216 patients with EF 50% on stable HF therapy and with reduced exercise tol- erance (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.

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## Cardiac Amyloidosis

#### *Diagnosis of Cardiac Amyloidosis*

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| **Recommendations for Diagnosis of Cardiac Amyloidosis Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. Patients for whom there is a clinical suspi- cion for cardiac amyloidosis\*1–5 should have screening for serum and urine monoclonal light chains with serum and urine immuno- fixation electrophoresis and serum free light chains.6 |
| **1** | **B-NR** | 2. In patients with high clinical suspicion for car- diac amyloidosis, without evidence of serum or urine monoclonal light chains, bone scintigra- phy should be performed to confirm the pres- ence of transthyretin cardiac amyloidosis.7 |
| **1** | **B-NR** | 3. In patients for whom a diagnosis of transthyre- tin cardiac amyloidosis is made, genetic testing with *TTR* gene sequencing is recommended to differentiate hereditary variant from wild-type transthyretin cardiac amyloidosis.8 |

\*LV wall thickness 14 mm in conjunction with fatigue, dyspnea, or edema, especially in the context of discordance between wall thickness on echocardiogram and QRS voltage on ECG, and in the context of aortic ste- nosis, HFpEF, carpal tunnel syndrome, spinal stenosis, and autonomic or sensory polyneuropathy.

## Synopsis

Cardiac amyloidosis is a restrictive cardiomyopathy with extracellular myocardial protein deposition, most com- monly monoclonal immunoglobulin light chains (amyloid

cardiomyopathy [AL-CM]) or transthyretin amyloidosis (ATTR-CM). ATTR can be caused by pathogenic vari- ants in the transthyretin gene *TTR* (variant transthyretin amyloidosis, ATTRv) or wild-type transthyretin (wild-type transthyretin amyloidosis, ATTRwt). A diagnostic ap- proach is outlined in Figure 13.9

## Recommendation-Specific Supportive Text

* + - 1. Diagnosis of ATTR-CM requires a high index of suspicion. LV thickening (wall thickness 14 mm) along with fatigue, dyspnea, or edema should trigger consideration of ATTR-CM, especially with discor- dance between wall thickness on echocardiogram and QRS voltage on ECG,10 or other findings such as apical sparing of LV longitudinal strain impair- ment on echocardiography and diffuse late-gado- linium enhancement on cardiac MRI. ATTR-CM is prevalent in severe aortic stenosis,1 HFpEF,2 car- pal tunnel syndrome,3 lumbar spinal stenosis,4 and autonomic or sensory polyneuropathy.5 Practically, screening for the presence of a monoclonal light chain and technetium pyrophosphate (99mTc-PYP) scan can be ordered at the same time for con- venience, but the results of the 99mTc-PYP scan are interpreted only on the context of a nega- tive monoclonal light chain screen. 99mTc-PYP scans may be positive even in AL amyloidosis7 and, thus, a bone scintigraphy scan alone, without con- comitant testing for light chains, cannot distinguish ATTR-CM from AL-CM. Serum free light chain (FLC) concentration and serum and urine immu- nofixation electrophoresis (IFE) are assessed to rule out AL-CM. IFE is preferred because serum plasma electrophoresis and urine plasma electro- phoresis are less sensitive. Together, measurement of serum IFE, urine IFE, and serum FLC is >99% sensitive for AL amyloidosis.6,11
      2. The use of 99mTc bone-avid compounds for bone scintigraphy allows for noninvasive diagnosis of ATTR-CM.7 99mTc compounds include PYP, 3,3-diphosphono-1,2-propanodicarboxylic acid, and hydromethylene diphosphonate, and PYP is used in the United States. In the absence of a light-chain abnormality, the 99mTc-PYP scan is diag- nostic of ATTR-CM if there is grade 2/3 cardiac uptake or an H/CL ratio of >1.5. In fact, the pres- ence of grade 2/3 cardiac uptake in the absence of a monoclonal protein in serum or urine has a very high specificity and positive predictive value for ATTR-CM.7 SPECT is assessed in all positive scans to confirm that uptake represents myocar- dial retention of the tracer and not blood pool or rib uptake signal.12

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##### Figure 13. Diagnostic and Treatment of Transthyretin Cardiac Amyloidosis Algorithm.

Colors correspond to COR in Table 2. AF indicates atrial fibrillation; AL-CM, amyloid cardiomyopathy; ATTR-CM, transthyretin amyloid cardiomyopathy; ATTRv, variant transthyretin amyloidosis; ATTRwt, wild-type transthyretin amyloidosis; CHA2DS2-VASc, congestive heart failure, hypertension, age 75 years, diabetes mellitus, stroke or transient ischemic attack (TIA), vascular disease, age 65 to 74 years, sex category; ECG, electrocardiogram; H/CL, heart to contralateral chest; HFrEF, heart failure with reduced ejection fraction; IFE, immunofixation electrophoresis; MRI, magnetic resonance imaging; NYHA, New York Heart Association; PYP, pyrophosphate; Tc, technetium; and TTR, transthyretin.

* + - 1. If ATTR-CM is identified, then genetic sequenc- ing of the *TTR* gene will determine if the patient has a pathological variant (ATTRv) or wild-type (ATTRwt) disease.12 Differentiating ATTRv from ATTRwt is important because confirmation

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of ATTRv would trigger genetic counseling and potential screening of family members and therapies, inotersen and patisiran, which are presently approved only for ATTRv with polyneuropathy.13,14

* + 1. ***Treatment of Cardiac Amyloidosis***

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| **Recommendations for Treatment of Cardiac Amyloidosis Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-R** | 1. In select patients with wild-type or variant trans- thyretin cardiac amyloidosis and NYHA class I to III HF symptoms, transthyretin tetramer sta- bilizer therapy (tafamidis) is indicated to reduce cardiovascular morbidity and mortality.1 |
| **Value Statement: Low Value (B-NR)** | | 2. At 2020 list prices, tafamidis provides low economic value (>$180000 per QALY gained) in patients with HF with wild-type or variant transthyretin cardiac amyloidosis.2 |
| **2a** | **C-LD** | 3. In patients with cardiac amyloidosis and AF, anticoagulation is reasonable to reduce the risk of stroke regardless of the CHA2DS2-VASc (congestive heart failure, hypertension, age  75 years, diabetes mellitus, stroke or transient ischemic attack [TIA], vascular disease, age 65 to 74 years, sex category) score.3,4 |

## Synopsis

For patients with ATTR-CM and EF 40%, GDMT may be poorly tolerated. The vasodilating effects of ARNi, ACEi, and ARB may exacerbate hypotension, especially with amyloid-associated autonomic dysfunction. Beta blockers may worsen HF symptoms as patients with AT- TR-CM rely on heart rate response to maintain cardiac output. The benefit of ICDs in ATTR-CM has not been studied in randomized trials, and a case-control study showed unclear benefit.5 CRT has not been studied in ATTR-CM with HFrEF. Disease-modifying therapies in- clude TTR silencers (disrupt hepatic synthesis via mRNA inhibition/degradation: inotersen and patisiran), TTR stabilizers (prevent misfolding/deposition: diflunisal and tafamidis), and TTR disruptors (target tissue clearance: doxycycline, tauroursodeoxycholic acid [TUDCA], and epigallocatechin-3-gallate [EGCG] in green tea). Light chain cardiac amyloidosis is managed by hematology- oncology specialists and beyond the scope of cardiolo- gists, but diagnosis is often made by cardiologists when cardiac amyloid becomes manifest (Figure 13). AL amy- loidosis is treatable, and patients with AL amyloidosis with cardiac involvement should promptly be referred to hematology-oncology for timely treatment. Inotersen and patisiran are associated with slower progression of am- yloidosis-related polyneuropathy in ATTRv-CM.6,7 There are ongoing trials of the impact of inotersen and pati- siran and newer generation mRNA inhibitors-degraders on cardiovascular morbidity or mortality. There is limited benefit of diflunisal,8 doxycycline plus TUDCA,9,10 and EGCG,11 on surrogate endpoints such as LV mass, but the impact of these agents on cardiovascular morbid- ity and mortality has not been assessed. Evaluation and management of autonomic dysfunction, volume status, and arrhythmia are important.

## Recommendation-Specific Supportive Text

* + - 1. Tafamidis is currently the only therapy to improve cardiovascular outcomes in ATTR-CM.1 Tafamidis binds the thyroxin-binding site of TTR. In the ATTR-ACT (Safety and Efficacy of Tafamidis in Patients With Transthyretin Cardiomyopathy) clinical trial, a randomized trial of patients with ATTRwt-CM or ATTRv-CM and NYHA class I to III symptoms, tafamidis had lower all-cause mor- tality (29.5% versus 42.9%) and lower cardio- vascular-related hospitalization (0.48 versus 0.70 per year) after 30 months.1 There was a higher rate of cardiovascular-related hospitalizations in patients with NYHA class III HF, potentially attrib- utable to longer survival during a more severe period of disease. Given that tafamidis prevents but does not reverse amyloid deposition, tafami- dis is expected to have greater benefit when administered early in the disease course. As the survival curves separate after 18 months, patients for whom noncardiac disease is not expected to limit survival should be selected. Benefit has not been observed in patients with class IV symp- toms, severe aortic stenosis, or impaired renal function (eGFR <25 mL·min−1·1.73 m−2 body surface area). Tafamidis is available in 2 formula- tions: tafamidis meglumine is available in 20-mg capsules; and the FDA-approved dose is 80 mg (4 capsules) once daily. Tafamidis is also avail- able in 61-mg capsules; the FDA-approved dose for this new formulation is 61 mg once daily.
      2. One model-based analyses used the results of the ATTR-ACT study1 to evaluate the cost-effec- tiveness of chronic tafamidis compared with no amyloidosis-specific therapy among patients with wild-type or variant transthyretin amyloidosis and NYHA class I to III HF.2 With assumptions that tafamidis remained effective beyond the clinical trial duration, they estimated tafamidis increased average survival by 1.97 years and QALY by 1.29. Despite these large clinical benefits, tafamidis (with an annual cost of $225 000) had an incre- mental cost-effectiveness ratio >$180 000 per QALY gained, the benchmark used by this guide- line for low value. The cost of tafamidis would need to decrease by approximately 80% for it to be intermediate value with a cost per QALY

<$180 000.

* + - 1. Intracardiac thrombosis occurs in approximately one-third of patients with cardiac amyloidosis, in some cases in the absence of diagnosed AF3,4,12 and regardless of CHA2DS2-VASc score.13 The use of anticoagulation reduced the risk of intra- cardiac thrombi in a retrospective study.4 The choice of direct oral anticoagulants (DOAC)

##### Table 16. ESC Definition of Advanced HF

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| All these criteria must be present despite optimal guideline-directed treatment: |
| 1. Severe and persistent symptoms of HF (NYHA class III [advanced] or IV) |
| 2. Severe cardiac dysfunction defined by 1 of these: |
| LVEF 30%  Isolated RV failure  Nonoperable severe valve abnormalities Nonoperable severe congenital heart disease  EF 40%, elevated natriuretic peptide levels and evidence of significant  diastolic dysfunction |
| 3. Hospitalizations or unplanned visits in the past 12 mo for episodes of: |
| Congestion requiring high-dose intravenous diuretics or diuretic com- binations  Low output requiring inotropes or vasoactive medications  Malignant arrhythmias |
| 4. Severe impairment of exercise capacity with inability to exercise or low  6-minute walk test distance (<300 m) or peak VO2 (<12–14 mL/kg/min) estimated to be of cardiac origin |
| Criteria 1 and 4 can be met in patients with cardiac dysfunction (as described in criterion 2) but who also have substantial limitations as a result of other conditions (eg, severe pulmonary disease, noncardiac cirrhosis, renal disease). The therapeutic options for these patients may be more limited. |

EF indicates ejection fraction; ESC, European Society of Cardiology; HF, heart failure; LVEF, left ventricular ejection fraction; NYHA, New York Heart As- sociation; RV, right ventricular; and VO2, oxygen consumption/oxygen uptake.

Adapted with permission from Crespo-Leiro et al.1

versus warfarin has not been studied in patients with ATTR, nor has the role of left atrial append- age closure devices. The risk of anticoagulation on bleeding risk in patients with ATTR-CM and AF has not been established. However, although patients with AL amyloidosis may have acquired hemostatic abnormalities, including coagulation factor deficiencies, hyperfibrinolysis, and platelet dysfunction, TTR amyloidosis is not associated with hemostatic defects.

# STAGE D (ADVANCED) HF

## Specialty Referral for Advanced HF

|  |  |  |
| --- | --- | --- |
| **Recommendation for Specialty Referral for Advanced HF** | | |
| **COR** | **LOE** | **Recommendation** |
| **1** | **C-LD** | 1. In patients with advanced HF, when consistent with the patient’s goals of care, timely refer- ral for HF specialty care is recommended to review HF management and assess suitability for advanced HF therapies (eg, LVAD, cardiac transplantation, palliative care, and palliative inotropes).1–6 |

**Synopsis**

A subset of patients with chronic HF will continue to progress and develop persistently severe symptoms despite maximum GDMT. Several terms have been

used to describe this population, including “end-stage,” “advanced,” and “refractory” HF. In 2018, the European Society of Cardiology updated its definition of advanced HF (Table 16), which now includes 4 distinct criteria.1 The revised definition focuses on refractory symptoms rather than cardiac function and more clearly acknowl- edges that advanced HF can occur in patients without severely reduced EF, including those with isolated RV dysfunction, uncorrectable valvular or congenital heart disease, and in patients with preserved and mildly re- duced EF.1,3 The INTERMACS (Interagency Registry for Mechanically Assisted Circulatory Support) has developed 7 profiles that further stratify patients with advanced HF (Table 17).7

Determining that HF and not a concomitant pulmonary disorder is the basis of dyspnea is important. Severely symptomatic patients presenting with a new diagnosis of HF can often improve substantially if they are initially stabilized. Patients should also be evaluated for nonad- herence to medications.8–11 Finally, a careful review of medical management should be conducted to verify that all therapies likely to improve clinical status have been considered.

## Recommendation-Specific Supportive Text

* + 1. Clinical indicators of advanced HF that should trig- ger possible referral to an advanced HF special- ist are shown in Table 18.1,2,12–14 Timely referral for review and consideration of advanced HF therapies is crucial to achieve optimal patient outcomes.15–17 Acronyms such as I-Need-Help…

I, Intravenous inotropes

N, New York Heart Association (NYHA) class IIIB to IV or persistently elevated natriuretic peptides

E, End-organ dysfunction E, EF 35%

D, Defibrillator shocks H, Hospitalizations >1

E, Edema despite escalating diuretics L, Low systolic BP 90, high heart rate

P, Prognostic medication; progressive intolerance or down-titration of GDMT

…have been developed to assist in decision-making for referral to advanced HF.18 Indications and contra- indications to durable mechanical support are listed in Table 19. After patients develop end-organ dysfunc- tion or cardiogenic shock, they may no longer qualify for advanced therapies.19,20 A complete assessment of the patient is not required before referral, because comprehensive, multidisciplinary assessment of car- diac disease and comorbid conditions is routinely performed when evaluating patients for advanced therapies.19,20 Decisions around evaluation and use of advanced therapies should be informed by the pa- tient’s values, goals, and preferences. Discussion with

##### Table 17. INTERMACS Profiles

|  |  |  |
| --- | --- | --- |
| **Profile\*** | **Profile Description** | **Features** |
| 1 | Critical cardiogenic shock | Life-threatening hypotension and rapidly escalating inotropic/pressor support, with critical organ hypoperfusion often confirmed by worsening acidosis and lactate levels. |
| 2 | Progressive decline | “Dependent” on inotropic support but nonetheless shows signs of continuing deterioration in nutrition, renal function, fluid retention, or other major status indicator. Can also apply to a patient with refractory volume over- load, perhaps with evidence of impaired perfusion, in whom inotropic infusions cannot be maintained because of tachyarrhythmias, clinical ischemia, or other intolerance. |
| 3 | Stable but inotrope dependent | Clinically stable on mild-moderate doses of intravenous inotropes (or has a temporary circulatory support de- vice) after repeated documentation of failure to wean without symptomatic hypotension, worsening symptoms, or progressive organ dysfunction (usually renal). |
| 4 | Resting symptoms on oral therapy at home | Patient who is at home on oral therapy but frequently has symptoms of congestion at rest or with activities of daily living (dressing or bathing). He or she may have orthopnea, shortness of breath during dressing or bath- ing, gastrointestinal symptoms (abdominal discomfort, nausea, poor appetite), disabling ascites, or severe lower extremity edema. |
| 5 | Exertion intolerant | Patient who is comfortable at rest but unable to engage in any activity, living predominantly within the house or housebound. |
| 6 | Exertion limited | Patient who is comfortable at rest without evidence of fluid overload but who is able to do some mild ac- tivity. Activities of daily living are comfortable, and minor activities outside the home such as visiting friends or going to a restaurant can be performed, but fatigue results within a few minutes or with any meaningful physical exertion. |
| 7 | Advanced NYHA class III | Patient who is clinically stable with a reasonable level of comfortable activity, despite a history of previous decompensation that is not recent. This patient is usually able to walk more than a block. Any decompensation requiring intravenous diuretics or hospitalization within the previous month should make this person a Patient Profile 6 or lower. |

ICD indicates implantable cardioverter-defibrillator; INTERMACS, Interagency Registry for Mechanically Assisted Circulatory Support; and NYHA, New York Heart Association.

Adapted from Stevenson et al,7 with permission from the International Society for Heart and Lung Transplantation.

\*Modifier options: Profiles 3 to 6 can be modified for patients with recurrent decompensations leading to frequent (generally at least 2 in past 3 mo or 3 in past 6 mo) emergency department visits or hospitalizations for intravenous diuretics, ultrafiltration, or brief inotropic therapy. Profile 3 can be modified in this manner if the patient is usually at home. If a Profile 7 patient meets the modification of frequent hospitalizations, the patient should be moved to Profile 6 or worse. Other modifier options include arrhythmia, which should be used in the presence of recurrent ventricular tachyarrhythmias contributing to the overall clinical course (eg, frequent ICD shocks or requirement of external defibrillation, usually more than twice weekly); or temporary circulatory support for hospitalized patients Profiles 1 to 3.

HF specialists and other members of the multidisci- plinary team may help ensure that the patient has ad- equate information to make an informed decision.

## Nonpharmacological Management: Advanced HF

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| **Recommendation for Nonpharmacological Management: Advanced HF** | | |
| **COR** | **LOE** | **Recommendation** |
| **2b** | **C-LD** | 1. For patients with advanced HF and hypona- tremia, the benefit of fluid restriction to reduce congestive symptoms is uncertain.1–4 |

**Synopsis**

Hyponatremia and diuretic-refractory congestion is common in advanced HF and is associated with poor clinical5,6 and patient-reported outcomes.7 Moreover,

tion has limited-to-no effect on clinical outcomes or diuretic use.4 Although HF nutritional counseling typi- cally focuses on restricting sodium and fluid, patients with advanced HF have the greatest risk of developing cachexia or malnutrition.11 Hence, dietary restrictions and recommendation should be both evidence-based and comprehensive.

## Recommendation-Specific Supportive Text

* + 1. In a registry study of hyponatremia in acute decom- pensated HF, fluid restriction only improved hypo- natremia marginally.1 A registered dietitian-guided fluid and sodium restriction intervention improved NYHA functional classification and leg edema in patients with HFrEF who were not in stage D HF,2 and fluid restriction improved QOL in a pilot RCT of patients with HFrEF and HFpEF (NYHA class I to

improvement in hyponatremia has been shown to improve clinical outcomes.8,9 Fluid restriction is com- monly prescribed for patients with hyponatremia in acute HF but only improves hyponatremia modestly.1 Although restricting fluid is a common recommenda- tion for patients with HF, evidence in this area is of low quality,10 and many studies have not included patients with advanced HF specifically. Moreover, fluid restric-

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IV).3 In a meta-analysis of RCTs on fluid restriction in HF in general, restricted fluid intake compared with free fluid consumption did not result in reduced hospitalization or mortality rates, changes in thirst, the duration of intravenous diuretic use, serum cre- atinine, or serum sodium levels.4 The validity of a previous trial supporting clinical benefits of fluid restriction in HF is in serious question.12

**Table 18. Clinical Indicators of Advanced HF1,2,12,13,23–37**

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|  |
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| Repeated hospitalizations or emergency department visits for HF in the past 12 mo. |
| Need for intravenous inotropic therapy. |
| Persistent NYHA functional class III to IV symptoms despite therapy. |
| Severely reduced exercise capacity (peak VO2, <14 mL/kg/min or <50% predicted, 6-min walk test distance <300 m, or inability to walk 1 block on level ground because of dyspnea or fatigue). |
| Intolerance to RAASi because of hypotension or worsening renal function. |
| Intolerance to beta blockers as a result of worsening HF or hypotension. |
| Recent need to escalate diuretics to maintain volume status, often reaching daily furosemide equivalent dose >160 mg/d or use of supplemental meto- lazone therapy. |
| Refractory clinical congestion. |
| Progressive deterioration in renal or hepatic function. |
| Worsening right HF or secondary pulmonary hypertension. |
| Frequent SBP 90 mm Hg. |
| Cardiac cachexia. |
| Persistent hyponatremia (serum sodium, <134 mEq/L). |
| Refractory or recurrent ventricular arrhythmias; frequent ICD shocks. |
| Increased predicted 1-year mortality (eg, >20%) according to HF survival models (eg, MAGGIC,21 SHFM22). |

HF indicates heart failure; ICD, implantable cardioverter-defibrillator; MAG- GIC, Meta-analysis Global Group in Chronic Heart Failure; NYHA, New York Heart Association; RAASi, renin-angiotensin-aldosterone system inhibitors; SBP, systolic blood pressure; SHFM, Seattle Heart Failure model; and VO2, oxygen consumption/oxygen uptake.

