

CLINICAL PRACTICE GUIDELINE

# 2017 AHA/ACC/HRS Guideline for Management of Patients With Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death



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

*Developed in Collaboration With the Heart Failure Society of America*

**Writing Committee Members**

Sana M. Al-Khatib, MD, MHS, FACC, FAHA, FHRS, *Chair*  
William G. Stevenson, MD, FACC, FAHA, FHRS,  
*Vice Chair\**

Michael J. Ackerman, MD, PhD\*†  
William J. Bryant, JD, LLM†  
David J. Callans, MD, FACC, FHRS\*‡  
Anne B. Curtis, MD, FACC, FAHA, FHRS\*†  
Barbara J. Deal, MD, FACC, FAHA†  
Timm Dickfeld, MD, PhD, FHRS\*†  
Michael E. Field, MD, FACC, FAHA, FHRS†  
Gregg C. Fonarow, MD, FACC, FAHA, FHFSA\*§  
Anne M. Gillis, MD, FHRS\*†  
Christopher B. Granger, MD, FACC, FAHA\*†

Stephen C. Hammill, MD, FACC, FHRS‡  
Mark A. Hlatky, MD, FACC, FAHA†  
José A. Joglar, MD, FACC, FAHA, FHRS||  
G. Neal Kay, MD†  
Daniel D. Matlock, MD, MPH†  
Robert J. Myerburg, MD, FACC†  
Richard L. Page, MD, FACC, FAHA, FHRS‡

\*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. ‡HRS Representative. §ACC/AHA Task Force on Performance Measures Liaison/HFSA Representative. ||ACC/AHA Task Force on Clinical Practice Guidelines Liaison.

This document was approved by the American College of Cardiology Clinical Policy Approval Committee, the American Heart Association Science Advisory and Coordinating Committee, and the Heart Rhythm Society in September 2017, and the American Heart Association Executive Committee in October 2017.

The American College of Cardiology requests that this document be cited as follows: Al-Khatib SM, Stevenson WG, Ackerman MJ, Bryant WJ, Callans DJ, Curtis AB, Deal BJ, Dickfeld T, Field ME, Fonarow GC, Gillis AM, Granger CB, Hammill SC, Hlatky MA, Joglar JA, Kay GN, Matlock DD, Myerburg RJ, Page RL. 2017 AHA/ACC/HRS guideline for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol* 2018;72:e91–220.

This article has been copublished in *Circulation* and *HeartRhythm*.

Copies: This document is available on the World Wide Web sites of the American College of Cardiology ([www.acc.org](http://www.acc.org)), the American Heart Association ([professional.heart.org](http://professional.heart.org)), and the Heart Rhythm Society ([www.hrsonline.org](http://www.hrsonline.org)). For copies of this document, please contact the Elsevier Inc. Reprint Department via fax (212-633-3820) or e-mail ([reprints@elsevier.com](mailto:reprints@elsevier.com)).

Permissions: Multiple copies, modification, alteration, enhancement, and/or distribution of this document are not permitted without the express permission of the American College of Cardiology. Requests may be completed online via the Elsevier site (<http://www.elsevier.com/about/policies/author-agreement/obtaining-permission>).

<b>ACC/AHA Task Force Members</b>	Glenn N. Levine, MD, FACC, FAHA, <i>Chair</i> Patrick T. O’Gara, MD, MACC, FAHA, <i>Chair-Elect</i> Jonathan L. Halperin, MD, FACC, FAHA, <i>Immediate Past Chair</i> ¶
	Sana M. Al-Khatib, MD, MHS, FACC, FAHA Joshua A. Beckman, MD, MS, FAHA Kim K. Birtcher, MS, PHARM, AAC Biykem Bozkurt, MD, PhD, FACC, FAHA¶ Ralph G. Brindis, MD, MPH, MACC¶ Joaquin E. Cigarroa, MD, FACC Anita Deswal, MD, MPH, FACC, FAHA Lesley H. Curtis, PhD, FAHA¶ Lee A. Fleisher, MD, FACC, FAHA