## Inotropic Support

|  |  |  |
| --- | --- | --- |
| **Recommendations for Inotropic Support**  **Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **2a** | **B-NR** | 1. In patients with advanced (stage D) HF refrac- tory to GDMT and device therapy who are eligible for and awaiting MCS or cardiac trans- plantation, continuous intravenous inotropic support is reasonable as “bridge therapy.”1–4 |
| **2b** | **B-NR** | 2. In select patients with stage D HF, despite optimal GDMT and device therapy who are ineligible for either MCS or cardiac transplan- tation, continuous intravenous inotropic sup- port may be considered as palliative therapy for symptom control and improvement in func- tional status.5–7 |
| **3: Harm** | **B-R** | 3. In patients with HF, long-term use of either continuous or intermittent intravenous inotropic agents, for reasons other than palliative care or as a bridge to advanced therapies, is potentially harmful.5,6,8–11 |

**Synopsis**

Despite improving hemodynamic compromise, positive inotropic agents have not shown improved survival in patients with HF in either the hospital or the outpatient setting.6 Regardless of their mechanism of action (eg, in- hibition of phosphodiesterase, stimulation of adrenergic

##### Table 19. Indications and Contraindications to Durable Me- chanical Support37

|  |
| --- |
| Indications (combination of these): |
| Frequent hospitalizations for HF |
| NYHA class IIIb to IV functional limitations despite maximal therapy |
| Intolerance of neurohormonal antagonists |
| Increasing diuretic requirement |
| Symptomatic despite CRT |
| Inotrope dependence |
| Low peak VO2 (<14–16) |
| End-organ dysfunction attributable to low cardiac output |
| Contraindications: |
| Absolute |
| Irreversible hepatic disease |
| Irreversible renal disease |
| Irreversible neurological disease |
| Medical nonadherence |
| Severe psychosocial limitations |
| Relative |
| Age >80 y for destination therapy |
| Obesity or malnutrition |
| Musculoskeletal disease that impairs rehabilitation |
| Active systemic infection or prolonged intubation |
| Untreated malignancy |
| Severe PVD |
| Active substance abuse |
| Impaired cognitive function |
| Unmanaged psychiatric disorder |
| Lack of social support |

CRT indicates cardiac resynchronization therapy; HF, heart failure; NYHA,

New York Heart Association; VO2, oxygen consumption; and PVD, peripheral vascular disease.

or dopaminergic receptors, calcium sensitization), paren- teral inotropes remain an option to help the subset of patients with HF who are refractory to other therapies and are suffering consequences from end-organ hypo- perfusion. In hospitalized patients presenting with doc- umented severe systolic dysfunction who present with low blood pressure and significantly low cardiac index, short-term, continuous intravenous inotropic support may be reasonable to maintain systemic perfusion and preserve end-organ performance.8,11,12 There continues to be lack of robust evidence to suggest the clear benefit of 1 inotrope over another.13 To minimize adverse effects, lower doses of parenteral inotropic drugs are preferred, although the development of tachyphylaxis should be ac- knowledged, and the choice of agent may need to be changed during longer periods of support. Similarly, the ongoing need for inotropic support and the possibility of discontinuation should be regularly assessed. Table 20 compares commonly used inotropes.

##### Table 20. Intravenous Inotropic Agents Used in the Management of HF

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Inotropic Agent** | **Dose (mcg/kg)** | |  | **Effects** | | | |  |  |
| **Bolus** | **Infusion (/min)** | **Drug Kinetics and Metabolism** | **CO** | **HR** | **SVR** | **PVR** | **Adverse Effects** | **Special Considerations** |
| Adrenergic agonists | | | | | | | | | |
| Dopamine | NA | 5–10 | t1/2: 2–20 min | ↑ | ↑ | ↔ | ↔ | T, HA, N,  tissue necrosis | Caution: MAO-I |
| NA | 10–15 | R, H, P | ↑ | ↑ | ↑ | ↔ |  |  |
| Dobutamine | NA | 2.5–20 | t1/2: 2–3 min H | ↑ | ↑ | ↔ | ↔ | ↑/↓BP, HA, T, N, F,  hypersensitivity | Caution: MAO-I; CI: sulfite allergy |
| PDE 3 inhibitor | | | | | | | | | |
| Milrinone | NR | 0.125–0.75 | t1/2: 2.5 h H | ↑ | ↑ | ↓ | ↓ | T, ↓BP | Accumulation may occur in setting of renal failure; monitor kidney function and LFTs |
| Vasopressors | | | | | | | | | |
| Epinephrine | NR | 5–15 mcg/min | t1/2: 2–3 min | ↑ | ↑ | ↑ (↓) | ↔ | HA, T | Caution: MAO-I |
| 15–20 mcg/min | t1/2: 2–3 min | ↑ | ↑↑ | ↑↑ | ↔ | HA, T | Caution: MAO-I |
| Norepinephrine | NR | 0.5–30 mcg/min | t1/2: 2.5 min | ↔ | ↑ | ↑↑ | ↔ | ↓ HR, tissue necrosis | Caution: MAO-I |

BP indicates blood pressure; CI, contraindication; CO, cardiac output; F, fever; H, hepatic; HA, headache; HF, heart failure; HR, heart rate; LFT, liver function test; MAO-I, monoamine oxidase inhibitor; N, nausea; NA, not applicable; NR, not recommended; P, plasma; PDE, phosphodiesterase; PVR, pulmonary vascular resistance; R, renal; SVR, systemic vascular resistance; T, tachyarrhythmias; and t1/2, elimination half-life.

Up arrow means increase. Side arrow means no change. Down arrow means decrease. Up/down arrow means either increase or decrease.

## Recommendation-Specific Supportive Text

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* + 1. More prolonged use of inotropes as “bridge” ther- apy for those awaiting either heart transplantation or MCS may have benefit in reducing pulmonary hypertension and maintaining end-organ perfusion beyond initial stabilization of patients.1–4
    2. The use of inotropes for palliation does carry with it risks for arrhythmias and catheter-related infections, although the presence of an ICD does decrease the mortality associated with arrhythmias. This risk should be shared with patients if there is planned use of inotropes in a patient without an ICD, or in whom the preference is to deactivate the ICD for pallia- tive purposes. The rate of inappropriate shocks for sinus tachycardia is relatively low, and the concomitant use of beta blockers may help in these patients. Patients may elect to have their shocking devices deactivated, especially if they receive numerous shocks.14,15
    3. With the currently available inotropic agents, the benefit of hemodynamic support and sta- bilization may be compromised by increased myocardial oxygen demand and increased arrhythmic burden. As newer agents are devel- oped, more options may not have these known risks. There are investigational inotropic agents that may provide more options for the manage- ment of patients with HF and represent differ- ent classes of agents.16

## Mechanical Circulatory Support

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| --- | --- | --- |
| **Recommendations for Mechanical Circulatory Support**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. In select patients with advanced HFrEF with NYHA class IV symptoms who are deemed to be dependent on continuous intravenous inotropes or temporary MCS, durable LVAD implantation is effective to improve functional status, QOL, and survival.1–18 |
| **2a** | **B-R** | 2. In select patients with advanced HFrEF who have NYHA class IV symptoms despite GDMT, durable MCS can be beneficial to improve symptoms, improve functional class, and reduce mortality.2,4,7,10,12–17,19 |
| **Value Statement: Uncertain Value (B-NR)** | | 3. In patients with advanced HFrEF who have NYHA class IV symptoms despite GDMT, durable MCS devices provide low to intermedi- ate economic value based on current costs and outcomes.20–24 |
| **2a** | **B-NR** | 4. In patients with advanced HFrEF and hemody- namic compromise and shock, temporary MCS, including percutaneous and extracorporeal ventricular assist devices, are reasonable as a “bridge to recovery” or “bridge to decision.”25–29 |

**Synopsis**

MCS is a therapeutic option for patients with advanced HFrEF to prolong life and improve functional capacity. Over the past 10 years, evolution and refinement of temporary and durable options has continued. MCS is differentiated by the implant location, approach, flow characteristics,

pump mechanisms, and ventricle(s) supported. It can be ef- fective for short-term support (hours to days) and for long- term management (months to years). There are anatomic and physiologic criteria that make durable MCS inappropri- ate for some patients; it is most appropriate for those with HFrEF and a dilated ventricle. With any form of MCS, the device will eventually be turned off, whether at the time of explant for transplantation or recovery, or to stop support in a patient who either no longer wishes to continue support, or in whom the continued functioning of an MCS prevents their death from other causes, such as a catastrophic neu- rologic event, or metastatic malignancy.30 This topic should be addressed a priori with patients before discussions about MCS. Particularly with temporary devices, the poten- tial need to either discontinue or to escalate support should be addressed at time of implantation.

**CLINICAL STATEMENTS AND GUIDELINES**

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## Recommendation-Specific Supportive Text

* + 1. Durable LVADs should be considered in selected patients with NHYA class IV symptoms who are deemed dependent on IV inotropes or tempo- rary MCS. The magnitude of the survival ben- efit for durable LVAD support in advanced NYHA class IV patients has progressively improved, with a 2-year survival >80% in recent trials with newer generation LVADs, which approaches the early survival after cardiac transplantation.2 The 2020 INTERMACS (Interagency Registry for Mechanically Assisted Circulatory Support) report showed that 87.6% of recent durable LVAD recipients were categorized as INTERMACS 1 to 3 before their implant surgery.10 It also showed improved mean survival, >4 years for the desti- nation LVAD cohort, and >5 years for bridge-to- transplant patients. Durable LVAD support has also achieved impressive functional improvement and QOL improvement in multiple trials,2,7,31 although patients remain tethered to external electrical power supplies via a percutaneous lead can limit this improvement. Most patients require rehospi- talization within the first year post-implant. These factors emphasize the need for a thorough evalu- ation and patient education before the decision to proceed with the treatment. Appropriate patient selection benefits from review by a multidisci- plinary team that typically includes an HF cardi- ologist, surgeon, social worker, nurse, pharmacist, dietician, and a palliative medicine specialist.
    2. Durable MCS should be considered in patients with NHYA class IV symptoms despite optimal medical therapy or those deemed dependent on IV inotropes. Destination therapy MCS pro- vides considerable survival advantage in addition to improvement in functional status and health- related QOL.1,7,12,32,33 There is no clear 1-risk model

to assess patient risk for complications, but fac- tors such as elevated central venous pressure, pulmonary hypertension, and coagulopathy have been linked to poorer outcomes.15,34–36 In patients who are unable to tolerate anticoagulation after repeated challenges, implantation of a durable MCS is associated with excess morbidity; incidents of pump thrombosis, hemolysis, and ischemic neu- rologic events have been linked to subtherapeu- tic international normalized ratios.37–41 In addition, implantation of MCS in patients with INTERMACS profile of 1 has been associated with poorer out- come, while those ambulatory patients with profiles 5 to 7 might be too well to have large significant benefit, depending on their symptom burden.19 For patients who are initially considered to be trans- plant ineligible because of pulmonary hyperten- sion, obesity, overall frailty, or other reasons, MCS can provide time to reverse or modify these con- ditions.35,42–44 Continuing and uptitrating GDMT in patients with durable MCS is recommended.45

* + 1. Multiple studies evaluated the cost-effectiveness of ventricular assist device implantation for advanced HF between 2012 and 2017.20,21,23 They consis- tently found device implantation was of low eco- nomic value, with incremental cost-effectiveness ratios of $200000 per QALY gained compared with medical therapy alone among patients who potentially underwent subsequent heart transplant and those who were ineligible for heart transplant. In these studies, costs after implantation remained high given high rates of complication and rehos- pitalization. However, these studies used earlier estimates of post-implant outcomes and compli- cation-related costs that have generally improved over time with better care and newer devices.46–48 Additionally, limited recent data suggest improve- ment in health care costs and intermediate eco- nomic value with LVAD among patients with advanced HF who are either eligible or ineligible for subsequent heart transplant.22,24 The improve- ment may result from lower complication rates, increased survival, lower implant costs, and higher estimated QOL. However, given the conflicting data and limited analyses of contemporary data, the current value of LVAD therapy is uncertain.
    2. Temporary MCS can help stabilize patients and allow time for decisions about the appropriateness of transitions to definitive management, such as durable MCS as a bridge or destination therapy, stabilization until cardiac transplantation or, in the case of improvement and recovery, suitability for device removal.45 These patients often present in cardiogenic shock that cannot be managed solely with IV inotropes and in whom other organ function is at risk. Temporary MCS is also appropriate for

use to allow patients to engage in decision-making for durable MCS or transplantation and for deter- mination of recovery of neurologic status.

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## Cardiac Transplantation

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| **Recommendation for Cardiac Transplantation** | | |
| **COR** | **LOE** | **Recommendation** |
| **1** | **C-LD** | 1. For selected patients with advanced HF despite GDMT, cardiac transplantation is indi- cated to improve survival and QOL.1–3 |
| **Value Statement: Intermediate Value (C-LD)** | | 2. In patients with stage D (advanced) HF despite GDMT, cardiac transplantation provides inter- mediate economic value.4 |

**Synopsis**

The evidence that cardiac transplantation provides a mortality and morbidity benefit to selected patients with stage D HF (refractory, advanced) is derived from observational cohorts. Datasets from the International Society for Heart and Lung Transplantation1 and Unit- ed Network of Organ Sharing2 document the median survival of adult transplant recipients to be now >12 years; the median survival of patients with stage D HF without advanced therapies is <2 years. For compari- son, the risk of death becomes greater than survival between 3 and 4 years on an LVAD, regardless of im- plant strategy (eg, bridge-to-transplant, bridge-to-de- cision, destination therapy).3 Improvements in pre- and posttransplant management have also increased more patients to be eligible for transplant, and treated rejec- tion rates in the first year after transplantation are now

<15%.1 Minimizing waitlist mortality while maximizing posttransplant outcomes continues to be a priority in heart transplantation and was addressed with the re- cent changes in donor allocation policy instituted in 2018.5 Several analyses6–11 have confirmed a decrease in waitlist mortality as well as an increase in the use of temporary circulatory support devices, graft ischemic times, and distances between donor and recipient hos- pitals. The impact on posttransplant survival remains uncertain. Multiorgan transplantation remains uncom- mon and reserved for highly selected candidates. In 2018, 7% of all heart transplants involved another or- gan, in addition to the heart.1

## Recommendation-Specific Supportive Text

* + 1. Cardiac transplantation is the established treat- ment for eligible patients with stage D HF refrac- tory to GDMT, device, and surgical optimization. The survival of adult recipients who received a transplantation between 2011 and 2013 at 1, 3, and 5 years is 90.3%, 84.7%, and 79.6%, respec- tively.2 Conditional survival now approaches 15

years.1 Cardiac transplantation also improves functional status and health-related QOL.12–15 Good outcomes can be achieved in patients not only with HF that is primarily cardiovascular in origin, including reversible pulmonary hyperten- sion,16 congenital heart disease,17 and hypertro- phic cardiomyopathy,18 but also in patients with systemic conditions complicated by HF, such as muscular dystrophy,19 sarcoidosis,20 and amyloi- dosis.21 CPET can refine candidate prognosis and selection.22–28 Appropriate patient selection should include integration of comorbidity burden, caretaker status, and goals of care. The listing criteria, evaluation, and management of patients undergoing cardiac transplantation are described by the International Society for Heart and Lung Transplantation.29 The United Network of Organ Sharing Heart Transplant Allocation Policy was revised in 2018 with a broader geographic shar- ing policy and a 6-tiered system to better priori- tize more unstable patients and minimize waitlist mortality.5–11

* + 1. One study evaluated the cost-effectiveness of heart transplantation compared with medical therapy among patients with inotrope-dependent advanced HF.30 This analysis found transplanta- tion was of intermediate value. The results were similar across a broad range of patient age, wait- list duration, and monthly mortality risk with medi- cal therapy.

# PATIENTS HOSPITALIZED WITH ACUTE DECOMPENSATED HF

## Assessment of Patients Hospitalized With Decompensated HF

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| **Recommendations for Assessment of Patients Hospitalized With Decompensated HF** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 1. In patients hospitalized with HF, severity of con- gestion and adequacy of perfusion should be assessed to guide triage and initial therapy.1–5 |
| **1** | **C-LD** | 2. In patients hospitalized with HF, the common precipitating factors and the overall patient trajectory should be assessed to guide appro- priate therapy.5,6 |
| Goals for Optimization and Continuation of GDMT | | |
| **1** | **C-LD** | 3. For patients admitted with HF, treatment should address reversible factors, establish optimal volume status, and advance GDMT toward targets for outpatient therapy.6 |

**Synopsis**

Initial triage includes clinical assessment of the hemo- dynamic profile for severity of congestion and adequa-

##### Table 21. Common Factors Precipitating HF Hospitalization With Acute Decompensated HF

**CLINICAL STATEMENTS AND GUIDELINES**

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|  |
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| ACS |
| Uncontrolled hypertension |
| AF and other arrhythmias |
| Additional cardiac disease (eg, endocarditis) |
| Acute infections (eg, pneumonia, urinary tract) |
| Nonadherence with medication regimen or dietary intake |
| Anemia |
| Hyper- or hypothyroidism |
| Medications that increase sodium retention (eg, NSAID) |
| Medications with negative inotropic effect (eg, verapamil) |

ACS indicates acute coronary syndrome; AF, atrial fibrillation; HF, heart fail- ure; and NSAID, nonsteroidal anti-inflammatory drug.

cy of perfusion.1–5 The diagnosis of cardiogenic shock warrants consideration of recommendations in Sec- tion 9.5, “Evaluation and Management of Cardiogenic Shock,” but any concern for worsening hypoperfusion should also trigger involvement of the multidisciplinary team for hemodynamic assessment and intervention. Initial triage includes recognition of patients with ACS for whom urgent revascularization may be indicated. In the absence of ischemic disease, recent onset with accelerating hemodynamic decompensation may rep- resent inflammatory heart disease, particularly when accompanied by conduction block or ventricular ar- rhythmias.7,8 However, most HF hospitalizations for de- compensation are not truly “acute” but follow a gradual increase of cardiac filling pressures on preexisting structural heart disease, often with precipitating fac- tors that can be identified3,6 (Table 21). Some patients present with pulmonary edema and severe hyperten- sion, which require urgent treatment to reduce blood pressure, more commonly in patients with preserved LVEF. Patients require assessment and management of ischemia, arrhythmia and other precipitating factors and comorbidities. The presenting profile, reversible factors, appropriate workup for the cause of HF includ- ing ischemic and nonischemic causes, comorbidities, and potential for GDMT titration inform the plan of care to optimize the disease trajectory.5

## Recommendation-Specific Supportive Text

* + 1. and 2. Most patients admitted with HF have clinical evidence of congestion without appar- ent hypoperfusion.1–5,9,10 Although elevations of right- and left-sided cardiac filling pressures are usually proportional in decompensation of chronic HF with low EF, up to 1 in 4 patients have a mismatch between right- and left-sided filling pressures.9–11 Disproportionate elevation

of right-sided pressures, particularly with TR, hinders effective decongestion. Disproportionate elevation of left-sided filling pressures may be underrecognized as the cause of dyspnea in the absence of jugular venous distention and edema. Elevated natriuretic peptides can help identify HF in the urgent care setting but with less utility in certain situations, including decreased sen- sitivity with obesity and HFpEF and decreased specificity in the setting of sepsis. Resting hypo- perfusion is often underappreciated in patients with chronic HF but can be suspected from nar- row pulse pressure and cool extremities1,9 and by intolerance to neurohormonal antagonists. Elevated serum lactate levels may indicate hypo- perfusion and impending cardiogenic shock.12 When initial clinical assessment does not sug- gest congestion or hypoperfusion, symptoms of HF may be a result of transient ischemia, arrhythmias, or noncardiac disease such as chronic pulmonary disease or pneumonia, and more focused hemodynamic assessment may be warranted. Assessment of arrhythmia, device profiles such as percent LV pacing versus RV pacing in patients with CRT, and device therapy and shocks in patients with ICD can provide important information.

3. Hospitalization for HF is a sentinel event that signals worse prognosis and the need to restore hemodynamic compensation but also provides key opportunities to redirect the disease trajec- tory. During the HF hospitalization, the approach to management should include and address precipitating factors, comorbidities, and previ- ous limitations to ongoing disease manage- ment related to social determinants of health.1 Patients require assessment and management of ischemia, arrhythmia, and other precipitat- ing factors and comorbidities. The presenting profile, reversible factors, appropriate work-up for cause of HF including ischemic and non- ischemic causes, comorbidities, disease trajec- tory, and goals of care should be addressed. Establishment of optimal volume status is a major goal, and patients with residual conges- tion merit careful consideration for further inter- vention before and after discharge, because they face higher risk for rehospitalization and death.2–5 The disease trajectory for patients hos- pitalized with reduced EF is markedly improved by optimization of recommended medical ther- apies, which should be initiated or increased toward target doses once the efficacy of diure- sis has been shown.13,14

## Maintenance or Optimization of GDMT During Hospitalization

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| **Recommendations for Maintenance or Optimization of GDMT During Hospitalization**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. In patients with HFrEF requiring hospitalization, preexisting GDMT should be continued and optimized to improve outcomes, unless contra- indicated.1–5 |
| **1** | **B-NR** | 2. In patients experiencing mild decrease of renal function or asymptomatic reduction of blood pressure during HF hospitalization, diuresis and other GDMT should not routinely be discontin- ued.6–11 |
| **1** | **B-NR** | 3. In patients with HFrEF, GDMT should be initi- ated during hospitalization after clinical stability is achieved.2,3,5,12–18 |
| **1** | **B-NR** | 4. In patients with HFrEF, if discontinuation of GDMT is necessary during hospitalization, it should be reinitiated and further optimized as soon as possible.19–22 |

**Synopsis**

Hospitalization for HFrEF is a critical opportunity to continue, initiate, and further optimize GDMT.23–25 Continuation of oral GDMT during hospitalization for HF has been shown in registries to lower risk of postdischarge death and readmission compared with discontinuation.1–5 Initiation of oral GDMT during hos- pitalization for HF is associated with numerous clini- cal outcome benefits.2,5,12,16,17 Based on data from the CHAMP-HF (Change the Management of Patients with Heart Failure) registry, however, only 73%, 66%, and 33% of eligible patients with HFrEF were prescribed ACEi-ARB-ARNi, beta blockers, and MRA therapy, respectively.19 Furthermore, based on information ob- tained from claims data, roughly 42% of patients are not prescribed any GDMT within 30 days postindex hospitalization,20 and 45% are prescribed either no oral GDMT or monotherapy within 1-year post-hospitaliza- tion.21 In the management of patients with HFrEF in the community, very few receive target doses of oral GDMT.6 Moreover, most patients with HFrEF have no changes made to oral GDMT over 12 months,21 despite being discharged on suboptimal doses or no GDMT.22 It cannot be assumed that oral GDMT will be initiated or optimized after hospitalization for HFrEF.