Federico Gentile, MD, FACC Samuel Gidding, MD, FAHA¶ Zachary D. Goldberger, MD, MS, FACC, FAHA Mark A. Hlatky, MD, FACC, FAHA John Ikonomidis, MD, PhD, FAHA José A. Joglar, MD, FACC, FAHA Laura Mauri, MD, MSc, FAHA Barbara Riegel, PhD, RN, FAHA Susan J. Pressler, PhD, RN, FAHA¶ Duminda N. Wijeysundera, MD, PhD
--

¶Former Task Force member; current member during the writing effort.

## TABLE OF CONTENTS

<b>PREAMBLE</b>	e93
<b>1. INTRODUCTION</b>	e95
1.1. Methodology and Evidence Review	e95
1.2. Organization of the Writing Committee	e95
1.3. Document Review and Approval	e95
1.4. Scope of the Guideline	e97
1.5. Abbreviations	e99
<b>2. EPIDEMIOLOGY</b>	e99
2.1. General Concepts	e99
2.1.1. Premature Ventricular Complexes and Non-sustained VT	e99
2.1.2. VT and VF During ACS	e101
2.1.3. Sustained VT and VF Not Associated With ACS	e102
2.2. Sudden Cardiac Death	e102
2.2.1. Incidence of SCD	e102
2.2.2. Population Subgroups and Risk Prediction	e102
<b>3. MECHANISMS OF VA</b>	e103
3.1. Cellular Mechanisms and Substrates	e103
3.2. Automaticity	e104
3.3. Triggered Activity	e104
3.4. Reentry	e104
<b>4. GENERAL EVALUATION OF PATIENTS WITH DOCUMENTED OR SUSPECTED VA</b>	e105
<b>4.1. History and Physical Examination</b>	e105
<b>4.2. Noninvasive Evaluation</b>	e106
4.2.1. 12-lead ECG and Exercise Testing	e106
4.2.2. Ambulatory Electrocardiography	e107
4.2.3. Implanted Cardiac Monitors	e108
4.2.4. Noninvasive Cardiac Imaging	e108
4.2.5. Biomarkers	e109
4.2.6. Genetic Considerations in Arrhythmia Syndromes	e109
<b>4.3. Invasive Testing</b>	e110
4.3.1. Invasive Cardiac Imaging: Cardiac Catheterization or CT Angiography	e110
4.3.2. Electrophysiological Study for VA	e111
<b>5. THERAPIES FOR TREATMENT OR PREVENTION OF VA</b>	e111
<b>5.1. Medication Therapy</b>	e111
5.1.1. Medications With Prominent Sodium Channel Blockade	e112
5.1.2. Beta Blockers	e112
5.1.3. Amiodarone and Sotalol	e112
5.1.4. Calcium Channel Blockers	e115
5.1.5. Nonantiarrhythmic Medications and Therapies	e115
<b>5.2. Preventing SCD With HF Medications</b>	e116
<b>5.3. Defibrillators for Treatment of VA and SCD</b>	e116
<b>5.4. Catheter Ablation</b>	e117
5.4.1. General Considerations	e117
5.4.2. VA in Patients With No Apparent Structural Heart Disease	e117
5.4.3. Scar-Related VT	e117