## Recommendation-Specific Supportive Text

* + 1. In OPTIMIZE-HF (Organized Program to Initiate Lifesaving Treatment in Hospitalized Patients with Heart Failure), discontinuation of beta blockers was associated with a higher risk for mortality compared with those continued on beta

blockers.1 In a meta-analysis of observational and trial data, discontinuation of beta blockers in hospitalized patients with HFrEF also was asso- ciated with a higher risk of in-hospital mortality, short-term mortality, and the combined endpoint of short-term rehospitalization or mortality.4 Withholding or reducing beta-blocker therapy should be considered in patients with marked volume overload or marginal low cardiac output. In the Get With The Guidelines-Heart Failure (GWTG-HF) registry, withdrawal of ACEi-ARB among patients hospitalized with HFrEF was associated with higher rates of postdischarge mortality and readmission.2 In the COACH (Coordinating study evaluating Outcomes of Advising and Counselling in Heart failure) study, continuation of spironolactone among hospital- ized patients with HFrEF was associated with lower 30-day mortality and HF rehospitaliza- tion.3 From the ARIC (Atherosclerosis Risk in Communities) study, discontinuation of any oral GDMT among patients hospitalized with HFrEF was associated with higher mortality risk.5 Oral GDMT should not be withheld for mild or tran- sient reductions in blood pressure6–9 or mild deteriorations in renal function.10,11 True contra- indications are rare, such as advanced degree atrioventricular block for beta blockers in the absence of pacemakers; cardiogenic shock that may preclude use of certain medications until resolution of shock state; or angioedema for ACEi or ARNi.

* + 1. In CHAMP-HF, very few patients with HF and SBP <110 mm Hg received target doses of beta blockers (17.5%) ACEi-ARB (6.2 %), or ARNi (1.8%).6 In PARADIGM-HF, patients with HF and lower SBP on sacubitril-valsartan had the same tolerance and relative benefit over enalapril compared with patients with higher SBP.7 From the SENIORS (Study of Effects of Nebivolol Intervention on Outcomes and Rehospitalization in Seniors With Heart Failure) trial, nebivolol had equivalent tolerance and benefits irrespective of SBP.8 In Val-HeFT (Valsartan Heart Failure Trial), decreases in SBP did not offset the beneficial effects of valsartan on HF morbidity.9 In patients with HF on oral GDMT, small to moderate worsening of renal function (defined as 20% decrease in eGFR in that study) was not asso- ciated with AKI.10 Moreover, it has been shown that spironolactone and beta blockers might be protective in patients with HF and worsening renal function.11
    2. In OPTIMIZE-HF, discharge use of carvedilol was associated with a reduction in 60- to 90-day mortality and composite risk of mortality

or rehospitalization compared with no carvedilol use.12,13 Discharge use of beta blockers is also associated with lower 30-day all-cause mortality and 4-year all-cause mortality/all-cause read- mission.14 Caution should be used when initiating beta blockers in patients who have required ino- tropes during hospitalization. In GWTG-HF, initia- tion of ACEi-ARB in patients hospitalized with HFrEF reduced 30-day and 1-year mortality.2 Among patients hospitalized with HFrEF, initia- tion of ACEi-ARB also is associated with lower risk of 30-day all-cause readmission and all- cause mortality.15 In a claims study, initiation of MRA therapy at hospital discharge was associ- ated with improved HF readmission but not mor- tality or cardiovascular readmission among older adults hospitalized with HFrEF.16 In COACH, initiating spironolactone among patients hospi- talized with HFrEF was associated with lower 30-day mortality and HF rehospitalization.3 In the PIONEER-HF trial, ARNi use was associ- ated with reduced NT-proBNP levels in patients hospitalized for acute decompensated HF with- out increased rates of adverse events (worsen- ing renal function, hyperkalemia, symptomatic hypotension, angioedema) when compared with enalapril.18 In the ARIC study, initiation of any oral GDMT was associated with reduced 1-year mor- tality among patients hospitalized with HFrEF.5 In SOLOIST-WHF, initiation of sotagliflozin before or shortly after discharge reduced cardiovascular mortality and hospitalization.17

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* + 1. Nearly half (46%) of patients with HFrEF have no changes made to oral GDMT in the 12 months after hospitalization despite many being discharged on suboptimal doses.21 From claims- based studies, 42% of patients with HFrEF are not prescribed any GDMT within 30 days post-index hospitalization,20 and 45% are pre- scribed either no oral GDMT or monotherapy within 1-year post-index hospitalization.21 From CHAMP-HF, initiation or dose increases of beta blockers, ACEi-ARB-ARNi, and MRA occur in

10% of patients with HFrEF within 1 year of hospitalization.22 Very few eligible patients with HFrEF receive target doses of beta blockers (18.7%), ACEi-ARB (10.8%), or ARNi (2.0%).6

Less than 1% of patients with HFrEF are on tar- get doses of ACEi-ARB-ARNi, beta blockers, and MRA within 12 months of an index hospitaliza- tion.22 For patients with HFrEF, there is a graded improvement in the risk of death or rehospitaliza- tion with monotherapy, dual therapy, and triple therapy compared with no GDMT after an index hospitalization in Medicare claims data.21

## Diuretics in Hospitalized Patients: Decongestion Strategy

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| **Recommendations for Diuretics in Hospitalized Patients: Decongestion Strategy**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. Patients with HF admitted with evidence of significant fluid overload should be promptly treated with intravenous loop diuretics to improve symptoms and reduce morbidity.1 |
| **1** | **B-NR** | 2. For patients hospitalized with HF, therapy with diuretics and other guideline-directed medica- tions should be titrated with a goal to resolve clinical evidence of congestion to reduce symp- toms and rehospitalizations.1–6 |
| **1** | **B-NR** | 3. For patients requiring diuretic treatment during hospitalization for HF, the discharge regimen should include a plan for adjustment of diuret- ics to decrease rehospitalizations.7 |
| **2a** | **B-NR** | 4. In patients hospitalized with HF when diuresis is inadequate to relieve symptoms and signs of congestion, it is reasonable to intensify the  diuretic regimen using either: a. higher doses of intravenous loop diuretics.1,3); or b. addition of a second diuretic.3 |

**Synopsis**

Intravenous loop diuretic therapy provides the most rapid and effective treatment for signs and symptoms of congestion leading to hospitalization for HF. Titration to achieve effective diuresis may require doubling of initial doses, adding a thiazide diuretic, or adding an MRA that has diuretic effects in addition to its cardiovascular bene- fits. A major goal of therapy is resolution of the signs and symptoms of congestion before discharge, as persistent congestion scored at discharge has been associated with higher rates of rehospitalizations and mortality. Most patients who have required intravenous diuretic therapy during hospitalization for HF will require prescription of loop diuretics at discharge to decrease recurrence of symptoms and hospitalization.

## Recommendation-Specific Supportive Text

* + 1. Diuretic therapy with oral furosemide was the cornerstone of HF therapy for >20 years before construction of the modern bases of evidence for HF therapies. The pivotal RCTs showing ben- efit in ambulatory HFrEF have been conducted on the background of diuretic therapy to treat and prevent recurrence of fluid retention. An RCT compared intravenous diuretic doses and infusion to bolus dosing during hospitalization for HF but without a placebo arm.1 Protocols for recent trials of other medications in patients hospitalized with HF have all included intravenous diuretic therapy

as background therapy.1–6,8,9 There are no RCTs for hospitalized patients comparing intravenous loop diuretics to placebo, for which equipoise is consid- ered unlikely.10

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**CLINICAL STATEMENTS AND GUIDELINES**

* + 1. Monitoring HF treatment includes careful measure- ment of fluid intake and output, vital signs, standing body weight at the same time each day, and clinical signs and symptoms of congestion and hypoperfu- sion. Daily laboratory tests during active medication adjustment include serum electrolytes, urea nitrogen, and creatinine concentrations. Signs and symptoms of congestion have been specified as inclusion cri- teria in recent trials of patients hospitalized for HF, in which resolution of these signs and symptoms has been defined as a goal to be achieved by hospital dis- charge,1–6,8,9 as it has in the recent HF hospitalization pathway consensus document.11 Evidence of persis- tent congestion at discharge has been reported in 25% to 50% of patients,4,5,12 who have higher rates of mortality and readmission and are more likely to have elevated right atrial pressures, TR, and renal dysfunc- tion. Diuresis should not be discontinued prematurely because of small changes in serum creatinine,13,14 because elevations in the range of 0.3 mg/dL do not predict worse outcomes except when patients are discharged with persistent congestion. Decongestion often requires not only diuresis but also adjustment of other guideline-directed therapies, because elevated volume status and vasoconstriction can contribute to elevated filling pressures.
    2. After discharge, ACEi-ARB, MRAs, and beta block- ers all may decrease recurrent congestion leading to hospitalization in HFrEF. Despite these therapies, most patients with recent HF hospitalization require continued use of diuretics after discharge to pre- vent recurrent fluid retention and hospitalization, as shown in a recent large observational analysis.7 Increases in diuretic doses are frequently required early after discharge even in patients on all other currently recommended therapies for HFrEF.8 It is unknown how increased penetration of therapy with ARNi and SGLT2i will, in the future, affect the dosing of diuretics after discharge with HFrEF.
    3. Titration of diuretics has been described in mul- tiple recent trials of patients hospitalized with HF, often initiated with at least 2 times the daily home diuretic dose (mg to mg) administered intrave- nously.1 Escalating attempts to achieve net diuresis include serial doubling of intravenous loop diuretic doses, which can be done by bolus or infusion, and sequential nephron blockade with addition of a thiazide diuretic, as detailed specifically in the pro- tocol for the diuretic arms of the CARRESS and ROSE trials.3,9 In the DOSE (Diuretic Optimization Strategies Evaluation) trial, there were no signifi- cant differences in patients’ global assessment of

symptoms or in the change in renal function when diuretic therapy was administered by bolus, com- pared with continuous infusion or at a high dose compared with a low dose. Patients in the low-dose group were more likely to require a 50% increase in the dose at 48 hours than were those in the high- dose group, and all treatment groups had higher doses of diuretics compared with baseline pread- mission doses, underlining the necessity to inten- sify and individualize diuretic regimen.1 MRAs have mild diuretics properties and addition of MRAs can help with diuresis in addition to significant cardio- vascular benefits in patients with HF. Addition of low-dose dopamine to diuretic therapy in the set- ting of reduced eGFR did not improve outcomes in a study that included patients with all EFs, but a subset analysis showed increased urine output and weight loss in patients with LVEF <0.40,9 with significant interaction of effect with LVEF. Bedside ultrafiltration initiated early after admission increased fluid loss, with decreased rehospitaliza- tions in some studies when compared with use of diuretics without systematic escalation.15,16and was also associated with adverse events related to the intravenous catheters required.3 Many aspects of ultrafiltration including patient selection, fluid removal rates, venous access, prevention of ther- apy-related complications, and cost require further investigation.

## 9.4a. Parenteral Vasodilation Therapy in Patients Hospitalized With HF

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| **Recommendation for Parenteral Vasodilation Therapy in Patients Hos- pitalized With HF**  **Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **2b** | **B-NR** | 1. In patients who are admitted with decompen- sated HF, in the absence of systemic hypoten- sion, intravenous nitroglycerin or nitroprusside may be considered as an adjuvant to diuretic therapy for relief of dyspnea.1,2 |

**Synopsis**

Vasodilators can be used in acute HF to acutely relieve symptoms of pulmonary congestion in selected patients. Although they may mitigate dyspnea and relieve pulmo- nary congestion, their benefits have not been shown to have durable effects for either rehospitalization or mor- tality benefit. In select patients who present with signs of hypoperfusion such as worsening renal function, even in the absence of hypotension, other escalation of care may need to be considered (see Section 8.3, “Inotropic Support,” and Section 9.5, “Evaluation and Management of Cardiogenic Shock”).

## Recommendation-Specific Supportive Text

**CLINICAL STATEMENTS AND GUIDELINES**

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1. The role for directed vasodilators in acute decom- pensated HF remains uncertain. Part of the rationale for their use is targeting pulmonary congestion, while trying to avoid some potential adverse consequences of loop diuretics. Patients with hypertension, coronary ischemia, or significant MR may be suitable candidates for the use of intravenous nitroglycerin. However, tachyphylaxis may develop within 24 hours, and up to 20% of those with HF may develop resistance to even high doses.3,4 Because of sodium nitroprusside’s potential for producing marked hypotension, invasive hemodynamic blood pressure monitoring (eg, an arte- rial line) is typically required, and nitroprusside is usu- ally used in the intensive care setting; longer infusions of the drug have been associated, albeit rarely, with thiocyanate and cyanide toxicity, particularly in the set- ting of renal insufficiency and significant hepatic dis- ease. Nitroprusside is potentially of value in severely congested patients with hypertension or severe MV regurgitation complicating LV dysfunction.5 Overall, there are no data that suggest that intravenous vaso- dilators improve outcomes in the patient hospitalized with HF; as such, use of intravenous vasodilators is limited to the relief of dyspnea in the hospitalized HF patient with intact or high blood pressure.6,7

## 9.4b. VTE Prophylaxis in Hospitalized Patients

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| **Recommendation for VTE Prophylaxis in Hospitalized Patients Referenced studies that support the recommendation are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendation** |
| **1** | **B-R** | 1. In patients hospitalized with HF, prophylaxis for VTE is recommended to prevent venous throm- boembolic disease.1–3 |

**Synopsis**

HF has long been recognized as affording additional risk for venous thromboembolic disease. When patients are hospitalized for decompensated HF, or when patients with chronic stable HF are hospitalized for other reasons, they are at increased risk for venous thromboembolic disease. The risk may be associated with higher HF symptom bur- den.4 This risk may extend for up to 2 years after hospi- talization but is greatest in the first 30 days.5,6 The use of anticoagulation with subcutaneous low-molecular-weight heparin, unfractionated heparin, fondaparinux, or approved DOAC are used for the prevention of clinically symptomatic deep vein thrombosis and pulmonary embolism.7,8

## Recommendation-Specific Supporting Text

1. Trials using available antithrombotic drugs often were not limited to patients with HF but included

patients with acute illnesses, severe respiratory dis- eases, or simply a broad spectrum of hospitalized medical patients.9–12 All included trials excluded patients perceived to have an elevated risk of bleeding complications or of toxicity from the spe- cific agent tested (eg, enoxaparin in patients with compromised renal function). In some trials, aspirin was allowed but not controlled for as a confounding variable. Despite the increased risk for the develop- ment of VTE in the 30 days after hospitalization, the data for extending prophylaxis to the immediate post-hospital period have shown decreased devel- opment of VTE but were associated with increased bleeding events and overall do not appear to pro- vide additional benefit.2,3,11 For patients admitted specifically for decompensated HF and with ade- quate renal function (creatinine clearance, >30 mL/min), randomized trials suggest that enoxaparin

40 mg subcutaneously once daily,1,13 unfraction- ated heparin 5000 units subcutaneously every 8 or 12 hours,14–17 or rivaroxaban 10 mg once daily11 will radiographically reduce demonstrable venous thrombosis. Effects on mortality or clinically sig- nificant pulmonary embolism rates are unclear. For obese patients, a higher dose of enoxaparin 60 mg once daily achieved target range of thromboprophy- laxis without increased bleeding.12

## Evaluation and Management of Cardiogenic Shock

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| **Recommendations for Evaluation and Management of Cardiogenic Shock Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. In patients with cardiogenic shock, intravenous inotropic support should be used to maintain systemic perfusion and preserve end-organ performance.1–8 |
| **2a** | **B-NR** | 2. In patients with cardiogenic shock, temporary MCS is reasonable when end-organ function cannot be maintained by pharmacologic means to support cardiac function.9–17 |
| **2a** | **B-NR** | 3. In patients with cardiogenic shock, manage- ment by a multidisciplinary team experienced in shock is reasonable.17–22 |
| **2b** | **B-NR** | 4. In patients presenting with cardiogenic shock, placement of a PA line may be considered to define hemodynamic subsets and appropriate management strategies.23–27 |
| **2b** | **C-LD** | 5. For patients who are not rapidly responding to initial shock measures, triage to centers that can provide temporary MCS may be consid- ered to optimize management.17–22 |

**Synopsis**

Cardiogenic shock is a commonly encountered clinical challenge with a high mortality and is characterized by

##### Table 22. Suggested Shock Clinical Criteria\*29

|  |
| --- |
| SBP <90 mm Hg for >30 min: |
| a. Or mean BP <60 mm Hg for >30 min |
| b. Or requirement of vasopressors to maintain systolic BP 90 mm Hg or mean BP 60 mm Hg |
| Hypoperfusion defined by: |
| c. Decreased mentation |
| d. Cold extremities, livedo reticularis |
| e. Urine output <30 mL/h |
| f. Lactate >2 mmol/L |

BP indicates blood pressure; and SBP, systolic blood pressure.

\*Systolic BP and hypoperfusion criteria need to be met for the shock diag- nosis.

a critical reduction in cardiac output manifest by end- organ dysfunction.28 Hypotension (eg, SBP <90 mm Hg) is the primary clinical manifestation of shock but is not sufficient for the diagnosis. Additionally, end-organ hypoperfusion should be present as a consequence of cardiac dysfunction (Tables 22, 23, 24).29 Causes can be broadly separated into acute decompensations of chronic HF, acute myocardial dysfunction without prec- edent HF, and survivors of cardiac arrest. In the case of acute MI, urgent revascularization is paramount. The approach to cardiogenic shock should include its early

##### Table 23. Suggested Shock Hemodynamic Criteria\*29

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| --- |
| 1. SBP <90 mm Hg or mean BP <60 mm Hg |
| 2. Cardiac index <2.2 L/min/m2 |
| 3. Pulmonary capillary wedge pressure >15 mm Hg |
| 1. Other hemodynamic considerations    1. Cardiac power output ([CO x MAP]/451) <0.6 W    2. Shock index (HR/systolic BP) >1.0    3. RV shock       1. Pulmonary artery pulse index [(PASP-PADP)/CVP] <1.0       2. CVP >15 mm Hg       3. CVP-PCW >0.6 |

BP indicates blood pressure; CO, cardiac output; CVP, central venous pres- sure; HR, heart rate; MAP, mean arterial pressure; PADP, pulmonary artery diastolic pressure; PASP, pulmonary artery systolic pressure; PCW, pulmonary capillary wedge; RV, right ventricular; and SBP, systolic blood pressure.

\*Diagnosis of shock requires 1 criteria to be present along with cardiac

index <2.0 L/min/m2 and SBP <90 mm Hg.

recognition, invasive hemodynamic assessment when there is insufficient clinical improvement to initial mea- sures and providing appropriate pharmacological and MCS to optimize end-organ perfusion and prevent meta- bolic complications. The evidence that supports the use of specific pharmacologic therapies and the nature of temporary MCS is primarily gleaned from observational retrospective datasets. Only a few randomized trials have been conducted to assess the most appropriate circula- tory support device, and they have been limited by small

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##### Table 24. Society for Cardiovascular Angiography and Interventions (SCAI) Cardiogenic Shock Criteria29

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| **Stage** | **Bedside Findings** | **Selected Laboratory Markers** | **Hemodynamics** |
| A: At risk Normotensive Normal perfusion  Cause for risk for shock such as large myocardial infarction or HF | Normal venous pressure Clear lungs  Warm extremities Strong palpable pulses  Normal mentation | Normal renal function Normal lactate | SBP >100 mm Hg  Hemodynamics: Normal |
| B: Beginning shock (“pre-shock”) Hypotension  Normal perfusion | Elevated venous pressure Rales present  Warm extremities  Strong pulses Normal mentation | Preserved renal function Normal lactate  Elevated BNP | SBP <90 mm Hg, MAP <60 mm Hg, or >30  mm Hg decrease from baseline SBP HR >100 bpm  Hemodynamics: CI 2.2 L/min/m2 |
| C: Classic cardiogenic shock Hypotension Hypoperfusion | Elevated venous pressure Rales present  Cold, ashen, livedo  Weak or nonpalpable pulses Altered mentation Decreased urine output Respiratory distress | Impaired renal function Increased lactate Elevated BNP Increased LFTs Acidosis | SBP <90 mm Hg; MAP <60 mm Hg; >30  mm Hg from baseline SBP despite drugs and temporary MCS  HR >100 bpm  Hemodynamics: CI 2.2 L/min/m2; PCW  >15 mm Hg; CPO <0.6 W; PAPi <2.0;  CVP-PCW >1.0 |
| D: Deteriorating Worsening hypotension  Worsening hypoperfusion | Same as stage C | Persistent or worsening values of stage C | Escalating use of pressors or MCS to main- tain SBP and end-organ perfusion in setting of stage C hemodynamics |
| E: Extremis  Refractory hypotension Refractory hypoperfusion | Cardiac arrest CPR | Worsening values of stage C laboratories | SBP only with resuscitation PEA  Recurrent VT/VF |

BNP indicates brain natriuretic peptide; CI, cardiac index; CPO, cardiac power output; CPR, cardiopulmonary resuscitation; CVP, central venous pressure; HR, heart rate; LFT, liver function test; MAP, mean arterial blood pressure; MCS, mechanical circulatory support; PAPi, pulmonary artery pulsatility index; PCW, pulmonary capillary wedge pressures; PEA, pulseless electrical activity; SBP, systolic blood pressure; VF, ventricular fibrillation; and VT, ventricular tachycardia.

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sample size, the inherent open-label study design, short follow-up, and surrogate endpoints.

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## Recommendation-Specific Supportive Text

1. Intravenous inotropic support can increase cardiac output and improve hemodynamics in patients presenting with cardiogenic shock. Despite their ubiquitous use for initial manage- ment of cardiogenic shock, there are few pro- spective data and a paucity of randomized trials to guide their use.1–8 However, their broad avail- ability, ease of administration, and clinician famil- iarity favor such agents as the first therapeutic consideration when signs of organ hypoperfu- sion persist despite empiric volume replacement and vasopressors. There is a lack of robust evi- dence to suggest the clear benefit of one inotro- pic agent over another in cardiogenic shock.30 In general, the choice of a specific inotropic agent is guided by blood pressure, concurrent arrhyth- mias, and availability of drug.
2. Despite the lack of direct comparative data, the use of short-term MCS has dramatically increased.9–16,31,32 The hemodynamic benefits of the specific devices vary, and few head-to-head randomized comparisons exist.33–39 Randomized clinical trials are underway that will address the risks and benefits of one modality over another. Vascular, bleeding, and neurologic complications are common to MCS devices, and the risk of such complications should generally be considered in the calculation to proceed with such support.40 As much as possible, an understanding of a patient’s wishes, overall prognosis and trajectory, and assessment of therapeutic risk should precede the use of invasive temporary MCS.
3. Team-based cardiogenic shock management pro- vides the opportunity for various clinicians to pro- vide their perspective and input to the patient’s management.17–22 The escalation of either phar- macological and mechanical therapies should be considered in the context of multidisciplinary teams of HF and critical care specialists, inter- ventional cardiologists, and cardiac surgeons. Such teams should also be capable of provid- ing appropriate palliative care. Most documented experiences have suggested outcomes improve after shock teams are instituted.17–22 In 1 such experience, the use of a shock team was asso- ciated with improved 30-day all-cause mortality (HR, 0.61; 95% CI, 0.41–0.93) and reduced in- hospital mortality (61.0% vs. 47.9%; *P*=0.041).19
4. If time allows, escalation to MCS should be guided by invasively obtained hemodynamic data (eg, PA catheterization). Several observational experiences

have associated PA catheterization use with improved outcomes, particularly in conjunction with short-term MCS.23–27,41 PA catheterization may also be useful when there is diagnostic uncertainty as to the cause of hypotension or end-organ dysfunc- tion, particularly when a patient in shock is not responding to empiric initial shock measures.42

1. Transfer to centers capable of providing such sup- port should be considered early in the assessment of a patient with cardiogenic shock and a trajec- tory of worsening end-organ malperfusion.17–22,43 The treatment of shock should be recognized as a temporizing strategy to support end-organ per- fusion and blood pressure until the cause of the cardiac failure has either been treated (eg, revas- cularization in ST-elevation MI) or recovery (eg, myocarditis) or a definitive solution to the cardiac failure can be accomplished (eg, durable LVAD or transplant). In many cases, pharmacological or MCS can provide sufficient time to address the appropriateness of more definitive therapies (eg, bridge-to-decision) with the patient, family, and the multidisciplinary shock team.

## Integration of Care: Transitions and Team-Based Approaches

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| **Recommendations for Integration of Care: Transitions and Team-Based Approaches**  **Referenced studies that support the recommendations are summarized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-R** | 1. In patients with high-risk HF, particularly those with recurrent hospitalizations for HFrEF, refer- ral to multidisciplinary HF disease management programs is recommended to reduce the risk of hospitalization.1–4 |
| **1** | **B-NR** | 2. In patients hospitalized with worsening HF, patient-centered discharge instructions with a clear plan for transitional care should be pro- vided before hospital discharge.5,6 |
| **2a** | **B-NR** | 3. In patients hospitalized with worsening HF, participation in systems that allow benchmark- ing to performance measures is reasonable to increase use of evidence-based therapy, and to improve quality of care.7–10 |
| **2a** | **B-NR** | 4. In patients being discharged after hospital- ization for worsening HF, an early follow-up, generally within 7 days of hospital discharge, is reasonable to optimize care and reduce rehos- pitalization.11,12 |

**Synopsis**

For patients with HF, the transition from inpatient to outpatient care can be an especially vulnerable period because of the progressive nature of the disease state, complex medical regimens, the large number of comor- bid conditions, and the multiple clinicians who may be

##### Table 25. Important Components of a Transitional Care Plan

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| A transitional care plan, communicated with the patient and their outpatient clinicians before hospital discharge, should clearly outline plans for: |
| Addressing any precipitating causes of worsening HF identified in the hos- pital; |
| Adjusting diuretics based on volume status (including weight) and electro- lytes; |
| Coordination of safety laboratory checks (eg, electrolytes after initiation or intensification of GDMT); |
| Further changes to optimize GDMT, including:  Plans for resuming medications held in the hospital; Plans for initiating new medications;  Plans for titration of GDMT to goal doses as tolerated; |
| Reinforcing HF education and assessing compliance with medical ther- apy and lifestyle modifications, including dietary restrictions and physical activity; |
| Addressing high-risk characteristics that may be associated with poor post- discharge clinical outcomes, such as:  Comorbid conditions (eg, renal dysfunction, pulmonary disease, diabetes, mental health, and substance use disorders);  Limitations in psychosocial support;  Impaired health literacy, cognitive impairment; |
| Additional surgical or device therapy, referral to cardiac rehabilitation in the future, where appropriate; |
| Referral to palliative care specialists and/or enrollment in hospice in selected patients. |

GDMT indicates guideline-directed medical therapy; and HF, heart failure.

involved. Patients are at highest risk for decompensation requiring readmission in the days and weeks post-hospi- tal discharge.13 Optimal transitions of care can decrease avoidable readmissions and improve patient satisfac- tion.14 Multidisciplinary systems of care that promote im- proved communication between health care profession- als, systematic use and monitoring of GDMT, medication reconciliation, and consistent documentation are exam- ples of patient safety standards that should be ensured for all patients with HF transitioning out of the hospital.

## Recommendation-Specific Supportive Text

* + 1. HF disease management programs can help to organize the patient’s care across settings. Potential team members may include cardiologists, primary care clinicians, HF nurses, pharmacists, dieticians, social workers, and community health workers. A Cochrane systematic review of 47 RCTs of disease management interventions after hospi- tal discharge found that interventions that use case management (case manager or nurse coordinates care for high-risk patients) or multidisciplinary approach may decrease all-cause mortality and rehospitalization.3 Disease management programs may comprise education, self-management, medi- cation optimization, device management, weight monitoring, exercise and dietary advice, facilitated access to care during episodes of decompensation, and social and psychological support.14 Disease

management programs coordinated by HF special- ists, including HF nurses, may be best suited for patients with HFrEF; however, there are far fewer data on the effectiveness of disease management programs in patients with HFpEF.2

* + 1. Although hospitalizations for worsening HF are often characterized by rapid changes in medical, surgical, and device therapy to optimize a patient’s clinical status, the patient’s journey with achieving optimal HF care continues beyond hospital dis- charge. Written discharge instructions or educa- tional material given to the patient, family members, or caregiver during the hospital stay or at discharge to home should address all of these: activity level, diet, discharge medications, follow-up appoint- ment, weight monitoring, cardiac rehabilitation, and what to do if symptoms worsen.14 Thorough dis- charge planning that includes special emphasis on ensuring adherence to an evidence-based medica- tion regimen is associated with improved patient outcomes.15,16 Details of the hospital course and the transitional plan of care, with special attention to changes in medications and new medical diag- noses, must be transmitted in a timely and clearly understandable form to all of the patient’s clini- cians who will be delivering follow-up care (Table 25). Any changes in prognosis that will require appropriate care coordination and follow-up post- discharge should be noted.
    2. Systems of care designed to support patients with HF as they move through the continuum of care can improve outcomes.7,14,17,18 Real-time feedback on performance measure benchmarks can improve use of evidence-based therapy and quality of care.8 Quality improvement programs designed to increase the prescription of appropriate discharge medications can increase GDMT prescription at discharge and decrease readmissions and mortal- ity.9 Electronic point-of-care reminders to prescribe GDMT in patients with HFrEF can improve use.10,19 Leveraging transparent health care analytics plat- forms for benchmarking and performance improve- ment may be helpful. There are ongoing studies to determine the most effective strategies to improve evidence-based care.20
    3. Early outpatient follow-up, a central element of transitional care, varies significantly across US hospitals.11 Early postdischarge follow-up may help minimize gaps in understanding of changes to the care plan or knowledge of test results and has been associated with a lower risk of subsequent rehos- pitalization.11,12 Transition of care interventions have often bundled timely clinical follow-up with other interventions, making it challenging to iso- late any unique intervention effects.21 A structured contact with the patient within 7 days of hospital

##### Table 26. Most Common Co-Occurring Chronic Conditions Among Medicare Beneficiaries With HF (N=4947 918), 2011

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| --- | --- | --- | --- | --- | --- |
| **Beneficiaries Age** **65 y (n=4376150)\*** | | | **Beneficiaries Age <65 y (n=571768)†** | | |
|  | **n** | **%** |  | **n** | **%** |
| Hypertension | 3 685373 | 84.2 | Hypertension | 461 235 | 80.7 |
| Ischemic heart disease | 3 145 718 | 71.9 | Ischemic heart disease | 365 889 | 64.0 |
| Hyperlipidemia | 2 623 601 | 60.0 | Diabetes | 338687 | 59.2 |
| Anemia | 2 200 674 | 50.3 | Hyperlipidemia | 325 498 | 56.9 |
| Diabetes | 2 027875 | 46.3 | Anemia | 284 102 | 49.7 |
| Arthritis | 1 901447 | 43.5 | CKD | 257 015 | 45.0 |
| CKD | 1 851 812 | 42.3 | Depression | 207082 | 36.2 |
| COPD | 1 311 118 | 30.0 | Arthritis | 201 964 | 35.3 |
| AF | 1 247748 | 28.5 | COPD | 191 016 | 33.4 |
| Alzheimer’s disease or dementia | 1 207704 | 27.6 | Asthma | 88 816 | 15.5 |

Data source: Centers for Medicare & Medicaid Services administrative claims data, January 2011 to December 2011, from the Chronic Condition Warehouse (CCW), [ccwdata.org.](http://ccwdata.org/)50

AF indicates atrial fibrillation; CKD, chronic kidney disease; COPD, chronic obstructive pulmonary disease; and HF, heart failure.

\*Mean No. of conditions is 6.1; median is 6.

†Mean No. of conditions is 5.5; median is 5.

discharge is a desired goal. Although historically this has been an in-person visit, telemedicine is being increasingly used for chronic management. A pragmatic randomized trial found that an initial telephone visit with a nurse or pharmacist to guide follow-up may reduce the need for in-person vis- its if they are constrained.22 Overall, the timing and method of delivery (in-person clinic versus virtual visit by video or telephone) should be individualized based on patient risk and available care delivery options. Clinical risk prediction tools may help to identify patients at highest risk of postdischarge adverse outcomes.23–25

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# COMORBIDITIES IN PATIENTS WITH HF

## Management of Comorbidities in Patients With HF

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| **Recommendations for the Management of Comorbidities in Patients With HF**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **Management of Anemia or Iron Deficiency** | | |
| **2a** | **B-R** | 1. In patients with HFrEF and iron deficiency with or without anemia, intravenous iron replace- ment is reasonable to improve functional status and QOL.1–4 |
| **3: Harm** | **B-R** | 2. In patients with HF and anemia, erythropoietin- stimulating agents should not be used to improve morbidity and mortality.5,6 |
| **Management of Hypertension** | | |
| **1** | **C-LD** | 3. In patients with HFrEF and hypertension, uptitration of GDMT to the maximally tolerated target dose is recommended.7,8 |

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| **Recommendations for the Management of Comorbidities in Patients With HF (Continued)** | | |
| **COR** | **LOE** | **Recommendations** |
| **Management of Sleep Disorders** | | |
| **2a** | **C-LD** | 4. In patients with HF and suspicion of sleep-dis- ordered breathing, a formal sleep assessment is reasonable to confirm the diagnosis and differentiate between obstructive and central sleep apnea.9,10 |
| **2a** | **B-R** | 5. In patients with HF and obstructive sleep apnea, continuous positive airway pressure may be reasonable to improve sleep quality and decrease daytime sleepiness.9,11–13 |
| **3: Harm** | **B-R** | 6. In patients with NYHA class II to IV HFrEF and central sleep apnea, adaptive servo-ventilation causes harm.11,12 |
| **Management of Diabetes** | | |
| **1** | **A** | 7. In patients with HF and type 2 diabetes, the use of SGLT2i is recommended for the man- agement of hyperglycemia and to reduce HF- related morbidity and mortality.14–17 |

## Synopsis

Multimorbidity is common in patients with HF, with >85% of patients having 2 additional chronic conditions.18,19 Hy- pertension, ischemic heart disease, diabetes, anemia, CKD, morbid obesity, frailty, and malnutrition are among the most common comorbid conditions in patients with HF (Table 26). These chronic conditions complicate the manage- ment of HF and have a significant impact on its prognosis. How to generate specific recommendations addressing many of these conditions in the setting of HF is challeng- ing given the current state of the evidence. For example, although depression is common in patients with HF and strongly impacts QOL and mortality, conventional thera- pies such as antidepressants have not been effective in

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##### Figure 14. Recommendations for Treatment of Patients With HF and Selected Comorbidities.

Colors correspond to COR in Table 2. Recommendations for treatment of patients with HF and select comorbidities are displayed. ACEi indicates angiotensin-converting enzyme inhibitor; AF, atrial fibrillation; ARB, angiotensin receptor blocker; AV, atrioventricular; CHA2DS2-VASc, congestive heart failure, hypertension, age 75 years, diabetes mellitus, stroke or transient ischemic attack [TIA], vascular disease, age 65 to 74 years, sex category; CPAP, continuous positive airway pressure; CRT, cardiac resynchronization therapy; EF, ejection fraction; GDMT, guideline-directed medical therapy; HF, heart failure; HFrEF, heart failure with reduced ejection fraction; IV, intravenous; LVEF, left ventricular ejection fraction; NYHA, New York Heart Association; SGLT2i, sodium-glucose cotransporter-2 inhibitor; and VHD, valvular heart disease. \*Patients with chronic HF with permanent-persistent-paroxysmal AF and a CHA2DS2-VASc score of 2 (for men) and 3 (for women).

improving outcomes.20–22 CKD and HF are closely inter- twined in pathophysiology and have a complex and bidirec- tional relationship.23 Renal dysfunction increases the risk of toxicities of HF therapies and impairs response to diuret- ics.23 The effectiveness of GDMT in patients with HF and concomitant kidney disease is uncertain, because data for treatment outcomes in this patient population are sparse.24 Recommendations surrounding the management of ane- mia, hypertension, diabetes, and sleep disorders that are attributable to the presence of evolving evidence for spe- cific treatment strategies in HF are discussed next. Other comorbidities not addressed in the recommendations are, of course, also important and warrant attention but, be- cause of lack of large-scale trial data, are not addressed as specific recommendations. Figure 14 summarizes COR 1 and 2a for management of select HF comorbidities.

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## Recommendation-Specific Supportive Text Anemia

* + 1. Routine baseline assessment of all patients with HF includes an evaluation for anemia. Anemia is independently associated with HF disease sever- ity and mortality,25 and iron deficiency appears to be uniquely associated with reduced exercise capacity.26 Iron deficiency is usually defined as ferritin level <100 μg /L or 100 to 300 μg/L, if the transferrin saturation is <20%. Intravenous repletion of iron has been shown to improve exer- cise capacity and QOL.1–3,27 The FAIR-HF (Ferric Carboxymaltose Assessment in Patients With Iron Deficiency and Chronic Heart Failure) trial showed significant improvement in NYHA classification,

the 6-minute walk test, and QOL of 459 outpa- tients with chronic HF who received weekly intra- venous ferric carboxymaltose until iron repletion.1 The improvement was independent of the pres- ence of anemia. These findings were confirmed in 2 more recent trials.2,3 The IRONOUT HF (Iron Repletion Effects on Oxygen Uptake in Heart Failure) trial, however, showed no such improve- ment with oral iron supplementation.28 This is attributed to the poor absorption of oral iron and inadequacy of oral iron to replete the iron stores in patients with HF. Therefore, oral iron is not ade- quate to treat iron deficiency anemia in patients with HF. Although these trials were underpowered to detect reductions in hard clinical endpoints, 2 meta-analyses have suggested intravenous iron is associated with a reduction in cardiovascular death and hospitalizations.27,29 Most recently, the AFFIRM-AHF multicenter trial, which included 1132 patients with EF <50% hospitalized for HF, showed a decrease in hospitalization for HF with intravenous ferric carboxymaltose compared to placebo (RR, 0.74; 95% CI, 0.58–0.94) but no reduction in cardiovascular death.4

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* + 1. Anemia in patients with HF is associated with impaired erythropoietin production, with low lev- els found to be associated with worse long-term outcomes.30,31 Although small studies examining the use of erythropoietin-stimulating agents for the treatment of anemia in patients with HF have suggested a trend toward improvement in func- tional capacity and reduction in hospitalization, a high-quality randomized trial of darbepoetin alpha in 2278 patients showed no benefit and an increase in thrombotic events, including stroke.5,6,32 A meta-analysis of 13 trials supports these find- ings.6 Accordingly, erythropoietin-stimulating agent therapy is not recommended for the treatment of anemia in patients with HF.

## Hypertension

* + 1. Clinical trials assessing the impact of goal blood pressure reduction on outcomes in patients with HFrEF and concomitant hypertension are lacking. The optimal blood pressure goal and antihyper- tensive regimen are not known. Antihypertensive therapy is associated with a decrease in the risk of incident HF in the general population,33,34 nota- bly with the more stringent SBP target <120 mm Hg.35 However, low blood pressure, not as a part of an antihypertensive treatment, has been associated with poor outcomes in patients with HFrEF.7,8 This observation may reflect the association between low cardiac output and low blood pressure, rather than the effects of treatment for hypertension.

Nevertheless, hypertension in patients with HFrEF represents an opportunity to maximize GDMT to goal blood pressures defined by the ACC/AHA hypertension guidelines.36

## Sleep Disorders

* + 1. In patients with HF, daytime sleepiness—typically a feature of obstructive sleep apnea—may not reflect the degree of underlying sleep-disordered breathing.37 Hence, the decision to refer a patient for a sleep study should be based on clinical judg- ment. Because the treatment of obstructive sleep apnea and central sleep apnea differ, and because obstructive sleep apnea and central sleep apnea can co-occur,9,11,12 sleep studies can inform clinical decision-making in patients with HF.
    2. In patients with HF and central sleep apnea, con- tinuous positive airway pressure is associated with better sleep quality and nocturnal oxygen- ation9 but has not been shown to affect sur- vival.38 In adults with HFrEF and sleep-disordered breathing, meta-analyses of RCTs have shown that positive airway pressure therapy results in a moderate reduction in BNP39 and improvement in blood pressure and LVEF.40
    3. Adaptive servo-ventilation was associated with increased mortality in 2 RCTs involving patients with HFrEF and central sleep apnea.11,12 Meta- analyses have supported these results.41,42 The weight of evidence does not support the use of adaptive servo-ventilation for central sleep apnea in HFrEF.