<b>5.5. Surgery and Revascularization Procedures in Patients With Ischemic Heart Disease . . . . .</b>	e117
5.5.1. Surgery for Arrhythmia Management . . . . .	e118
<b>5.6. Autonomic Modulation . . . . .</b>	e118
<b>6. ACUTE MANAGEMENT OF SPECIFIC VA . . . . .</b>	e119
<b>7. ONGOING MANAGEMENT OF VA AND SCD RISK RELATED TO SPECIFIC DISEASE STATES . . . . .</b>	e123
<b>7.1. Ischemic Heart Disease . . . . .</b>	e123
7.1.1. Secondary Prevention of SCD in Patients With Ischemic Heart Disease . . . . .	e123
7.1.2. Primary Prevention of SCD in Patients With Ischemic Heart Disease . . . . .	e126
7.1.3. Treatment and Prevention of Recurrent VA in Patients With Ischemic Heart Disease . . . . .	e128
<b>7.2. Nonischemic Cardiomyopathy . . . . .</b>	e131
7.2.1. Secondary Prevention of SCD in Patients With NICM . . . . .	e132
7.2.2. Primary Prevention of SCD in Patients With NICM . . . . .	e133
7.2.3. Treatment of Recurrent VA in Patients With NICM . . . . .	e134
<b>7.3 Arrhythmogenic Right Ventricular Cardiomyopathy . . . . .</b>	e135
<b>7.4. Hypertrophic Cardiomyopathy . . . . .</b>	e139
<b>7.5. Myocarditis . . . . .</b>	e142
<b>7.6. Cardiac Sarcoidosis . . . . .</b>	e143
7.6.1. Other Infiltrative Cardiomyopathies . . . . .	e145
<b>7.7. Heart Failure . . . . .</b>	e146
7.7.1. HF With Reduced Ejection Fraction . . . . .	e146
7.7.2. HF With Preserved Ejection Fraction . . . . .	e146
7.7.3. Left Ventricular Assist Device . . . . .	e147
7.7.4. ICD Use After Heart Transplantation . . . . .	e147
<b>7.8. Neuromuscular Disorders . . . . .</b>	e148
<b>7.9. Cardiac Channelopathies . . . . .</b>	e149
7.9.1. Specific Cardiac Channelopathy Syndromes . . . . .	e150
<b>8. VA IN THE STRUCTURALLY NORMAL HEART . . . . .</b>	e160
<b>8.1. Outflow Tract and Atrioventricular Annular VA . . . . .</b>	e161
<b>8.2. Papillary Muscle VA . . . . .</b>	e162
<b>8.3. Interfascicular Reentrant VT (Belhassen Tachycardia) . . . . .</b>	e162
<b>8.4. Idiopathic Polymorphic VT/VF . . . . .</b>	e163
<b>9. PVC-INDUCED CARDIOMYOPATHY . . . . .</b>	e164
<b>10. VA AND SCD RELATED TO SPECIFIC POPULATIONS . . . . .</b>	e165
<b>10.1. Athletes . . . . .</b>	e165
<b>10.2. Pregnancy . . . . .</b>	e166
<b>10.3. Older Patients With Comorbidities . . . . .</b>	e166
<b>10.4. Chronic Kidney Disease . . . . .</b>	e167
<b>10.5. Valvular Heart Disease . . . . .</b>	e167
<b>10.6. Sex-Related Differences in the Risk of SCD . . . . .</b>	e168
<b>10.7. Medication-Induced Arrhythmias . . . . .</b>	e168
<b>10.8. Adult Congenital Heart Disease . . . . .</b>	e170
<b>11. DEFIBRILLATORS OTHER THAN TRANSVENOUS ICDs . . . . .</b>	e175
<b>11.1. Subcutaneous Implantable Cardioverter-Defibrillator . . . . .</b>	e175
<b>11.2. Wearable Cardioverter-Defibrillator . . . . .</b>	e176
<b>11.3. Automated External Defibrillator . . . . .</b>	e177
<b>12. SPECIAL CONSIDERATIONS FOR CATHETER ABLATION . . . . .</b>	e177
<b>13. POSTMORTEM EVALUATION OF SCD . . . . .</b>	e178
<b>14. TERMINAL CARE . . . . .</b>	e179
<b>15. SHARED DECISION-MAKING . . . . .</b>	e180
<b>16. COST AND VALUE CONSIDERATIONS . . . . .</b>	e181
<b>17. QUALITY OF LIFE . . . . .</b>	e182
<b>18. EVIDENCE GAPS AND FUTURE RESEARCH NEEDS . . . . .</b>	e182
<b>APPENDIX 1</b>	
Author Relationships With Industry and Other Entities (Relevant) . . . . .	e214
<b>APPENDIX 2</b>	
Reviewer Relationships With Industry and Other Entities (Comprehensive) . . . . .	e216
<b>PREAMBLE</b>	
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 methods to evaluate and classify evidence, provide a cornerstone for quality cardiovascular care. The ACC and AHA sponsor the development and publication of guidelines without	