## Diabetes

* + 1. The American Diabetes Association guidelines recommend the use of SGLT2i as first-line agent for the treatment of hyperglycemia in patients with diabetes with HF or at high risk of HF.43 SGLT2i are the first class of glucose-lowering agents to receive approval from the FDA for the treatment of HFrEF. Treatment of patients with type 2 diabetes with SGLT2i, including canagliflozin, dapagliflozin, empagliflozin, and sotagliflozin, is associated with a reduction in major adverse cardiovascular events, including hospitalization for HF and cardiovascular death.44 The mechanisms underlying the improve- ment in cardiovascular outcomes attributed to SGLT2i are, however, unknown but appear to be only partially related to the glucosuric effect.45 Two RCTs totaling 8474 patients with NYHA class II, III, or IV HF and EF 40%—DAPA-HF assessing dapagliflozin and EMPEROR-Reduced assessing empagliflozin—showed significant reductions in the relative risk of all-cause death (13%), cardiovascular

death (14%), hospitalization for HF (26%), and renal outcomes (38%) with SGLT2i treatment.14–17 Benefits were consistent across age, sex, and in patients with or without diabetes. Whether dapa- gliflozin or empagliflozin improves outcomes spe- cifically in patients with HFpEF is being studied.46,47 The SOLOIST-WHF trial extends the benefits of SGLT2i to patients with diabetes and acutely decompensated HF.48 Patients on SGLT2i should be closely monitored for potential risks, including severe genitourinary infections and, less commonly, diabetic ketoacidosis.49

## Management of AF in HF

atrioventricular nodal ablation with placement of a CRT device can be considered. Patients with HF, and difficult to control rates, may benefit from atrioventricular node ablation and implantation of a permanent pacemaker if other rate and rhythm control measures fail. If their LVEF is >50%, there is no current evidence that CRT is ben- eficial compared with RV pacing.15,21

## Recommendation-Specific Supportive Text

* + 1. The efficacy of long-term warfarin for the preven- tion of stroke in patients with AF is well established; randomized trials have shown reduced embolic rates and mortality. The AHA/ACC/Heart Rhythm Society guidelines for AF recommend use of the CHA2DS2-VASc score (history of hypertension, age 75 [doubled weight], diabetes mellitus, previ- ous stroke or transient ischemic attack or thrombo- embolism [doubled weight], vascular disease, age 65 to 74 years, sex category) to assess patient risk for adverse outcomes before initiating anticoagu- lation therapy.1,27,28 Regardless of whether patients receive rhythm or rate control, anticoagulation is recommended for patients with HF and AF for stroke prevention with a CHA DS -VASc score of

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| --- | --- | --- |
| **Recommendations for Management of AF in HF**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **A** | 1. Patients with chronic HF with permanent-per- sistent-paroxysmal AF and a CHA2DS2-VASc score of 2 (for men) and 3 (for women) should receive chronic anticoagulant therapy.1–5 |
| **1** | **A** | 2. For patients with chronic HF with permanent- persistent-paroxysmal AF, DOAC is recom- mended over warfarin in eligible patients.2–10 |
| **2a** | **B-R** | 3. For patients with HF and symptoms caused by AF, AF ablation is reasonable to improve symp- toms and QOL.11–14 |
| **2a** | **B-R** | 4. For patients with AF and LVEF 50%, if a rhythm control strategy fails or is not desired, and ventricular rates remain rapid despite medi- cal therapy, atrioventricular nodal ablation with implantation of a CRT device is reasonable.15–22 |
| **2a** | **B-NR** | 5. For patients with chronic HF and permanent- persistent-paroxysmal AF, chronic anticoagulant therapy is reasonable for men and women with- out additional risk factors.23–26 |

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## Synopsis

The interplay between AF and HF is complex. It is clear that AF may worsen HF but also that HF increases the risk of AF. Data from randomized trials support the use of anticoagulation among those with HF and AF but not in patients with HF without AF. Anticoagulation may be accomplished with DOAC or with warfarin when favored because of other indications, cost or drug-drug interac- tions (the DOAC are generally preferred). The choice between rate or rhythm control strategy reflects both patient symptoms and the likelihood of better ventricular function with sinus rhythm. RCTs of rhythm control with antiarrhythmic agents versus rate control have not shown a benefit of rhythm control. More recent RCTs with abla- tion show that ablation may be preferable to antiarrhyth- mic drugs for a rhythm control strategy. Patients thought to have a cardiomyopathy resulting from rapid AF despite attempts at rate control should be aggressively treated to maintain sinus rhythm and, if that is not successful,

2 (for men) and 3 (for women).2–5

* + 1. Trials of DOAC have compared the efficacy and safety with warfarin therapy rather than placebo. Several DOAC are available, including the factor Xa inhibitors apixaban, rivaroxaban, edoxaban, and the direct thrombin inhibitor dabigatran.2–5 These drugs do not need routine anticoagulation monitor- ing or dose adjustment. The fixed dosing together with fewer interactions may simplify patient man- agement, particularly with the polypharmacy com- monly seen in HF, but cost for some patients can be prohibitive when not covered by insurance. These drugs have a potential for an improved ben- efit–risk profile compared with warfarin, which may increase their use in practice, especially in those at increased bleeding risk.6–9 In a meta-analysis of 4 trials examining efficacy and safety of DOAC in patients with and without HF, DOAC more effec- tively reduced the rate of stroke or systemic embo- lism, major bleeding, and intracranial bleeding compared with warfarin, with no treatment hetero- geneity by HF status.10
    2. The 2 largest RCTs of AF ablation in HF showed a benefit in hospitalizations and mortality with ablation.11,12although other smaller trials did not. In the AATAC (Ablation Versus Amiodarone for Treatment of Persistent Atrial Fibrillation in Patients with Congestive Heart Failure and an Implanted Device) trial, 203 patients with persis- tent AF, LVEF <40%, and NYHA class II to III HF, ablation improved the likelihood of maintaining

normal sinus rhythm at 24 months compared with amiodarone and, in addition, had a 45% decrease in hospitalization and decrease in mor- tality (8% vs. 18%).11 The CASTLE AF (Catheter Ablation for Atrial Fibrillation with Heart Failure) trial randomized 363 patients with paroxysmal or persistent AF, LVEF <35%, NYHA class II to IV HF, and ICD to ablation versus standard medi- cal care.12 The composite endpoint of death or rehospitalization was lower in ablation (28.5%) compared with standard care (44.6%). In addi- tion, there was a lower mortality in the ablation group. In a meta-analysis of 11 RCTs comparing rhythm versus rate control, patients undergoing catheter ablation had improved survival (49% relative risk reduction) and reduced hospitaliza- tions (56% relative risk reduction).13

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* + 1. If a rhythm control strategy fails or is undesired, and ventricular rates remain rapid despite medi- cal therapy after all other options are exhausted, atrioventricular nodal ablation with implantation of a CRT device can be considered as a treatment option. Ablate and pace is an old strategy for diffi- cult to rate control AF. Early studies with RV pacing showed benefit.15,16 However, when RV pacing was compared with cardiac resynchronization in more recent trials, especially in those with reduced LVEFs, CRT generally produced more benefit than RV pac- ing.17–21 The PAVE (Left Ventricular-Based Cardiac Stimulation post AV Nodal Ablation Evaluation) and the BLOCK-HF (Biventricular versus Right Ventricular Pacing in Patients with AV block) trials included patients with LVEF >35%, with mean EF 46%22 in PAVE and 40% in BLOCK-HF (enrolled

50%). In both of these trials, patients undergoing CRT had improved outcomes.

* + 1. HF is a hypercoagulable state and serves as an independent risk factor for stroke, systemic embolism, and mortality in the setting of AF.23,24 There are compelling data to support the use of anticoagulation in most patients with HF and con- comitant AF, barring contraindications. In patients with HF and a CHA2DS2-VASc score of 1, those with AF had a 3-fold higher risk compared with individuals without concomitant AF.25 In a post hoc analysis of 2 contemporary HF trials, parox- ysmal and new onset AF were associated with a greater risk for hospitalization caused by HF or stroke.26 In a recent registry study, the risk of stroke was particularly higher in the initial period after diagnosis of HF among patients with preva- lent AF.29 Because HF is a risk factor, additional risk factors may not be required to support the use of anticoagulation in patients with HF, and the decision to anticoagulate can be individual- ized according to risk versus benefit.

# SPECIAL POPULATIONS

## Disparities and Vulnerable Populations\*

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| --- | --- | --- |
| **Recommendations for Disparities and Vulnerable Populations Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 1. In vulnerable patient populations at risk for health disparities, HF risk assessments and multidisciplinary management strategies should target both known risks for CVD and social determinants of health, as a means toward elimination of disparate HF outcomes.1–6 |
| **1** | **C-LD** | 2. Evidence of health disparities should be moni- tored and addressed at the clinical practice and the health care system levels.7–13 |

\*This section crosslinks to Section 7.1.1, “Stage C Nonpharmacological In- terventions and Self-Care Support in HF,” where screening and interventions for social determinants of health are now addressed.

## Synopsis

There are important differences in HF incidence, risk factors, clinical care needs, and outcomes between spe- cific patient populations2,3,14,15 (Table 27). It is essential that HF clinicians be aware of the biological factors, social determinants of health, and implicit biases that impact the burden of disease, clinical decision-making, and effective delivery of GDMT.9,16–18 Women generally present with HF later in life, with more comorbidities and lower patient-reported health status than men.10,19 Sur- vival for women with HF is generally more favorable,20 although access to specialty care may be lower.21–24 The highest incident of HF is consistently observed in self-identified Black patients.25,26 HF hospitalization and mortality rates for Black patients are also higher than for White patients, with the gap increasing over time for young men.2,4,27 These differences are driven mostly by social circumstances; a biological premise or genetic ex- planation for disease or disease severity should not be inferred by race or ethnicity.28 Older patients with HF are especially vulnerable to polypharmacy, multimorbid- ity, cognitive decline, and frailty.29,30 Important strate- gies to remove biases within health care professionals and systems impacting minority and socioeconomically disadvantaged patient populations include implicit bias training, recruiting a diverse workforce, and promoting broad access to HF care.28,31–35

## Recommendation-Specific Supportive Text

* + 1. Hypertension is significantly more prevalent in Black patients, compared with White patients, pop- ulations in the United States, with a younger age of onset and greater attributable cardiovascular risks.36,37 An estimated 50 000 to 350 000 immi- grants to the United States from Mexico and Central America may have asymptomatic *Trypanosoma*

##### Table 27. Risk of HF and Outcomes in Special Populations

|  |  |  |
| --- | --- | --- |
| **Vulnerable Population** | **Risk of HF** | **HF Outcomes** |
| Women | The lifetime risk of HF is equivalent between sexes, but HFpEF risk is higher in women—in FHS participants with new-onset HF, odds of HFpEF (EF >45%) are 2.8-fold higher in women than in men.66  Sex-specific differences in the predictive value of cardiac bio- markers for incident HF.67  Nontraditional cardiovascular risk factors, including anxiety, de- pression, caregiver stress, and low household income may contrib- ute more toward incident heart disease in women than men.68 | Overall, more favorable survival with HF than men. In the OPTI- MIZE-HF registry, women with acute HF had a lower 1-y mortality (HR, 0.93; 95% CI, 0.89–0.97), although women are more likely not to receive optimal GDMT.20,69–71  Lower patient-reported quality of life for women with HFrEF, com- pared with men.10,71  Greater transplant waitlist mortality for women but equivalent sur- vival after heart transplantation or LVAD implantation.24,52 |
| Older adults | Per FHS, at 40 y of age, the lifetime risk of incident HF is 20% for both sexes; at 80 y of age, the risk remains 20% for men and women despite the shorter life expectancy.72  LVEF is preserved in at least two-thirds of older adults with the diagnosis of HF.73 | Among 1233 patients with HF aged 80 y, 40% mortality during mean 27-mo follow-up; survival associated with prescription of GDMT.74 |
| Lower socioeconomic status populations | Among 27078 White and Black adults of low income (70% earned <$15 000/y) participating from 2002–2009 in the Southern Community Cohort Study, a 1 interquartile increase in neighborhood deprivation index was associated with a 12% in- crease in risk of HF (adjusted HR, 1.12; 95% CI, 1.07–1.18).46 | Age-adjusted 1999–2018 HF mortality (deaths/100 000; mean and 95% CI) was higher with increasing quartiles of ADI, which is based on 17 indicators of employment, poverty, and education:  Quartile 1, 20.0 (19.4–20.5);  Quartile 2, 23.3 (22.6–24.0);  Quartile 3, 26.4 (25.5–27.3);  Quartile 4, 33.1 (31.8–34.4).6 |
| Black populations | In MESA, patients of Black race had highest risk of incident HF (4.6/1000 person-years) and highest proportion of nonisch- emic incident HF.26  Higher prevalence of HF risk factors including hypertension, obesity, and diabetes, compared with White populations.75 | CDC data show race-based differences in HF mortality over time: Black men had a 1.16-fold versus 1.43-fold higher age-adjusted HF-related CVD death rate compared with White men in 1999 ver- sus 2017; Black women had a 1.35-fold versus 1.54-fold higher age-adjusted HF-related CVD death rate compared with White women in 1999 versus 2017.27  Gap in outcomes is more pronounced among younger adults (35– 64 y of age) versus older adults (65–84 y of age); age-adjusted HF-related CVD death rates were 2.60-fold and 2.97-fold higher in young Black versus White men and women, respectively.27  Higher rates of hospitalization3 and mortality among patients with HFpEF.76  Lower 5-year survival after heart transplant.77–79 |
| Hispanic populations | MESA study showed higher HF incidence in Hispanic com- pared with non-Hispanic White groups (3.5 versus 2.4 per 1000 person-years) but lower than for African Americans (4.6/1000 person-years).7,26,80 | Despite higher rates of hospitalization for HF compared with non- Hispanic Whites, Hispanic patients with HF have shown lower short-term mortality rates.81  In GWTG, Hispanic patients with HFpEF had lower mortality (OR, 0.50; 95% CI, 0.31–0.81) than non-Hispanic Whites, but this was not the case for Hispanic patients with HFrEF (OR, 0.94; 95% CI, 0.62–1.43).82  Lower risk of developing AF in the setting of HF, compared with White patients.83 |
| Asian and Pacific Islander populations | Limited population-specific data for Asian and Pacific Islander subgroups in the United States.84,85 | High rates of preventable HF hospitalization observed in some Asian and Pacific Islander populations.13  Lower mortality rates from HF for Asian subgroups when listed as the primary cause of death, compared with non-Hispanic White groups.86 |
| Native American and Alaskan Native populations | Limited population-specific data, with cardiovascular risk fac- tor trends best characterized by the Strong Heart Study and Strong Heart Family Study, demonstrating high rates of hyper- tension and diabetes.11,87 | Limited data suggest HF mortality rates in American Indians and Alaska Natives are similar to those in White populations.88 |

CDC indicates Centers for Disease Control and Prevention; CVD, cardiovascular disease; FHS, Framingham Heart Study; GDMT, guideline-directed medical therapy; GWTG, Get With The Guidelines registry; HF, heart failure; HFpEF, heart failure with preserved ejection fraction; HFrEF, heart failure with reduced ejection fraction; HR, hazard ratio; LVAD, left ventricular assist device; LVEF, left ventricular ejection fraction; MESA, Multi-Ethnic Study of Atherosclerosis; OPTMIZE-HF, Organized Program To Initiate Lifesaving Treatment In Hospitalized Patients With Heart Failure; and OR, odds ratio.

*cruzi*, with 20% progressing to Chagas cardiomy- opathy.38 Diabetes is highly prevalent in Southeast Asian and Pacific Islander populations and more strongly associated with poor HF outcomes.39,40 Among patients with established HF, social and medical vulnerabilities can impede successful

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delivery of GDMT and are associated with poorer outcomes.5,41 Among older adults, low income, social isolation, and lack of caregiver support increase HF mortality and low QOL.9,18,42 Nursing home residents, and elderly inpatients with acute HF, are at risk of inadequate GDMT prescription,

although interventions in nursing facilities can improve care delivery for HF.30,43–45 Lower socio- economic status is associated with HF incidence and HF mortality.6,46,47 Homelessness,48 substance use, food insecurity, and lack of transportation each represent potential barriers to optimal dis- ease management.49 Case management and social work services are essential to the comprehensive multidisciplinary HF team approach for coordinat- ing complex medical, psychiatric, and social needs across multiple sectors.

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* + 1. Health care system factors are a potential source of disparate HF care delivery and outcomes. Women are less likely to receive discharge instructions for HF,50 less likely to be referred to specialty care,21,22 and less likely to receive a heart transplantation,51–54 compared with men. Patients with HF of Black race have been identified as less likely to receive care from a cardiologist during an ICU admission for HF,55 have less access to specialized inpatient HF care,12 and may be vul- nerable to clinician biases during evaluation for advanced HF therapies.11,53 Hispanic patients are disproportionately noninsured in the United States,56 may experience language barriers to quality care,7,57 and also have less access to spe- cialized inpatient HF care.12 Native American and Alaskan Native populations experience particu- lar challenges in specialty care access because Indian Health Service facilities are often small and rural.11 Engaging patients in medical care within culturally tailored environments has proven suc- cessful.58,59 HF written educational materials for patients and caregivers should be delivered at or below the sixth grade reading level.60 Workplace interventions that improve cultural competency and address implicit biases are increasingly avail- able. Many aspects of GDMT have been inade- quately studied by population subgroups, largely as a result of clinical trial underrepresentation.61–65

## Cardio-Oncology

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| **Recommendations for Cardio-Oncology**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. In patients who develop cancer therapy–related cardiomyopathy or HF, a multidisciplinary discussion involving the patient about the risk- benefit ratio of cancer therapy interruption, dis- continuation, or continuation is recommended to improve management.1,2 |
| **2a** | **B-NR** | 2. In asymptomatic patients with cancer therapy– related cardiomyopathy (EF <50%), ARB, ACEi, and beta blockers are reasonable to prevent progression to HF and improve cardiac function.2–4 |

|  |  |  |
| --- | --- | --- |
| **Recommendations for Cardio-Oncology (Continued)** | | |
| **COR** | **LOE** | **Recommendations** |
| **2a** | **B-NR** | 3. In patients with cardiovascular risk factors or known cardiac disease being considered for potentially cardiotoxic anticancer therapies, pretherapy evaluation of cardiac function is rea- sonable to establish baseline cardiac function and guide the choice of cancer therapy.2,5–16 |
| **2a** | **B-NR** | 4. In patients with cardiovascular risk factors or known cardiac disease receiving potentially cardiotoxic anticancer therapies, monitoring of cardiac function is reasonable for the early iden- tification of drug-induced cardiomyopathy.2,4,6,8 |
| **2b** | **B-R** | 5. In patients at risk of cancer therapy–related cardiomyopathy, initiation of beta blockers and ACEi/ARB for the primary prevention of drug-induced cardiomyopathy is of uncertain benefit.17–28 |
| **2b** | **C-LD** | 6. In patients being considered for potentially car- diotoxic therapies, serial measurement of car- diac troponin might be reasonable for further risk stratification.29–32 |

## Synopsis

Advances in cancer therapy and an aging population have led to a growing number of cancer patients with comorbid CVD receiving treatment for cancer.33,34 Car- diovascular complications of cancer therapy, notably cardiomyopathy and HF, can result in significant mor- bidity and interruption of treatment, impacting both short- and long-term survival.35,36 Because drug devel- opment in cancer therapeutics grows at an exponential pace, establishing a unified framework for the man- agement of cancer therapy–related cardiomyopathy— commonly defined as a decrease in LVEF of at least 10% to <50%—is necessary to mitigate the cardio- vascular risks of established novel therapies. Cardio- oncology is the practice of precancer therapy cardio- vascular risk stratification, prevention, early detection, and treatment of cardiovascular complications.36,37 The evidence from which guideline recommendations in cardio-oncology have emerged has been based on studies of anthracycline and trastuzumab-induced car- diomyopathy. Cancer therapy–related cardiomyopathy is, however, a heterogeneous disease, with a wide range of presentations—from asymptomatic LV dys- function to cardiogenic shock—and drug-dependent pathophysiologic mechanisms that are often poorly understood. Until sufficient high-quality, drug-specific evidence and cost-effectiveness analyses for screen- ing and monitoring are available, these recommen- dations are applicable across potentially cardiotoxic therapies (Table 28).

## Recommendation-Specific Supportive Text

* + 1. HF secondary to cancer therapy–related cardio- myopathy is associated with significantly worse

**Table 28. Cancer Therapies Known to Be Associated With Cardiomyopathy**

|  |  |  |  |
| --- | --- | --- | --- |
| **Class** |  | **Cardiac Function Monitoring Often Performed in Clinical Practice** | |
| **Agent(s)** | **Pretherapy** | **Serial** |
| Anthracyclines55–57 | Doxorubicin, epirubicin | X | X |
| Alkylating agents58–60 | Cyclophosphamide, ifosfamide, melphalan | X |  |
| Antimicrotubule agents.61,62 | Docetaxel |  |  |
| Antimetabolites63–72 | Fluorouracil, capecitabine, fludarabine, decitabine |  |  |
| Anti-HER2 agents73–76 | Trastuzumab, pertuzumab | X | X |
| Monoclonal antibodies77 | Rituximab |  |  |
| Tyrosine-kinase inhibitors78–100 | Dabrafenib, dasatinib, lapatinib, pazopanib, ponatinib, sorafenib, trametinib, sunitinib, vandetanib, imatinib, vandetanib |  |  |
| Immune checkpoint inhibitors39,40,101 | Nivolumab, ipilimumab, pembrolizumab |  |  |
| Protease inhibitors102–106 | Bortezomib, carfilzomib |  |  |
| Endocrine therapy107–111 | Goserelin, leuprolide, flutamide, bicalutamide, nilutamide |  |  |
| Chimeric antigen receptor T-cell therapy.112,113 | Tisagenlecleucel, axicabtagene ciloleucel | X |  |
| Hematopoietic stem cell transplantation7,44,114–119 | Hematopoietic stem cell transplantation | X |  |
| Radiation7,44,114–119 | Chest |  |  |

outcomes.1,2,38 Patients who develop HF while receiving potentially cardiotoxic therapies should have these therapies discontinued while a diagnostic workup is undertaken to ascertain the cause of HF and initiate GDMT. The com- plex decision to resume, modify, or permanently discontinue therapy by the primary oncologist should be done in a patient-centered approach in concert with a cardiovascular specialist in cardio-oncology. Factors impacting the decision include the severity of cancer therapy–related cardiomyopathy and the response to neurohor- monal blockade, the offending agent’s specific mechanism of injury, the patient’s comorbid con- ditions and cancer-related prognosis and, lastly, the availability of alternative noncardiotoxic treatment options. However, the clinical signifi- cance of asymptomatic cancer therapy–related cardiomyopathy that is identified on routine monitoring is less clear. This is most apparent in patients receiving trastuzumab in whom asymp- tomatic decreases in LVEF can occur in >10% of patients yet result in a high recovery rate and low rate of discontinuation of therapy.1,2 Accordingly, trastuzumab is often continued in patients deemed low risk while neurohormonal blockade is initiated. Conversely, patients diagnosed with immune checkpoint-related myocarditis typically have the offending agents discontinued indefi- nitely, given the associated high mortality.39,40

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* + 1. Studying the effectiveness of neurohormonal therapies specifically in patients with the *CTRC* gene is challenging given the relative infrequency of events, heterogeneity of offending agents, the poorly understood pathophysiology, and the

overlap with comorbid CVD. Available data in patients with anthracycline and trastuzumab- induced cardiomyopathy suggest beta blockers and ACEi are effective in improving LV dysfunc- tion.2–4 Given the dearth of data specific to can- cer therapy–related cardiomyopathy for other GDMT, their use should align with the HFrEF management guidelines. Initiation and uptitration of standard HF therapies remains the mainstay of treatment in patients with cancer therapy–related cardiomyopathy or LVEF <50%, with close moni- toring of cardiac function to guide discussions with oncology on the resumption of, or choice of, subsequent cancer therapies.2

* + 1. Pretherapy quantification of LVEF in patients receiving potentially cardiotoxic cancer thera- pies serves 4 purposes: 1) pretherapy risk stratification and diagnosis of preexisting car- diomyopathy, 2) establish a reference baseline to which reevaluations can be compared, 3) initi- ate cardioprotective medications before cancer therapy, and 4) guide choice of cancer therapy. Echocardiography is recommended as the first- line modality for LVEF assessment given its availability, safety, relatively low cost, and its abil- ity to provide structural and functional informa- tion beyond LVEF.2,5–16,41–47 The risk of cancer therapy–related cardiomyopathy varies greatly across cancer therapies and is modified by pre- existing cardiovascular risk factors (Table 29). Pretherapy LVEF is a strong predictor of major adverse cardiovascular events in patients receiv- ing potentially cardiotoxic therapies.2,5–10,42–47 The clinical use and cost-effectiveness of systematic screening in all patients, however, is unclear.11–16

##### Table 29. Risk Factors for Cancer Therapy–Related Cardiomyopathy

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|  |
| --- |
| Age 60 y |
| Black race |
| CAD |
| Hypertension |
| Diabetes |
| Preexisting cardiomyopathy |
| Previous exposure to anthracyclines |
| Previous chest radiation |
| Elevated troponin pretherapy |

CAD indicates coronary artery disease.