commercial support, and members of each organization volunteer their time to the writing and review efforts. Guidelines are official policy of the ACC and AHA.

### Intended Use

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

### Clinical Implementation

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

### Methodology and Modernization

The ACC/AHA Task Force on Clinical Practice Guidelines (Task Force) continuously reviews, updates, and modifies guideline methodology on the basis of published standards from organizations including the Institute of Medicine ([P-1,P-2](#)) and on the basis of internal reevaluation. Similarly, the presentation and delivery of guidelines are reevaluated and modified on the basis of evolving technologies and other factors to facilitate optimal dissemination of information at the point of care to healthcare professionals.

Toward this goal, this guideline heralds the introduction of an evolved format of presenting guideline recommendations and associated text called the “modular knowledge chunk format”. Each modular “chunk” includes a table of related recommendations, a brief synopsis, recommendation-specific supportive text and, when appropriate, flow diagrams or additional tables. References are provided within the modular chunk itself to facilitate quick review. This format also will facilitate seamless updating of guidelines with focused updates as new evidence is published, and content tagging for rapid electronic retrieval of related recommendations on a topic of interest. This evolved format was instituted when this guideline was near completion; therefore, the current document represents a transitional formatting that best suits the text as written. Future guidelines will fully implement this format, including provisions for limiting the amount of text in a guideline.

Recognizing the importance of cost-value considerations in certain guidelines, when appropriate and feasible, an analysis of the value of a medication, device, or intervention may be performed in accordance with the ACC/AHA methodology ([P-3](#)).

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

### Selection of Writing Committee Members

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

### Relationships With Industry and Other Entities

The ACC and AHA have rigorous policies and methods to ensure that guidelines are developed without bias or improper influence. The complete relationships with industry and other entities (RWI) policy can be found [online](#). [Appendix 1](#) of the current document lists writing committee members' relevant RWI. For the purposes of full transparency, writing committee members' comprehensive disclosure information is available [online](#), as is the [comprehensive disclosure information for the Task Force](#).

### Evidence Review and Evidence Review Committees

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

An independent evidence review committee (ERC) is commissioned when there are  $\geq 1$  questions deemed of utmost clinical importance that merit formal systematic review. This systematic review will strive to determine which patients are most likely to benefit from a test, medication, device, or treatment strategy and to what

degree. Criteria for commissioning an ERC and formal systematic review include: a) the absence of a current authoritative systematic review; b) the feasibility of defining the benefit and risk in a time frame consistent with the writing of a guideline; c) the relevance to a substantial number of patients; and d) the likelihood that the findings can be translated into actionable recommendations. ERC members may include methodologists, epidemiologists, healthcare providers, and biostatisticians. When a formal systematic review has been commissioned, the recommendations developed by the writing committee on the basis of the systematic review are marked with “<sup>SR</sup>”.

### **Guideline-Directed Management and Therapy**

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

### **Class of Recommendation and Level of Evidence**

The Class of Recommendation (COR) indicates the strength of the recommendation, encompassing the estimated magnitude and certainty of benefit in proportion to risk. The Level of Evidence (LOE) rates the quality of scientific evidence that supports the intervention on the basis of the type, quantity, and consistency of data from clinical trials and other sources (Table 1) (P-4,P-6,P-8).

Glenn N. Levine, MD, FACC, FAHA

Chair, ACC/AHA Task Force on Clinical Practice Guidelines

## **1. INTRODUCTION**

### **1.1. Methodology and Evidence Review**

The recommendations listed in this clinical practice guideline are, whenever possible, evidence-based. An initial extensive evidence review, which included literature derived from research involving human subjects, published in English, and indexed in MEDLINE (through PubMed), EMBASE, the Cochrane Library, the Agency for Healthcare Research and Quality, and other selected databases relevant to this guideline, was conducted from April 2016 to September 2016. Key search words included, but were not limited, to the following: sudden cardiac death, ventricular tachycardia, ventricular fibrillation, premature ventricular contractions, implantable cardioverter-defibrillator, subcutaneous implantable cardioverter-defibrillator, wearable cardioverter-defibrillator, and catheter ablation. Additional relevant studies published through March 2017, during the

guideline writing process, were also considered by the writing committee, and added to the evidence tables when appropriate. The final evidence tables are included in the *Online Data Supplement* and summarize the evidence used by the writing committee to formulate recommendations. Additionally, the writing committee reviewed documents related to ventricular arrhythmias (VA) and sudden cardiac death (SCD) previously published by the ACC, AHA, and the Heart Rhythm Society (HRS). References selected and published in this document are representative and not all-inclusive.

As noted in the Preamble, an independent ERC was commissioned to perform a formal systematic review of 2 important clinical questions for which clear literature and prior guideline consensus were felt to be lacking or limited (Table 2). The results of the ERC review were considered by the writing committee for incorporation into this guideline. Concurrent with this process, writing committee members evaluated other published data relevant to the guideline. The findings of the ERC and the writing committee members were formally presented and discussed, then guideline recommendations were developed. The “Systematic Review for the 2017 AHA/ACC/HRS Guideline for Management of Patients With Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death” is published in conjunction with this guideline (S1.4-1).

The ACC and AHA have acknowledged the importance of value in health care and have called for eventual development of a Level of Value for clinical practice recommendations (S1.4-2). Available cost-effectiveness data were determined to be sufficient to support 2 specific recommendations in this guideline (see Sections 7.1.1 and 7.1.2). As a result, a Level of Value was assigned to those 2 recommendations on the basis of the “ACC/AHA Statement on Cost/Value Methodology in Clinical Practice Guidelines and Performance Measures,” as shown in Table 3 (S1.4-2). Available quality of life (QoL) data were deemed to be insufficient to support specific recommendations in this guideline.

### **1.2. Organization of the Writing Committee**

The writing committee consisted of cardiac electrophysiologists (including those specialized in pediatrics), general adult and pediatric cardiologists (including those specialized in critical care and acute coronary syndromes [ACS], genetic cardiology, heart failure, and cost-effectiveness analyses), a geriatrician with expertise in terminal care and shared decision-making, and a lay representative, in addition to representatives from the ACC, AHA, HRS, and the Heart Failure Society of America (HFSA).

### **1.3. Document Review and Approval**

This document was reviewed by 2 official reviewers nominated by the ACC, AHA, and HRS; 1 official lay