Patients with cancer and preexisting cardio- vascular risk factors are at significantly higher risk of cancer therapy–related cardiomyopathy, representing a population in which pretherapy evaluation would have a significantly higher yield.2,5–10,42–47

* + 1. The purpose of serial monitoring of LVEF in patients receiving potentially cardiotoxic anti- cancer agents is to identify subclinical cardiac injury, initiate cardioprotective agents, and con- sider temporary or permanent interruption of the offending agent.2,4,6,8,48 The practice of LVEF monitoring has mostly been implemented in patients receiving anthracyclines, trastuzumab, or both (Table 28). In a study of 2625 patients receiving anthracyclines for breast cancer or lymphoma who underwent serial LVEF moni- toring, cancer therapy–related cardiomyopathy occurred in 9% of patients, of whom 81% had mild symptoms (NYHA class I to II).4 Beta block- ers and ACEi-ARB were initiated in all patients, with 86% having at least partial recovery of LVEF.4 Patients with recovered LVEF had a lower incidence of cardiac events than those that did not.4 The clinical significance of an asymp- tomatic decrease in LVEF and the optimal fre- quency and duration of monitoring is less clear and likely depend on patient risk, the anticancer agent used, and its cumulative dose. Although a one-size-fits-all approach to monitoring for cancer therapy–related cardiomyopathy may be easier to implement systematically, it may not be the most cost-effective. Until additional data are available, limiting the monitoring to patients at higher risk of cancer therapy–related cardiomy- opathy (Table 29) is a reasonable strategy.
    2. Whether the preemptive use of ACEi-ARB, spi- ronolactone, or selected beta blockers such as carvedilol and nebivolol is effective in reducing the risk of cancer therapy–related cardiomyopathy

has been investigated in a number of small clinic trials, with conflicting findings.17–27,49 The most supportive of this practice is a study that randomized 114 patients receiving high-dose chemotherapy and having a posttreatment tropo- nin rise >0.07 ng/mL to enalapril or placebo.20 None of the patients in the enalapril arm met the primary endpoint (>10% decrease in LVEF to below 50%), while 43% of patients in the stan- dard of care group had a significant decrease in LVEF.20 Although other studies have shown similar findings, the magnitude of the difference in LVEF between arms was often small (<5%) and of questionable clinical significance.19,22 Not all studies have replicated these findings.18,21,24,26 Most importantly, none of the studies have assessed whether preemptive use of HF thera- pies in patients at risk for cancer therapy–related cardiomyopathy improves clinical outcomes, such as mortality or hospitalization for HF. Additional studies are needed to define the appropriate cri- teria and patient population in whom to initiate medical therapies for the primary prevention of cancer therapy–related cardiomyopathy.

* + 1. Cardiovascular biomarkers, notably troponin, have been studied for cardiovascular risk stratifica- tion in patients undergoing potentially cardiotoxic therapies.29–32 A study of 452 patients with breast cancer showed that an elevated pretreatment level (>14 ng/L) was associated with a 4-fold increase in the risk of cancer therapy–related car- diomyopathy.32 Other smaller studies have found no advantage in measuring troponin or natriuretic peptides pretherapy.50–53 Overall, these biomarker studies were observational and small in sample size and number of events.54 Serial biomarkers may be more useful in risk stratification. For exam- ple, in a study of 703 patients receiving anthra- cyclines, an increase in troponin within 72 hours of chemotherapy and 1 month after the comple- tion of treatment course were associated with a greater risk of cancer therapy–related cardiomy- opathy.29 The clinical use of measuring biomark- ers was assessed in 1 trial in which 114 patients with posttreatment increase in troponin to >0.07 ng/mL were randomized to enalapril or standard of care.20 None of the patients in the enalapril group had a decrease in LVEF, compared with 43% in the standard of care group.20 Data for the use of natriuretic peptides are limited. In practice, biomarkers could provide rapid risk stratification in patients for which echocardiographic findings are equivocal and help determine whether symp- toms are cardiovascular in origin.

## HF and Pregnancy

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| **Recommendations for HF and Pregnancy**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 1. In women with a history of HF or cardiomyopathy, including previous peripartum cardiomyopathy, patient-centered counseling regarding contra- ception and the risks of cardiovascular deteriora- tion during pregnancy should be provided.1–8 |
| **2b** | **C-LD** | 2. In women with acute HF caused by peripartum cardiomyopathy and LVEF <30%, anticoagula- tion may be reasonable at diagnosis, until 6 to 8 weeks postpartum, although the efficacy and safety are uncertain.9–12 |
| **3: Harm** | **C-LD** | 3. In women with HF or cardiomyopathy who are pregnant or currently planning for pregnancy, ACEi, ARB, ARNi, MRA, SGLT2i, ivabradine, and vericiguat should not be administered because of significant risks of fetal harm.13–15 |

**Synopsis**

HF may complicate pregnancy either secondary to an ex- isting prepregnancy cardiomyopathy, or as a result of peri- partum cardiomyopathy.16–18 Peripartum cardiomyopathy is defined as systolic dysfunction, typically LVEF <45%, often with LV dilation, occurring in late pregnancy or early postpar- tum with no other identifiable cardiomyopathy cause.14,19–21 Peripartum cardiomyopathy occurs globally,22,23 with the highest incidences in Nigeria, Haiti, and South Africa. Inci- dence in the United States is 1 in 1000 to 8000 deliver- ies and has risen over time.24,25 Peripartum cardiomyopathy risk factors include maternal age >30 years, African an- cestry, multiparity, multigestation, preeclampsia/eclampsia, anemia, diabetes, obesity, and prolonged tocolysis.22,23,26–30 A genetic contribution is recognized,31–33 particularly titan gene mutations.34,35 Most women present with HF within 1 month postpartum; cardiogenic shock, arrhythmias, or venous-arterial thromboembolism are all possible. Treat- ment includes GDMT adjusted for pregnancy or breast- feeding status and anticoagulation consideration16; iden- tification of a pathogenic 16-kDa prolactin led to trials of the dopamine-agonist bromocriptine.36–41 Patient-centered multidisciplinary planning is essential, including early insti- tution of mechanical support for shock42 (Table 30). Prog- nosis is related to initial LVEF, LV thrombosis, RV involve- ment, preeclampsia, geographic region, and race.7,43–48 LV recovery and survival is generally favorable in developed countries11,25,49; a 100-patient US registry showed 93% transplant/LVAD-free 1-year survival.46

## Recommendation-Specific Supportive Text

* + 1. Pregnancy is generally well-tolerated in women with cardiomyopathy and NYHA class I prepregnancy. However, clinical deterioration can occur, so pre- pregnancy counseling and shared decision-making

are essential.1,3,50 Among women with non–peri- partum cardiomyopathy, major cardiovascular events occurred in 39% (United States) and 35% (Canada) of pregnancies, with 1% and 7% mortal- ity, respectively.51,52 Previous cardiac events, NYHA class III to IV, or LVEF <40% markedly increased maternal and fetal risks.51–55 The ROPAC (Registry of Pregnancy and Cardiac disease) study describes pregnancy outcomes for 1321 women with struc- tural heart disease: Women with prepregnancy or peripartum cardiomyopathy had the highest mortal- ity rate (2.4%).2,22 ROPAC was used to validate the modified WHO risk classification56; the ZAHARA I (Zwangerschap bij Aangeboren Hartafwijkingen I) and CARPREG II (CARdiac disease in PREGnancy) scores also support shared decision-making.51,57,58 Subsequent pregnancies for women with previous peripartum cardiomyopathy have been associated with further decreases in LV function, maternal death, and adverse fetal outcomes.43,58 The stron- gest prognostic determinant is LVEF <50% before a subsequent pregnancy.6–8 An international sys- tematic review that included 93 subsequent preg- nancies with persistent LV dysfunction reported 48% further LVEF deterioration, 49% HF symp- toms, and 16% mortality, whereas among 98 with recovered LV function presubsequent pregnancy, these rates were 27%, 32%, and 0%, respectively.5

* + 1. Pregnancy is a hypercoagulable state even in the absence of cardiovascular complications. In the set- ting of acute HF, particularly when there is LV blood stasis because of severely reduced systolic function, the risk of intracardiac thrombus formation is sig- nificant. The incidence of intracardiac thrombi dur- ing acute HF caused by peripartum cardiomyopathy has been reported to be around 16% to 17%.9,10 with 9% thromboembolic events in 2 separate cohorts11,12 Women with an intracardiac thrombus or a thromboembolic event receive anticoagula- tion as per standard of care. Women with severely depressed LVEF (<30%) in the setting of acute HF caused by peripartum cardiomyopathy can be considered for anticoagulation, especially in the first 6 to 8 weeks postpartum, when hypercoagulabil- ity is most pronounced. If bromocriptine is used for postpartum women with severe acute HF caused by peripartum cardiomyopathy and LVEF <35%, it should be accompanied by at least prophylactic- dosed anticoagulation, because of the potential association with thromboembolic events.6 However, the efficacy and safety of bromocriptine for acute peripartum cardiomyopathy treatment currently remains uncertain, and further randomized placebo- controlled trials are required to define the role of this therapy, particularly in the setting of contemporary HF GDMT and cardiogenic shock management.

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##### Table 30. HF Management Strategies Across the Pregnancy Continuum

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| --- | --- | --- | --- |
|  | **Preconception** | **During Pregnancy** | **Postpartum** |
| Nonpharmacological strategies | Preconception genetic counseling and testing for potentially heritable cardiac conditions.  Use of pregnancy cardiovascular risk tools,51,56–58 and echocardiography for myo- cardial structure and function assessment, to provide information that facilitates informed counseling.  For women planning a pregnancy, provide per- sonalized counseling that promotes the auton- omy and goals of the patient (and her partner, as applicable), the patient’s ability for self-care and risk awareness, and ensures adequate psychosocial support for decision-making.3  For women not currently planning a pregnancy but who might conceive, discuss HF-specific considerations regarding pregnancy and refer to gynecology or primary care for contracep- tive counseling. | Close maternal monitoring for HF signs or symp- toms or other cardiovascular instability by cardiol- ogy and obstetric and maternal-fetal medicine teams; close fetal monitoring by the obstetric and maternal-fetal medicine teams.  Consideration of routine echocardiographic screening in the third trimester for reassess- ment of myocardial structure and function before labor; echocardiography for any signifi- cant changes in HF symptoms or signs during pregnancy, or if HF medications are reduced or discontinued.18  BNP or NT-proBNP monitoring during preg- nancy may have some value for prediction of cardiovascular events.73,74  Close maternal monitoring by obstetrics and maternal-fetal medicine teams for preeclamp- sia, which has shared risk factors and patho- genesis with PPCM.47,75  For women presenting with decompensated HF or cardiogenic shock, hemodynamic moni- toring and MCS, as appropriate, within a mul- tidisciplinary collaborative approach that sup- ports prompt decision-making about the timing and mechanism of delivery. | Multidisciplinary recommendations from ob- stetrics and neonatology and pediatrics teams and shared decision-making regarding the maternal and neonatal risks and benefits of breastfeeding.  For women presenting with decompensated HF or cardiogenic shock, HF management should include hemodynamic monitoring and mechanical circulatory support as appropriate |
| Pharmacological strat- egies | Review of all current medications.  For women planning pregnancy imminently, modification of HF pharmacotherapy includ- ing. discontinuation of any ACEi, ARB, ARNi, MRA, or SGLT2i or ivabradine medications; within a construct of multidisciplinary shared decision-making, continuation of a beta block- er (most commonly metoprolol), hydralazine, and nitrates; adjustment of diuretic dosing  to minimize the risk of placental hypoperfu- sion.13–15  Ideally, repeat echocardiography approximately 3 mo after preconception HF medication adjustments to ensure stability of myocardial structure and function before conception. | Close monitoring of maternal blood pressure, heart rate, and volume status, with adjustment of the modified HF regimen as appropriate  to avoid hypotension (systemic vasodilation peaks in the second trimester) and placental hypoperfusion.  For women with HF or cardiomyopathy presenting during pregnancy without precon- ception counseling and assessment, urgent discontinuation of any GDMT pharmacothera- pies with fetal toxicities; within a construct  of multidisciplinary shared decision-making, continuation of a beta blocker (most com- monly metoprolol succinate), hydralazine, and nitrates; adjustment of diuretic dosing to mini- mize the risk of placental hypoperfusion. | For women with acute HF caused by PPCM and LVEF <30%, consideration of anticoagula- tion until 6–8 wk postpartum, although the ef- ficacy and safety remain uncertain at this time.  For postpartum women with severe acute HF caused by PPCM and LVEF <35%, in GDMT pharmacotherapy and prophylactic anticoagula- tion, to improve LVEF recovery6,31,3641,76; the efficacy and safety of bromocriptine for acute PPCM treatment remains uncertain at this time, particularly in the setting of contemporary HF GDMT and cardiogenic shock management.\*  For women who choose to breastfeed, review medications with neonatology and pediatrics teams for neonatal safety during lactation, ideally with pharmacist consultation if available.  Within a construct of multidisciplinary shared decision-making, medications that may be ap- propriate during breastfeeding include ACEi (enalapril or captopril preferred, monitor neona- tal weight), beta blockers (metoprolol preferred, monitor neonatal heart rate).15  Diuretics can suppress lactation, but with neo- natal follow-up the use of furosemide may be appropriate.15 |
| Multidisciplinary care beyond the cardiology team | Consultation with genetics, gynecology, and maternal-fetal medicine teams, as appropriate to the outcome of shared decision-making. | Multidisciplinary management with obstetrics and maternal-fetal medicine teams during pregnancy.  For women with decompensated HF or evi- dence of hemodynamic instability antepartum, delivery planning will include obstetrics and maternal-fetal medicine, anesthesia, and neo- natology teams. | Multidisciplinary management with obstetrics, maternal-fetal medicine, neonatology, and pe- diatrics teams, especially for multidisciplinary recommendations regarding lactation.  Consultation with gynecology team for ongoing contraceptive planning. |

ACEi indicates angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; ARNi, angiotensin receptor-neprilysin inhibitor; BNP, B-natriuretic peptide; GDMT, guideline-directed medical therapy; HF, heart failure; LVEF, left ventricular ejection fraction; MCS, mechanical circulatory support; MRA, mineralocorticoid receptor antagonist; NT-proBNP, N-terminal prohormone of brain natriuretic peptide; PPCM, peripartum cardiomyopathy; RCT, randomized controlled trial; RV, right ventricular; and SGLT2i, sodium- glucose cotransporter-2 inhibitor.

\*An initial open-label pilot RCT in South Africa suggested addition of bromocriptine to GDMT was associated with greater LVEF improvement and a lower rate of the composite endpoint at 6 mo.37 Among 96 women with acute PPCM in a Burkina Faso RCT, 4 wk of bromocriptine was associated with LVEF recovery and lower mortality (16.6% versus 29.1%; *P*<0.001).39 A multicenter German study randomized 63 patients to 1 versus 8 wk of bromocriptine (no placebo, as deemed unethical),38 with LVEF recovery 50% in 52% and 68% of the 1- and 8-wk groups, respectively, and no deaths. A substudy also showed high rates of RV recovery.41 Two retrospective cohorts (Germany, Canada) and a multicenter cohort of subsequent pregnancies also suggested greater LVEF recovery with bromocriptine.31,38,40 Bromocriptine may currently be most justified in women with LVEF

<25% or cardiogenic shock. The downsides of prohibiting breastfeeding should be considered. Bromocriptine should be accompanied by at least prophylactic-dosed anticoagula- tion, because of potential hypercoagulability.38 The European Society of Cardiology endorses “BOARD” (Bromocriptine, Oral HF therapy, Anticoagulation, vasoRelaxing agents, Diuretics) for acute PPCM management.13,14

* + 1. In 2015, the FDA adopted the Pregnancy and Lactation Labeling Rule, which retired the previous pregnancy risk categories A through X and, instead, assigned a descriptive risk summary to aid medication counseling for pregnant and breastfeeding women. ACEi and ARB are associated with second- and third- trimester renal and tubular dysplasia, oligohydramnios, fetal growth restriction, ossification disorders of the skull, lung hypoplasia, contractures, large joints, ane- mia, and intrauterine fetal death and are, therefore, strictly contraindicated.59–61 There are no specific data for ARNi or ivabradine. For spironolactone, there is sufficient information regarding dose-dependent feminization of male rabbit and rat offspring to raise concern62; data are limited for eplerenone. HFrEF medications considered acceptable during preg- nancy,15 within a construct of multidisciplinary shared decision-making regarding benefits and potential risks, are furosemide, beta blockers (most com- monly metoprolol),63–65 hydralazine, and nitrates.13,14,19 Women with peripartum cardiomyopathy were his- torically counseled against breastfeeding because of metabolic demands and prolactin stimulation, but breastfeeding may even be associated with LV recov- ery.66–70 Postpartum women who breastfeed can start ACEi (enalapril or captopril preferred), and metoprolol remains the preferred beta blocker.66,71 The National Library of Medicine hosts LactMed [(https://www.ncbi.](https://www.ncbi.nlm.nih.gov/books/NBK501922/) [nlm.nih.gov/books/NBK501922/).](https://www.ncbi.nlm.nih.gov/books/NBK501922/)72

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# QUALITY METRICS AND REPORTING

## Performance Measurement

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| **Recommendations for Performance Measurement**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **B-NR** | 1. Performance measures based on professionally developed clinical practice guidelines should be used with the goal of improving quality of care for patients with HF.1–7 |
| **2a** | **B-NR** | 2. Participation in quality improvement programs, including patient registries that provide bench- mark feedback on nationally endorsed, clinical practice guideline–based quality and perfor- mance measures can be beneficial in improving the quality of care for patients with HF.1,2,5,6 |

**Synopsis**

The ACC/AHA Task Force on Performance Measures (Task Force) distinguishes quality measures from perfor- mance measures. Performance measures are selected from the most important ACC/AHA clinical practice guideline recommendations with the strongest evidence. These measures are suitable for public reporting or pay for performance. Quality measures are those metrics that may be useful for local quality improvement but do not

reach the performance measure standard. Performance measures of the ACC/AHA focus on process of care measures that measure the quality of care by the clini- cian, facility, and health system. Patient registries that track such measures can provide feedback to partici- pants, which may help with improvement in quality.