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

CLASS (STRENGTH) OF RECOMMENDATION		LEVEL (QUALITY) OF EVIDENCE†
<b>CLASS I (STRONG)</b>	<b>Benefit &gt;&gt; Risk</b>	<b>LEVEL A</b>
Suggested phrases for writing recommendations:		<ul style="list-style-type: none"> <li>■ Is recommended</li> <li>■ Is indicated/useful/effective/beneficial</li> <li>■ Should be performed/administered/other</li> <li>■ Comparative-Effectiveness Phrases‡:           <ul style="list-style-type: none"> <li>○ Treatment/strategy A is recommended/indicated in preference to treatment B</li> <li>○ Treatment A should be chosen over treatment B</li> </ul> </li> </ul>
<b>CLASS IIa (MODERATE)</b>	<b>Benefit &gt;&gt; Risk</b>	<b>LEVEL B-R</b> (Randomized)
Suggested phrases for writing recommendations:		<ul style="list-style-type: none"> <li>■ Is reasonable</li> <li>■ Can be useful/effective/beneficial</li> <li>■ Comparative-Effectiveness Phrases‡:           <ul style="list-style-type: none"> <li>○ Treatment/strategy A is probably recommended/indicated in preference to treatment B</li> <li>○ It is reasonable to choose treatment A over treatment B</li> </ul> </li> </ul>
<b>CLASS IIb (WEAK)</b>	<b>Benefit ≥ Risk</b>	<b>LEVEL B-NR</b> (Nonrandomized)
Suggested phrases for writing recommendations:		<ul style="list-style-type: none"> <li>■ May/might be reasonable</li> <li>■ May/might be considered</li> <li>■ Usefulness/effectiveness is unknown/unclear/uncertain or not well established</li> </ul>
<b>CLASS III: No Benefit (MODERATE)</b> (Generally, LOE A or B use only)	<b>Benefit = Risk</b>	<b>LEVEL C-LD</b> (Limited Data)
Suggested phrases for writing recommendations:		<ul style="list-style-type: none"> <li>■ Randomized or nonrandomized observational or registry studies with limitations of design or execution</li> <li>■ Meta-analyses of such studies</li> <li>■ Physiological or mechanistic studies in human subjects</li> </ul>
<b>CLASS III: Harm (STRONG)</b>	<b>Risk &gt; Benefit</b>	<b>LEVEL C-EO</b> (Expert Opinion)
Suggested phrases for writing recommendations:		Consensus of expert opinion based on clinical experience

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

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

\* The outcome or result of the intervention should be specified (an improved clinical outcome or increased diagnostic accuracy or incremental prognostic information).

† For comparative-effectiveness recommendations (COR I and IIa; LOE A and B only), studies that support the use of comparator verbs should involve direct comparisons of the treatments or strategies being evaluated.

‡ The method of assessing quality is evolving, including the application of standardized, widely used, and preferably validated evidence grading tools; and for systematic reviews, the incorporation of an Evidence Review Committee.

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

**TABLE 2** Systematic Review Questions on SCD Prevention

Question Number	Question	Section Number
1	For asymptomatic patients with Brugada syndrome, what is the association between an abnormal programmed ventricular stimulation study and SCD and other arrhythmia endpoints?	7.9.1.3
2	What is the impact of ICD implantation for primary prevention in older patients and patients with significant comorbidities?	10.3

ICD indicates implantable cardioverter-defibrillator; and SCD, sudden cardiac death.

**TABLE 3 Proposed Integration of Level of Value Into Clinical Practice Guideline Recommendations\***

**Level of Value**

**High value:** Better outcomes at lower cost or ICER <\$50,000 per QALY gained

**Intermediate value:** \$50,000 to <\$150,000 per QALY gained

**Low value:** ≥\$150,000 per QALY gained

**Uncertain value:** Value examined but data are insufficient to draw a conclusion because of no studies, low-quality studies, conflicting studies, or prior studies that are no longer relevant

**Not assessed:** Value not assessed by the writing committee

Proposed abbreviations for each value recommendation:

*Level of Value: H to indicate high value; I, intermediate value; L, low value; U, uncertain value; and NA, value not assessed*

\*Dollar amounts used in this table are based on U.S. GDP data from 2012 and were obtained from WHO-CHOICE Cost-Effectiveness Thresholds (S1.4-3).

GDP indicates gross domestic product; ICER, incremental cost-effectiveness ratio; QALY, quality-adjusted life-years; and WHO-CHOICE, World Health Organization Choosing Interventions that are Cost-Effective.

Reproduced from Anderson, et al. (S1.4-2).

reviewer nominated by the AHA; 1 organizational reviewer nominated by the HFSA; and 28 individual content reviewers. Reviewers' RWI information was distributed to the writing committee and is published in this document (Appendix 2).

This document was approved for publication by the governing bodies of the ACC, the AHA, and the HRS; and endorsed by the HFSA.