## Recommendation-Specific Supportive Text

* + 1. The current ACC/AHA performance and qual- ity measures (based on the 2013 ACC/AHA HF guideline and the 2017 ACC/AHA/HFSA guideline supplement) are displayed in Table 31.8 The perfor- mance measures are derived from the most defini- tive guideline recommendations (ie, NYHA class I and class III recommendations). Observational data suggest that hospitals that receive feedback on their HF care improve over time.1–7
    2. Hospitals that perform well on medication-related performance measures have better HF mortality rates than hospitals with poorer performance.3,4 Other observational data suggest that hospitals that par- ticipate in registries have better process of care and outcomes compared with hospitals that do not partic- ipate.5,6 Randomized studies of audit and feedback of performance, in many different patient groups, have, in general, showed improvement in care.7 However, public reporting of HF measures in Ontario, Canada, did not clearly improve care during a randomized trial.9

# GOALS OF CARE

## Palliative and Supportive Care, Shared Decision-Making, and End-of-Life

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| **Recommendations for Palliative and Supportive Care, Shared Decision- Making, and End-of-Life**  **Referenced studies that support the recommendations are summa- rized in the** [**Online Data Supplements**](https://www.ahajournals.org/doi/suppl/10.1161/CIR.0000000000001063)**.** | | |
| **COR** | **LOE** | **Recommendations** |
| **1** | **C-LD** | 1. For all patients with HF, palliative and supportive care—including high-quality communication, conveyance of prognosis, clarifying goals of care, shared decision-making, symptom man- agement, and caregiver support—should be provided to improve QOL and relieve suffering.1 |
| **1** | **C-LD** | 2. For patients with HF being considered for, or treated with, life-extending therapies, the option for discontinuation should be anticipated and dis-  cussed through the continuum of care, including at the time of initiation, and reassessed with changing medical conditions and shifting goals of care.2,3 |
| **2a** | **B-R** | 3. For patients with HF—particularly stage D HF patients being evaluated for advanced  therapies, patients requiring inotropic support or temporary mechanical support, patients experiencing uncontrolled symptoms, major medical decisions, or multimorbidity, frailty, and cognitive impairment—specialist palliative care consultation can be useful to improve QOL and relieve suffering.4–6 |

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##### Table 31. ACC/AHA 2020 HF Clinical Performance, Quality, and Structural Measures8

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| --- | --- | --- | --- | --- |
| **Measure No.** | **Measure Title** | **Care Setting** | **Attribution** | **Measure Domain** |
| PM-1 | LVEF assessment | Outpatient | Individual practitioner Facility | Diagnostic |
| PM-2 | Symptom and activity assessment | Outpatient | Individual practitioner Facility | Monitoring |
| PM-3 | Symptom management | Outpatient | Individual practitioner Facility | Treatment |
| PM-4 | Beta-blocker therapy for HFrEF | Outpatient Inpatient | Individual practitioner Facility | Treatment |
| PM-5 | ACEi, ARB, or ARNi therapy for HFrEF | Outpatient Inpatient | Individual practitioner Facility | Treatment |
| PM-6 | ARNi therapy for HFrEF | Outpatient Inpatient | Individual practitioner Facility | Treatment |
| PM-7 | Dose of beta blocker therapy for HFrEF | Outpatient | Individual practitioner Facility | Treatment |
| PM-8 | Dose of ACEi, ARB, or ARNi therapy for HFrEF | Outpatient | Individual practitioner Facility | Treatment |
| PM-9 | MRA therapy for HFrEF | Outpatient Inpatient | Individual practitioner Facility | Treatment |
| PM-10 | Laboratory monitoring in new MRA therapy | Outpatient Inpatient | Individual practitioner Facility | Monitoring |
| PM-11 | Hydralazine and isosorbide dinitrate therapy for HFrEF in those patients self-identified as Black or African American | Outpatient Inpatient | Individual practitioner Facility | Treatment |
| PM-12 | Counseling regarding ICD placement for patients with HFrEF on GDMT | Outpatient | Individual practitioner Facility | Treatment |
| PM-13 | CRT implantation for patients with HFrEF on GDMT | Outpatient | Individual practitioner Facility | Treatment |
| QM-1 | Patient self-care education | Outpatient | Individual practitioner Facility | Self-care |
| QM-2 | Measurement of patient-reported outcome-health status | Outpatient | Individual practitioner Facility | Monitoring |
| QM-3 | Sustained or improved health status in HF | Outpatient | Individual practitioner Facility | Outcome |
| QM-4 | Post-discharge appointment for patients with HF | Inpatient | Individual practitioner, facility | Treatment |
| SM-1 | HF registry participation | Outpatient Inpatient | Facility | Structure |
| Rehabilitation PMs Related to HF (From the 2018 ACC/AHA performance measures for cardiac rehabilitation10 | | | | |
| Rehab PM-2 | Exercise training referral for HF from inpatient setting | Inpatient | Facility | Process |
| Rehab PM-4 | Exercise training referral for HF from outpatient setting | Outpatient | Individual practitioner Facility | Process |

ACEi indicates angiotensin-converting enzyme inhibitor; ACC, American College of Cardiology; AHA, American Heart Association; ARB, angiotensin receptor blocker; ARNi, angiotensin receptor-neprilysin inhibitor; CRT, cardiac resynchronization therapy; GDMT, guideline-directed medical therapy; HF, heart failure; HFrEF, heart failure with reduced ejection fraction; ICD, implantable cardioverter-defibrillator; LVEF, left ventricular ejection fraction; MRA, mineralocorticoid receptor antago- nist; PM, performance measure; QM, quality measure; and SM, structural measure.

## Synopsis

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| **Recommendations for Palliative and Supportive Care, Shared Decision- Making, and End-of-Life (Continued)** | | |
| **COR** | **LOE** | **Recommendations** |
| **2a** | **C-LD** | 4. For patients with HF, execution of advance care directives can be useful to improve docu- mentation of treatment preferences, delivery of patient-centered care, and dying in preferred place.7 |
| **2a** | **C-LD** | 5. In patients with advanced HF with expected survival <6 months, timely referral to hospice can be useful to improve QOL.8 |

Palliative care—defined as patient- and family-cen- tered care that optimizes health-related QOL by an- ticipating, preventing, and treating suffering—should be integrated into the care of all patients with HF.9 Palliative care includes high-quality communication, estimation of prognosis, anticipatory guidance, ad- dressing uncertainty; shared decision-making about medically reasonable treatment options; advance care planning; attention to physical, emotional, spiritual, and

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**Figure 15. A Depiction of the Clinical Course of HF With Associated Types and Intensities of Available Therapies Over Time.12** CHF indicates congestive heart failure; HF, heart failure; and MCS, mechanical circulatory support. Adapted with permission of the American Thoracic Society. Copyright © 2021 American Thoracic Society. All rights reserved. The American Journal of Respiratory and Critical Care Medicine is an official journal of the American Thoracic Society.13 Readers are encouraged to read the entire article for the correct context at <https://www.atsjournals.org/doi/abs/10.1164/rccm.200605-587ST>. The authors, editors, and The American Thoracic Society are not responsible for errors or omissions in adaptations. Adapted with permission from the World Health Organization.14 Copyright 1990 World Health Organization.

psychological distress; relief of suffering; and inclu- sion of family caregivers in patient care and attention to their needs during bereavement.10 Other support- ive needs include home and case management assis- tance, transportation, and care coordination.11 Pallia- tive and supportive care has a role across the stages of HF, starting early in the course of illness, intensify- ing in end-stage disease, and extending into caregiver bereavement (Figure 15).12 Many palliative care needs can and should be addressed by the patient’s interdis- ciplinary care team (primary palliative care), including clarifying their core values, health outcome goals, and therapeutic preferences.1 Specialty palliative care cli- nicians (secondary palliative care) may be consulted to collaboratively care for patients and their families with more challenging needs.7 Barriers to the receipt of palliative care include reluctance of health care pro- fessionals to address death and dying and a propen- sity for patients and caregivers to equate palliation and hospice as hastening death.15

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## Recommendation-Specific Supportive Text

* + 1. Palliative and supportive approaches to the care of patients with HF is inherent to their overall care and should be incorporated throughout the course of ill- ness by all health care professionals.9 The applica- tion of the principles embraced have been shown to improve various processes of care and patient outcomes (Table 32). Palliative and supportive care discussions do not imply that a formal palliative care consultation is needed for each patient but that team members should integrate palliative and sup- portive considerations into routine care.
    2. As overall illness progresses, major decisions are increasingly made regarding the initiation, con- tinued use, and discontinuation of potentially life-sustaining therapies, including intravenous ino- tropes, ICDs, MCS, and renal replacement therapy. Dependence on, and deactivation of, potentially life-sustaining therapies should be anticipated and

##### Table 32. Palliative and Supportive Care Domains to Improve Processes of Care and Patient Outcomes

|  |  |
| --- | --- |
| **Palliative and Supportive Domains of Care** | **What Palliative Care Adds to Overall HF Management** |
| High-quality communication | Central to palliative care approaches are communication and patient-caregiver engagement techniques.16 |
| Conveyance of prognosis | Palliative care specifically addresses patient and caregiver understanding of disease, treatment, and prognosis. Research suggests that patients tend to overestimate their survival17 and overestimate the potential benefits of treatment.18 Objective risk models can calibrate expectations, but discussion of uncertainty should accompany prognostic conversations, often sum- marized as “hope for the best, plan for the worst.” |
| Clarifying goals of care | Management of patients with HF as their disease becomes end-stage and death seems near includes decisions about when to discontinue treatments designed primarily to prolong life (eg, ICD, hospitalization, tube feeding), decisions on when to initiate treatments to reduce pain and suffering that may hasten death (eg, narcotics), and decisions about the location of death, home services, and hospice care. Exploring patients’ expressed preferences, values, needs, concerns, means and desires through clinician-led discussion can clarify values-treatment concordance and improve medical deci- sion-making.12 |
| Shared decision-making | Shared decision-making is a process by which patients and clinicians work together to make optimal health care decisions from medically reasonable options that align with what matters most to patients. Shared decision-making re- quires: unbiased medical evidence about the risks, benefits, and burdens of each alternative, including no intervention; clinician expertise in communication and tailoring that evidence for individual patients; and patient goals and informed preferences.12 |
| Symptom management | Dyspnea, fatigue, pain, nausea, depression, anxiety, and other symptoms of HF refractory to cardiovascular therapies can be partially remediated through palliative and supportive approaches in addition to GDMT.5 |
| Caregiver support | Care of the patient with heart failure should extend to their loved ones, including beyond their death, to offer support to fami- lies and help them cope with loss. |

GDMT indicates guideline-directed medical therapy; HF, heart failure; and ICD, implantable cardioverter-defibrillator.

discussed at the time of initiation and reconsidered serially with changing medical realities and evolv- ing goals of care.12 Patients have a right to decline or withdraw care at any time, consistent with the principle of respect for autonomy.19 Failure to pro- actively address topics such as deactivation of ICD and LVAD therapies can lead to suffering at the end of life.2,3

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* + 1. Although a range of clinicians caring for patients with HF are able to manage many palliative care needs, formal palliative care consultation may be particularly helpful for patients with these: 1) refractory symptoms; 2) major medical decisions (eg, in the United States, inclusion of a palliative care specialist on the team is mandatory for pay- ment from Medicare for LVAD implantation); and 3) multimorbidity, frailty, or cognitive impairment (mul- tiple validated frailty and cognitive measures are available). A growing body of evidence supports the inclusion of specialty palliative care into the management of patients diagnosed with a range of advanced diseases,20 including HF. An interdis- ciplinary palliative care intervention in patients with advanced HF showed greater benefits in QOL, anxiety, depression, and spiritual well-being com- pared with usual care alone (PAL-HF [Palliative Care in Heart Failure]).4 However, other trials have been mixed,5,6 and many negative,21–23 such that formal palliative care interventions should be tai- lored to patient and caregiver wants and needs.
    2. Advance care planning is a process that sup- ports understanding and sharing of patients’ per- sonal values, life goals, and preferences regarding future medical care. Key domains include discuss- ing patients’ values, documenting plans for medi- cal treatments, designating a surrogate decision maker, and revisiting this process over time.24 Familiarity with local and state laws is needed relat- ing to advance care planning, decisions regarding life-sustaining treatments, and evolving treatments with legal ramifications, especially when caring for vulnerable populations.19 Few patients with HF have formally defined their care goals and desig- nated a surrogate decision maker.25
    3. Hospice is a specific model of subspecialty pallia- tive care that is offered to patients with a terminal disease who are at the end of life when curative or life-prolonging therapy is no longer the focus of treatment.10 Historically, hospice use has been low among patients dying with HF and, among those engaging in hospice, the duration of time in hospice was short, suggesting late referral. Low hospice referral rates and high-intensity care at end of life often reflects health care professional biases and limitations in models of care rather than patient values.26 This appears to be changing in the United States, where CDC data from 2003 to 2017 on US site of death show that the proportion of cardiovascular deaths related to HF occurring in hospice facilities rose from 0.2% to 8.2% and deaths at home rose from 20.6% to 30.7%.27

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##### Table 33. Evidence Gaps and Future Research Directions

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| Definition |
| Consensus on specific classifications of HFrEF, HFpEF, HFmrEF, and HFimpEF or whether a 2-category definition of HFrEF and HF with normal EF, or an addi- tional category of HFimpEF is needed separately for HFpEF; and whether these approaches can be uniformly applied to clinical trials and practice. |
| Definitions, detection, and management of myocarditis and myocardial injury, especially in the context of rapidly evolving concepts, such as COVID-19 infection and cardiotoxicity. |
| Definition and classification of cardiomyopathies. |
| Screening |
| Cost-effectiveness of different strategies to screen for HF. |
| Prediction of higher risk for HF among patients with traditional risk factors (eg, which patients with diabetes would be at a higher risk HF, warranting preventive treatment for HF). |
| Diagnostics and monitoring |
| Individualized treatment targeting specific causes. |
| Advanced role of precision medicine with incorporation of genetic, personalized, and individualized factors in medical management of HF. |
| High-value methods to use biomarkers in the optimization of medical therapy. |
| Ability to use integrated systems biology models, including biomarkers, molecular markers, omics, diagnostic modalities, and genetic variables for diagnosis, prognosis, and targeting therapies. |
| Ability to monitor and adjust therapy to individual changes over time. |
| Nonmedical strategies |
| Efficacy and safety of specific dietary interventions, sodium restriction, and fluid restriction to prevent and treat HF. |
| Efficacy and safety of cardiac rehabilitation in patients with HFpEF and HFmrEF. |
| Medical therapies |
| Effective management strategies for patients with HFpEF. |
| Evidence for specific treatment strategies for HFmrEF. |
| Research on causes and targeted therapies for cardiomyopathies such as peripartum cardiomyopathy. |
| Treatment of asymptomatic LV dysfunction to prevent transition to symptomatic HF. |
| Therapies targeting different phenotypes of HF; patients with advanced HF, persistent congestion, patients with profiles excluded from clinical trials such as those with advanced kidney failure or hypotension. |
| Studies on targets for optimal decongestion; treatment and prevention of cardiorenal syndrome and diuretic resistance. |
| Diagnostic and management strategies of RV failure. |
| Efficacy and safety of hydralazine isosorbide in non–African American patients with HF and also in African American patients on GDMT including SGLT2i and ARNi. |
| Efficacy and safety of vericiguat in patients with HFrEF and markedly elevated natriuretic peptide levels. |
| Efficacy and safety of omecamtiv mecarbil in patients with stage D (advanced HF) HFrEF. |
| Additional efficacy and safety of SGLT2i therapies in patients with HFpEF or patients with HFmrEF, efficacy and safety of combined SGLT2i and SGLT1i in HFrEF, HFmrEF, or HFpEF. |
| Additional efficacy and safety of SGLT2i studies in hospitalized patients with acute decompensated HF with and without diabetes. |
| Efficacy and safety of nonsteroidal, selective MRA in patients with HF. |
| Efficacy and safety of ARNi in pre-HF stage (stage B). |
| Effective management strategies for combined post- and precapillary pulmonary hypertension. |
| Novel treatments for ATTR cardiomyopathy. |
| Treatment strategies targeting downstream processes such as fibrosis, cardiac metabolism or contractile performance in dilated cardiomyopathies and HFpEF. |
| Comparative effectiveness and safety of different initiation and titration of GDMT at the same time or in different sequences, optimal strategies for sequencing and titration of therapies for HFrEF and HFpEF. |
| Studies on prediction of patient response; studies on how to incorporate patient preferences. |
| Efficacy and safety of optimal BP target in patients with established HF and hypertension. |
| Optimal BP target while optimizing GDMT in patients with HFrEF and HFpEF. |
| Appropriate management of electrolyte abnormalities in HF (eg, hyperkalemia or hypokalemia). |
| Role of potassium binders in optimization of GDMT and clinical outcomes in patients with HF. |
| Efficacy and safety of pirfenidone and other targeted treatment strategies for maladaptive fibrosis in patients with HFpEF. |
| AF risk in patients treated with PUFA for patients at risk for HF or with HF. |

(*Continued* )

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##### Table 33. Continued

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| Device management and advanced therapies |
| Optimal and timely selection of candidates for percutaneous interventions, MCS, or cardiac transplantation. |
| Interventional approaches to recurrent, life-threatening ventricular tachyarrhythmias. |
| Comparative effectiveness of His-bundle pacing or multisite pacing to prevent progression of HF. |
| Safety and efficacy of cardiac contractility modulation, vagal nerve stimulation, autonomic modulation, and renal denervation in patients with HF. |
| Safety and efficacy of splanchnic nerve ablation splanchnic nerve ablation to reduce splanchnic vasoconstriction and volume redistribution in HF. |
| Safety and efficacy of interatrial shunt, pericardiectomy, baroreceptor and neuromodulation, and renal denervation in HFpEF. |
| Safety and efficacy of percutaneous or surgical interventions for tricuspid regurgitation. |
| Clinical outcomes |
| Impact of therapies in patient-reported outcomes, including symptoms and QOL. |
| Studies addressing patient goals about care and care intensity as it intersects with disease trajectory. |
| Real-world evidence data to characterize generalization of therapies in HF populations who may not have been represented in trials. |
| Systems of care and social determinants of health |
| Implementation studies on how to develop a structured approach to patient participation in informed decision-making and goal setting through the continuum of HF care. |
| Implementation science for adoption and optimization of GDMT by clinicians on how to initiate multiple or sequenced GDMT, how to integrate these into learning health systems and networks, and how to increase patient education and adherence. |
| Pragmatic studies on multidisciplinary new care models (eg, cardiac teams for structural and valve management, shock teams, cardiometabolic clinics, telemedi- cine, digital health, cardiac rehabilitation at home or postdischarge, and palliative care). |
| Studies on strategies to eliminate structural racism, disparities, and health inequities in HF care. |
| Studies addressing evidence gaps in women, racial, and ethnic populations. |
| Management strategies for palliative care. |
| Identification of factors that lead to unwarranted variations in HF care. |
| Identify characteristics of systems of care (eg, disciplines and staffing, electronic health records, and models of care) that optimize GDMT before and after the discharge of hospitalized patients. |
| Comorbidities |
| Further studies on rhythm control versus ablation in AF. |
| Appropriate patient selection in evolving percutaneous approaches in VHD (eg, timing and appropriate patient selection for TAVI, Mitraclip, tricuspid valve inter- ventions). |
| Effective and safe treatment options in CKD, sleep-disordered breathing, chronic lung disease, diabetes, depression, cognitive disorders, and iron deficiency. |
| Efficacy and safety of transvenous stimulation of the phrenic nerve or role of nocturnal supplemental oxygen for treatment of central sleep apnea in patients with HF. |
| Efficacy and safety of weight loss management and treatment strategies in patients with HF and obesity. |
| Efficacy and safety of nutritional and food supplementation in patients with HF and frailty and malnutrition. |
| Efficacy and safety of GDMT in end-stage renal disease or in patients with eGFR <30 mL/min/1.73 m2. |
| Future/novel strategies |
| Pharmacological therapies targeting novel pathways and endophenotypes. |
| New device therapies, including percutaneous and durable mechanical support devices. |
| Invasive (eg, pulmonary artery pressure monitoring catheter) or noninvasive remote monitoring. |
| Studies on telehealth, digital health, apps, wearables technology, and artificial intelligence. |
| Role of enrichment trials, adaptive trials, umbrella trials, basket trials, and machine learning–based trials. |
| Therapies targeting multiple cardiovascular, cardiometabolic, renovascular, and pathobiological mechanisms. |
| Novel dissemination and implementation techniques to identify patients with HF (eg, natural language processing of electronic health records and automated analysis of cardiac imaging data) and to test and monitor proven interventions. |

AF indicates atrial fibrillation; ARNi, angiotensin receptor-neprilysin inhibitor; ATTR, transthyretin amyloidosis; BP, blood pressure; CKD, chronic kidney disease; COVID-19, coronavirus disease 2019; eGFR, estimated glomerular filtration rate; GDMT, guideline-directed medical therapy; HF, heart failure; HFimpEF, heart failure with improved ejection fraction; HFmrEF, heart failure with mildly reduced ejection fraction; HFpEF, heart failure with preserved ejection fraction; HFrEF, heart failure with reduced ejection fraction; LV, left ventricular; MCS, mechanical circulatory support; MRA, mineralocorticoid receptor antagonist; PUFA, polyunsaturated fatty acid; QOL, quality of life; RV, right ventricular; SGLT1i, sodium-glucose cotransporter-1 inhibitors; SGLT2i, sodium-glucose cotransporter-2 inhibitors; TAVI, transcath- eter aortic valve implantation; and VHD, valvular heart disease.

# RECOMMENDATION FOR PATIENT- REPORTED OUTCOMES AND EVIDENCE GAPS AND FUTURE RESEARCH DIRECTIONS

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## Patient-Reported Outcomes

|  |  |  |
| --- | --- | --- |
| **Recommendation for Patient-Reported Outcomes** | | |
| **COR** | **LOE** | **Recommendation** |
| **2a** | **C-LD** | 1. In patients with HF, standardized assessment of patient-reported health status using a vali- dated questionnaire can be useful to provide incremental information for patient functional status, symptom burden, and prognosis.1–19 |

**Synopsis**

Health status encapsulates symptoms, functional status, and health-related QOL. Understanding health status is important for treatment decisions and counseling. Clini- cians traditionally evaluate health status based on the clinical interview and exam, summarizing it as the NYHA functional classification. Additionally, patient-reported health status can be ascertained using standardized questionnaires, such as the Kansas City Cardiomyopathy Questionnaire or the Minnesota Living with Heart Fail- ure Questionnaire. Previous studies found discordance between patient-reported health status and clinician as- sessment using NYHA classification.20,21 Patient-reported health status may have higher reliability and better sensi- tivity for clinical changes than NYHA classification and is moderately correlated with CPET and the 6-minute walk test.1–8 Patient-reported health status is an independent predictor of hospitalization and mortality.9–19 There are minimal data regarding the effect of incorporating patient- reported health status assessment into routine care. How- ever, these assessments provide valuable incremental information beyond the standard evaluation. Increasing the patient’s voice in clinical assessment and decision- making is important in its own right. Additionally, there is substantial variation in risk-adjusted health status across practices.22 Future efforts should focus on expanding the use of patient-reported health status in routine care while researching its implementation and impact.

## Recommendation-Specific Supportive Text

* + 1. Standardized patient-reported health status ques- tionnaires provide reliable measures of health status correlated to other functional status mea- sures1–8 and independently associated with clinical outcomes.9–19 HF-specific health status assessments (eg, Kansas City Cardiomyopathy Questionnaire, Minnesota Living with Heart Failure Questionnaire, PROMIS-Plus-HF [Patient- Reported Outcomes Measurement Information System-Plus-Heart Failure]) are preferable because they are more sensitive to changes in

disease status and more responsive to HF therapy than generic health status measures.1 Although select clinics have successfully implemented patient-reported health status in clinical practice,23 there are minimal data regarding the impact of such efforts. However, there are potential advantages to routine assessment. First, better understanding of symptom burden and prognosis may improve the quality of treatment decisions and, subsequently, QOL. Health status can be improved via guide- line-recommended therapies.24–31 Although some therapies are recommended for mortality benefit, symptom assessment can identify patients need- ing additional interventions (eg, diuretic escala- tion). Second, routine assessment can facilitate population health management by identifying high- risk patients needing closer monitoring or refer- ral to specialized centers. Third, patient-reported health status assessment increases the patient’s role, which can motivate initiation and uptitration of medical therapy. However, routine assessment of patient-reported status increases the burden of data collection for patients and health systems and underscores the need for future studies evalu- ating the impact of assessment.

## Evidence Gaps and Future Research Directions

Significant gaps exist despite evolving evidence and treatment strategies in patients with HF. Table 33 pro- vides selected, common issues that should be addressed in future clinical research.

# ACC/AHA JOINT COMMITTEE MEMBERS

Joshua A. Beckman, MD, MS, FAHA, FACC, Chair; Pat- rick T. O’Gara, MD, MACC, FAHA, Immediate Past Chair\*; Sana M. Al-Khatib, MD, MHS, FACC, FAHA\*; Anastasia

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\*Former Task Force member; current member during the writing effort.

# PRESIDENTS AND STAFF

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and Digital Strategy

Grace D. Ronan, Team Leader, Clinical Policy Publications Timothy W. Schutt, MA, Clinical Practice Guidelines

Analyst

## American College of Cardiology/American Heart Association

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Abdul R. Abdullah, MD, Director, Guideline Science and Methodology

## American Heart Association

Donald M. Lloyd-Jones, MD, ScM, FAHA, President Nancy Brown, Chief Executive Officer

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Paul St. Laurent, DNP, RN, Senior Science and Medicine Advisor, Office of Science, Medicine and Health

Jody Hundley, Production and Operations Manager, Scientific Publications, Office of Science Operations

### ARTICLE INFORMATION

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### Class of Recommendation and Level of Evidence

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### Classification of HF by Left Ventricular Ejection Fraction (LVEF)

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#### *Management of Stage C HF: Activity, Exercise* Prescription, and Cardiac Rehabilitation

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### HF and Pregnancy

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##### Appendix 1. Author Relationships With Industry and Other Entities (Relevant)—2022 AHA/ACC/HFSA Guideline for the Management of Heart Failure

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| **Committee Member** | **Employment** | **Consultant** | **Speakers Bureau** | **Ownership/ Partnership/ Principal** | **Personal Research** | **Institutional, Organizational, or Other Financial Benefit** | **Expert Witness** | **Salary** |
| Paul A. Heidenreich, Chair | Stanford University School of Medi- cine—Professor and Vice-Chair for Quality, Department of Medicine | None | None | None | None | None | None | None |
| Biykem Bozkurt, Vice Chair | Baylor College of Medicine and De- Bakey VA Medical Center Cardiology Department—Mary and Gordon Cain Chair; W.A. “Tex” and Deborah Mon- crief, Jr., Chair; Professor of Medicine Medical Care Line Executive, DeBakey VA Medical Center; Director, Winters Center for Heart Failure Research; Associate Director, Cardiovascular Research Institute; Vice-Chair of Medi- cine, Baylor College of Medicine | * Abbott\* * Amgen * Baxter * Bristol-Myers Squibb\* * E.R. Squibb & Sons, L.L.C.\* * Relypsa * Sanofi-aventis\* * scPharmaceuticals * Vifor | None | None | None | * Abbott\* * Amgen * Relypsa * Respicardia * Sanofi- Aventis‡ * Past Presi- dent, HFSA (2019-2020) | None | None |
| David Aguilar | University of Kentucky—Professor of Medicine, Department of Medicine, Division of Cardiovascular Medicine | None | None | None | None | None | None | None |
| Larry A. Allen | University of Colorado School of Medicine, Anschutz Medical Campus—Professor of Medicine,  Department of Medicine, Division of Cardiology | * Abbott * ACI Clinical\* * Amgen\* * Boston Scientific\* * Cytokinetics * Novartis | None | None | None | * Abbott‡ * Amgen * Boston Sci- entific\*‡ * Janssen Phar- maceuticals * Medtronic Vascular Inc. * Novartis‡ | None | None |
| Joni J. Byun | Penultimate PR—President | None | None | None | None | None | None | None |
| Monica M. Colvin | University of Michigan—Professor of Medicine, Department of Medicine, Cardiovascular Division;  Associate Director, Heart Transplant Program, Advanced Heart Failure, Transplant, and MCS Natera | None | None | None | * CareDX | * Abbott‡ | None | None |
| Anita Deswal | UT MD Anderson Cancer Center— Ting Tsung and Wei Fong Chao Distinguished Chair, Professor of Medicine, and Chair of Cardiology | None | None | None | None | None | None | None |
| Mark H. Drazner | UT Southwestern Medical Cen- ter—Professor and Clinical Chief of Cardiology, Department of Internal Medicine, Cardiology | None | None | None | None | None | None | None |
| Shannon M. Dunlay | Mayo Clinic—Professor of Health Services Research and Medicine, De- partment of Cardiovascular Medicine | None | None | None | None | None | None | None |
| Linda R. Evers | Stevens & Lee—Shareholder and Chair of Stevens & Lee’s Energy, Communi- cations and Public Utility Group | None | None | None | None | None | None | None |
| James C. Fang | University of Utah—Professor of Medicine, Division of Cardiovascular Medicine | * Boston Scientific† | None | None | None | * ACI Clinical (Adjudication Committee)\* * Amgen (Steering Committee) * AstraZeneca (Steering Committee) * Boston Scientific† * Novartis (Executive Committee) | None | None |

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| **Committee Member** | **Employment** | **Consultant** | **Speakers Bureau** | **Ownership/ Partnership/ Principal** | **Personal Research** | **Institutional, Organizational, or Other Financial Benefit** | **Expert Witness** | **Salary** |
| Savitri E. Fedson | Michael E. DeBakey Medical Center—Professor, Medical Director, Advanced Heart Failure and Trans- plantation, Section of Cardiology | None | None | None | None | None | None | None |
| Gregg C. Fonarow | Geffen School of Medicine at UCLA—Professor of Cardiovascular Medicine, Chief, UCLA Division of Cardiology, Department of Medicine | * Abbott\* * Amgen * AstraZeneca * CHF Solutions * Cytokinetics * Edwards Life- sciences\* * Janssen Pharmaceuticals * Medtronic * Merck\* * Novartis\* * Regeneron | None | None | None | * Boston Scientific * Novartis\* * Medtronic | None | None |
| Salim S. Hayek | University of Michigan in Ann Arbor— Assistant Professor, Department of Medicine, Division of Cardiology | None | None | None | None | None | None | None |
| Adrian F. Hernandez | Duke University School of Medi- cine—Vice Dean of Clinical Research | * Amgen * AstraZeneca * Bayer * BioFourmis * Boehringer Ingelheim\* * Boston Scientific\* * Cytokinetics * Eli Lilly * Merck\* * Myokardia * Novartis\* * Pfizer * Relypsa * Sanofi-aventis\* | None | None | * American Regent * Amgen * Boston Scientific * AstraZen- eca\* * Boehringer Ingelheim * Daiichi Sankyo * Genentech * GlaxoS- mithKline\* * Janssen Pharma- ceuticals\* * Merck\* * Novartis\* * Verily\* | None | None | None |
| Prateeti Kha- zanie | University of Colorado—Assistant Professor of Medicine, Department of Medicine, Division of Cardiology | None | None | None | None | None | None | None |
| Michelle M. Kittleson | Smidt Heart Institute Cedars-Sinai— Professor of Medicine, Cardiology | None | None | None | None | None | None | None |
| Christopher  S. Lee | Boston College, William F. Connell School of Nursing—Professor and Associate Dean for Research | None | None | None | None | None | None | None |
| Mark S. Link | UT Southwestern Medical Center, Dallas—Professor of Medicine and Director, Cardiac Electrophysiology; Laurence and Susan Hirsch/Centex Distinguished Chair in Heart Dis- ease, Department of Internal Medi- cine, Division of Cardiology | None | None | None | None | None | None | None |
| Carmelo A. Milano | Duke University Medical Cen- ter—Professor of Surgery, Surgery Department | * Abbott\* | None | None | * Abbott\* * Medtronic\* * NuPulse† | * Abbott\* * AbioMed† * Allergan† * CryoLife† * Ethicon† * LivaNova† | None | None |

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| **Committee Member** | **Employment** | **Consultant** | **Speakers Bureau** | **Ownership/ Partnership/ Principal** | **Personal Research** | **Institutional, Organizational, or Other Financial Benefit** | **Expert Witness** | **Salary** |
| Lorraine C. Nnacheta§ | American Heart Association/Ameri- can College of Cardiology—Guide- line Advisor | None | None | None | None | * AHA/ACC salaried employee | None | None |
| Alexander T. Sandhu | Stanford University—Instructor of Medicine, Division of Cardiology, Department of Medicine | None | None | None | None | None | None | None |
| Lynne Warner Stevenson | Vanderbilt University Medical Center, Vanderbilt Heart and Vascular Insti- tute—Director of Cardiomyopathy and Lisa M. Jacobson Professor of Cardiovascular Medicine | * Novartis | None | None | None | * Abbott‡ * Biotronik * Boston Scientific * Bristol-Myers Squibb‡ * Endotronic† * Gore Medical† * Johnson & Johnson | None | None |
| Orly Vardeny | Minneapolis VA Health Care System and University of Minnesota— Associate Professor of Medicine, Department of Medicine | * Amgen * Novartis * Sanofi-Pasteur | None | None | * AstraZen- eca\* * Bayer\* | * AstraZeneca‡ | None | None |
| Amanda R. Vest | Tufts Medical Center, Cardiovascu- lar Center—Assistant Professor of Medicine | None | None | None | None | * Boehringer Ingelheim‡\* * CareDx‡ * Corvia‡ * Transmedics‡ | None | None |
| Clyde W. Yancy | Northwestern University, Feinberg School of Medicine; Northwestern Memorial Hospital—Professor of Medicine (Cardiology); Professor of Medical Social Sciences; Chief of Cardiology; Vice-Dean, Diversity & Inclusion; Associate Director Car- diovascular Institute Internal Medi- cine/Cardiology & Medical Social Sciences | None | None | None | None | * Abbott† * *JAMA Cardiology*, Deputy Editor | None | None |

This table represents the relationships of committee members with industry and other entities that were determined to be relevant to this document. These relation- ships 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 5% of the voting stock or share of the business entity, or ownership of $5000 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 ad- dressed 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.

†No financial benefit.

‡This disclosure was entered under the Clinical Trial Enroller category in the ACC’s disclosure system. To appear in this category, the author acknowledges that there is no *direct* or *institutional* relationship with the trial sponsor as defined in the (ACCF or AHA/ACC) Disclosure Policy for Writing Committees.

§Lorraine Nnacheta is an AHA/ACC joint staff member and acts as the guideline advisor for the “2022 AHA/ACC/HFSA Guideline for the Management of Heart Failure.” No relevant relationships to report. Non-voting author on recommendations and not included/counted in the RWI balance for this committee.

ACC indicates American College of Cardiology; ACCF, American College of Cardiology Foundation; AHA, American Heart Association; HFSA, Heart Failure Society of America; RWI, relationships with industry and other entities; UCLA, University of California, Los Angeles; UT, University of Texas; and VA, Veterans Affairs.

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##### Appendix 2. Reviewer Relationships With Industry and Other Entities (Comprehensive)—2022 AHA/ACC/HFSA Guideline for the Management of Heart Failure (June 2021)

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| **Reviewer** | **Representation** | **Employment** | **Consultant** | **Speakers Bureau** | **Ownership/ Partnership/ Principal** | **Personal Research** | **Institutional, Organizational,**  **or Other Financial Benefit** | **Expert Witness** |
| Anastasia Armbruster | Content Reviewer— Joint Committee on Clinical Practice Guidelines | University of Health Sciences & Pharmacy in St. Louis | None | * AstraZen- eca Phar- maceuticals | None | None | None | None |
| Alison Bailey | Content Reviewer— ACC | Centennial Heart at Parkridge | None | None | None | None | * American Society of Preventive Cardiology† * OptumRx | None |
| Joshua A. Beckman | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Vanderbilt University | * Amgen * JanOne * Janssen Pharmaceuticals\* | None | * EMX† * JanaCare† | * Bayer (DSMB) * Novartis | * Vascular Interven- tional Advances\* * NovoNordisk‡ | None |
| Claudio Bravo | Content Reviewer— AHA | University of Washington | None | None | None | None | None | None |
| Patricia Chang | Content Reviewer— AHA/ACC | University of North Carolina | None | None | None | None | None | None |
| Richard Cheng | Content Reviewer— AHA | University of Washington | None | None | None | * Eidos * Ionis | None | None |
| Lisa de las Fuentes | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Washington University in St. Louis | * Acceleron * Aerovate * Altavant * Arena * Bayer * Complexa\* * Express Scripts * Gossamer * Johnson&Johnson * Phase Bio * Sommetrics * V-wave * Vaderis * WebMD\* | * Simply Speaking\* | None | * Acceleron\* * Bayer * Complexa\* * Johnson&Johnson\* * Liquidia\* * Medtronic\* * NIH\* * Trio Analytics * United Therapeutics\* * University of Kentucky (DSMB)† * University of Toronto (DSMB)† | * ACC† * AHA† * *Circulation* Journals * Foundation for the NIH† * Pulmonary Hypertension Association\* | None |
| Akshay Desai | Content Reviewer— AHA/ACC | Brigham and Women’s Hospital | * Abbott Laboratories\* * Alnylam\* * Amgen\* * AstraZeneca Pharmaceuticals\* * Biofourmis\* * Boehringer Ingelheim * Boston Scientific\* * Corvidia Therapeutics\* * Cytokinetics * Dalcor Pharma\* * Lupin Pharma * Merck * Novartis\* * Regeneron\* * Relypsa\* * Sun Pharma | None | None | * Alnylam\* * AstraZeneca Pharmaceuticals\* * Bayer† * Myokardia† * Novartis\* | * Baim Clinical Research Institute\* * TIMI Study Group\* | None |
| Howard Eisen | Official Reviewer— AHA | Penn State Health | None | None | None | None | None | None |
| Mona Fiuzat | Content Reviewer— AHA/ACC | Duke University | * Fire1 | None | None | * Roche\* | None | None |
| Bulent Gorenek | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Eskisehir Os- mangazi University | * AstraZeneca Pharmaceuticals * Bayer * Daiichi Sankyo * Roche * Sandoz * Sanofi | None | None | None | None | None |

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| José A. Joglar | Content Reviewer— Joint Committee on Clinical Practice Guidelines | UT  Southwestern Medical Center | None | None | None | None | None | None |
| W. Schuyler Jones | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Duke University | * Bayer\* * Janssen Pharmaceuticals | None | None | * Boehringer Ingelheim * Bristol Myers Squibb * PCORI | None | None |
| Daniel Judge | Content Reviewer— AHA/ACC | The Medical University of South Carolina | * ADRx * Cytokinetics * Pfizer * Tenaya Therapeutics | None | None | * Capricor (DSMB) * TRiNDS (DSMB) | * Array Biopharma‡ * Eidos Therapeutics‡ * Myokardia‡ | None |
| Kimberly Ketter | Lay Reviewer | Morris Cardiovascular & Risk Reduc- tion Center | None | None | None | None | * Verilogue | None |
| Dharam Kumbhani | Content Reviewer— AHA/ACC | UT  Southwestern Medical Center | * ACC\* | None | None | None | * *Circulation*, Associate Editor\* | None |
| Daniel Mark | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Duke University | * Novartis | None | None | None | * Elsevier\* * Heartflow\* * Merck\* * NIH\* | None |
| Paul Mather | Content Reviewer— AHA/ACC | University of Pennsylvania | None | None | None | * Novartis\* | None | None |
| Shweta Motiwala | Content Reviewer— AHA/ACC | Harvard University | * Baim Institute for Clinical Research\* * Eli Lilly * Relay Therapeutics\* | None | * Relay   Therapeu- tics\* | * Puma Biotechnology\* * Relypsa† | * American Regent‡ * Boehringer Ingelheim‡ * Edwards Lifesciences‡ * NuPulse CV‡ | None |
| Debabrata Mukherjee | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Texas Tech University | * ACC\* | None | None | None | None | None |
| Patrick T. O’Gara | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Brigham and Women’s Hospital | None | None | None | None | * Edwards Lifesciences† * Medtronic† * *JAMA*\* * NIH\* | None |
| Francis D. Pagani | Official Reviewer— AHA | University of Michigan | None | None | None | None | None | None |
| Gurusher Panjrath | Content Reviewer— ACC | George Washington University | * CVRx | * Pfizer\* | None | None | * Abbott Laboratories‡ | None |
| Mariann Piano | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Vanderbilt University | None | None | None | None | None | None |
| Sean Pinney | Content Reviewer— AHA/ACC | University of Chicago | * Abbott Laboratories * CareDX, Inc. * Medtronic * NuPulse * Procyrion * Transmedics | None | None | None | None | None |
| Bunny Pozehl | Content Reviewer— AHA/ACC | University of Nebraska | None | None | None | None | None | None |
| Tanveer Rab | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Emory University | None | None | None | None | * ABIM * ACC† * American Bifurcation Club† | None |

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| Nosheen Reza | Content Reviewer— ACC | University of Pennsylvania | None | None | None | None | None | None |
| Jo E.  Rodgers | Content Reviewer— AHA | University of North Carolina | None | None | None | * Novartis† | * Duke Clinical Research Institute‡ * GlaxoSmithKline | None |
| Chris Salerno | Content Reviewer— ACC | St. Vincent Hospital | * Abbott | * Medtronic | None | None | * Abbott‡ * Medtronic‡ | None |
| Sanjiv Shah | Official Reviewer— HFSA | Northwestern University | * Abbott * ABIM * Amgen * Aria * AstraZeneca Pharmaceuticals\* * Axon * Bayer * Boehringer Ingelheim * Boston Scientific * Bristol Myers Squibb * Cardiora * CVRx * Cyclerion * Cytokinetics * Eisai * Ekoi.ai * GlaxoSmithKline * Imara * Ionis * Ironwood * Janssen Pharmaceuticals * Keyto * Eli Lilly * Merck * Myokardia * Novartis\* * NovoNordisk * Pfizer * Prothena * Regeneron * Sanofi * Shifamed * Tenax * United Therapeutics | * Pulmonary Hypertension Association | None | * Actelion\* * AHA\* * Covia\* * NIH\* | None | None |
| Erica S. Spatz | Official Reviewer, Joint Committee on Clinical Practice Guidelines | Yale University | None | None | None | None | * Centers for Medi- care & Medicaid Services\* * US Food and Drug Administration\* | None |
| Nancy Sweitzer | Official Reviewer— HFSA | University of Arizona | * Medscape\* * Myokardia | None | None | * NIH\* * Merck * Novartis† | * AHA\* * Amgen‡ * Array BioPharma‡ * Cellular Logistics† * Corvia Medical‡ * CVRx‡ * NIH\*‡ * University of Arizona\* | None |
| Jacqueline  E. Tamis- Holland | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Mount Sinai | * Gaffney Events Educational Trust | None | None | None | * AHA† * Bronx Lebanon Hospital, Cardiology Fellowship Program Director† * NYS† * The NGS Predict Study‡ | None |

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| Jennifer Thibodeau | Content Reviewer— AHA | UT  Southwestern Medical Center | None | None | None | None | * CareDX‡ * Cytokinetics‡ * Eidos Therapeutics‡ * NIH‡ | None |
| Sanjeev Trehan | Official Reviewer— ACC (Board of Governors) | Saint Francis Health System | None | None | None | None | None | None |
| Mary Norine Walsh | Content Reviewer— AHA/ACC | Ascension Medical Group | None | None | None | None | * Amgen‡ * EBR Systems * PCORI‡ * Thoratec‡ * Uppsala University‡ | None |
| Barbara Wiggins | Content Reviewer— ACC | Medical University of South Carolina | * Lexicomp | None | None | None | * ACC† * *American Journal of Cardiovascular* Drugs† * PERT Consortium Clinical Protocols† | None |
| Y. Joseph Woo | Content Reviewer— Joint Committee on Clinical Practice Guidelines | Stanford University School of Medicine | None | None | None | None | * Associate Editor, *Journal of Thoracic and Cardiovascular Disease* * NIH\* | None |

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ABIM indicates American Board of Internal Medicine; ACC, American College of Cardiology; ACCF, American College of Cardiology Foundation; AHA, American Heart As- sociation; DSMB, Data and Safety Monitoring Board; JAMA, Journal of the American Medical Association; NGS, Next-generation sequencing; NIH, National Institutes of Health; NYS, New York State; PCORI, Patient-Centered Outcomes Research Institute; PERT, Pulmonary Embolism Response Team; TIMI, Thrombolysis in Myocardial Infarction; and UT, University of Texas.

##### Appendix 3. Appendix for Tables 3 and 4: Suggested Thresholds for Structural Heart Disease and Evidence of Increased Filling Pressures

|  |  |
| --- | --- |
| Morphology | * LAVI 29 mL/m2 * LVMI >116/95 g/m2 * RWT >0.42 * LV wall thickness 12 mm |
| Ventricular systolic function | * LVEF <50% * GLS <16% |
| Ventricular diastolic function | * Average E/eʹ 15 for increased filling pressures * Septal eʹ <7 cm/s * Lateral eʹ <10 cm/s * TR velocity >2.8 m/s * Estimated PA systolic pressure >35 mm Hg |
| Biomarker | * BNP 35 pg/mL\* * NT-proBNP 125 pg/mL\* |

AF indicates atrial fibrillation; BNP, brain natriuretic peptide; CKD, chronic kidney disease; GLS, global longitudinal strain; HF, heart failure; LAVI, left atrial volume index; LV, left ventricular; LVEF, left ventricular ejection fraction; LVMI, left ventricular mass index; NT-proBNP, natriuretic peptide tests; PA, pulmonary artery; RWT, relative wall thickness; and TR, tricuspid regurgitation.

\*Cutoffs provided for natriuretic peptide levels may have lower specificity, especially in older patients or in patients with AF or CKD. Usually, higher cutoff values are recommended for the diagnosis of HF in these patients. Natriuretic peptide cutoffs selected for population screening for pre-HF (stage B HF) may be <99% reference limits and need to be defined according to the population at risk.