#### 1.4. Scope of the Guideline

The purpose of this AHA/ACC/HRS document is to provide a contemporary guideline for the management of adults who have VA or who are at risk for SCD, including diseases and syndromes associated with a risk of SCD from VA. This guideline supersedes the "ACC/AHA/ESC 2006 Guidelines for Management of Patients With Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death" (S1.4-4). It also supersedes some sections of the "ACC/AHA/HRS 2008 Guidelines for Device-Based Therapy of Cardiac Rhythm Abnormalities" (S1.4-5), specifically those sections on indications for the implantable cardioverter-defibrillator (ICD); and, it updates the SCD prevention recommendations in the "2011 ACCF/AHA Guideline for the Diagnosis and Treatment of Hypertrophic Cardiomyopathy" (S1.4-6). Some recommendations from the earlier guidelines have been updated as warranted by new evidence or a better understanding of existing evidence, and irrelevant or overlapping recommendations were deleted or modified.

In the current guideline, sudden cardiac arrest (SCA) is defined as the "sudden cessation of cardiac activity so that the victim becomes unresponsive, with no normal breathing and no signs of circulation" (S1.4-7). If corrective measures are not taken rapidly, this condition progresses to SCD. Cardiac arrest is used to signify an event that can be reversed, usually by cardiopulmonary resuscitation (CPR), administration of medications and/or defibrillation or cardioversion. SCA and SCD can result from causes other than VA, such as bradycardia, electromechanical dissociation, pulmonary embolism, intracranial hemorrhage, and aortic dissection; however,

the scope of this document includes only SCA and SCD due to VA.

This guideline includes indications for ICDs for the treatment of VA and prevention of SCD, but it does not delve into details on individual device selection and programming, including considerations relevant to cardiac resynchronization therapy (CRT), bradycardia pacing, and hemodynamic monitoring. These important aspects of ICD management have been covered in an HRS expert consensus statement (S1.4-8). An AHA science advisory discusses the use of wearable cardioverter-defibrillators (S1.4-9). The findings of that document were reviewed; however, recommendations on this topic were developed independently of that document. This guideline includes indications for catheter ablation of VA, but does not provide recommendations on specific techniques or ablation technologies, which were beyond the scope of this document.

Recommendations for interventional therapies, including ablation and the implantation of devices, apply only if these therapies can be implemented by qualified clinicians, such that outcomes consistent with published literature are a reasonable expectation. The writing committee agreed that a high degree of expertise was particularly important for performance of catheter ablation of VA, and this point is further emphasized in relevant sections. In addition, all recommendations related to ICDs require that meaningful survival of >1 year is expected; meaningful survival means that a patient has a reasonable quality of life and functional status.

Although this document is aimed at the adult population (≥18 years of age) and offers no specific recommendations for pediatric patients, some of the literature on pediatric patients was examined. In some cases, the data from pediatric patients beyond infancy helped to inform this guideline.

The writing committee recognized the importance of shared decision-making and patient-centered care and, when possible, it endeavored to formulate recommendations relevant to these important concepts. The

importance of a shared decision-making process in which the patient, family, and clinicians discuss risks and benefits of diagnostic and treatment options and consider the patients' personal preferences is emphasized (see [Section 15](#)).

In developing this guideline, the writing committee reviewed previously published guidelines and related statements. [Table 4](#) contains a list of guidelines and statements deemed pertinent to this writing effort and is intended for use as a resource, obviating repetition of existing guideline recommendations.

During final production review of the guidelines, several recommendations were refined to better reflect

the data and current recommended medical practice. These refinements were reviewed and approved by the writing committee, the Task Force, and ACC, AHA, and HRS organizational leadership. These recommendations were:

- [Section 7.1.1](#), recommendation 1
- [Section 7.1.3](#), recommendation 2
- [Section 7.2.1](#), recommendation 1
- [Section 7.9.1.4](#), recommendation 2
- [Section 10.8](#), recommendation 6

Readers should refer to these sections for the updated text.

**TABLE 4 Associated Guidelines and Statements**

Title	Organization	Publication Year (Reference)
<b>Guidelines</b>		
Syncpe	ACC/AHA/HRS	2017 ( <a href="#">S1.4-10</a> )
Heart failure	ACCF/AHA	2017 ( <a href="#">S1.4-11</a> ) 2016 ( <a href="#">S1.4-12</a> ), and 2013 ( <a href="#">S1.4-13</a> )
Valvular heart disease	AHA/ACC	2017 ( <a href="#">S1.4-14</a> ) and 2014 ( <a href="#">S1.4-15</a> )
Supraventricular tachycardia	ACC/AHA/HRS	2015 ( <a href="#">S1.4-16</a> )
Ventricular arrhythmias and the prevention of sudden cardiac death	ESC	2015 ( <a href="#">S1.4-17</a> )
Guidelines for cardiopulmonary resuscitation and emergency cardiovascular care	AHA	2015 ( <a href="#">S1.4-18</a> )
Atrial fibrillation	AHA/ACC/HRS	2014 ( <a href="#">S1.4-19</a> )
Non-ST-elevation acute coronary syndromes	AHA/ACC	2014 ( <a href="#">S1.4-20</a> )
Assessment of cardiovascular risk	ACC/AHA	2013 ( <a href="#">S1.4-21</a> )
ST-elevation myocardial infarction	ACCF/AHA	2013 ( <a href="#">S1.4-22</a> )
Acute myocardial infarction in patients presenting with ST-segment elevation	ESC	2012 ( <a href="#">S1.4-23</a> )
Device-based therapies for cardiac rhythm abnormalities	ACCF/AHA/HRS	2012 ( <a href="#">S1.4-24</a> )
Coronary artery bypass graft surgery	ACCF/AHA	2011 ( <a href="#">S1.4-25</a> )
Hypertrophic cardiomyopathy	ACCF/AHA	2011 ( <a href="#">S1.4-6</a> )
Percutaneous coronary intervention	ACCF/AHA/SCAI	2011 ( <a href="#">S1.4-26</a> )
Secondary prevention and risk reduction therapy for patients with coronary and other atherosclerotic vascular disease	AHA/ACCF	2011 ( <a href="#">S1.4-27</a> )
<b>Scientific Statements</b>		
Wearable cardioverter-defibrillator therapy for the prevention of sudden cardiac death	AHA	2016 ( <a href="#">S1.4-9</a> )
Optimal implantable cardioverter defibrillator programming and testing	HRS/EHRA/APHRS/SOLAECE	2016 ( <a href="#">S1.4-8</a> )
Treatment of cardiac arrest: current status and future directions: strategies to improve cardiac arrest survival	IOM	2015 ( <a href="#">S1.4-28</a> )
Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities	ACC/AHA	2015 ( <a href="#">S1.4-29</a> )
Ventricular arrhythmias	EHRA/HRS/APHRS	2014 ( <a href="#">S1.4-30</a> )
Arrhythmias in adult congenital heart disease	PACES/HRS	2014 ( <a href="#">S1.4-31</a> )
Implantable cardioverter-defibrillator therapy in patients who are not included or not well represented in clinical trials	HRS/ACC/AHA	2014 ( <a href="#">S1.4-32</a> )
Cardiac sarcoidosis	HRS	2014 ( <a href="#">S1.4-33</a> )
Inherited primary arrhythmia syndromes	HRS/EHRA/APHRS	2013 ( <a href="#">S1.4-34</a> )

ACC indicates American College of Cardiology; ACCF, American College of Cardiology Foundation; AHA, American Heart Association; APHRS, Asia Pacific Heart Rhythm Society; EHRA, European Heart Rhythm Association; ESC, European Society of Cardiology; HRS, Heart Rhythm Society; PACES, Pediatric and Congenital Electrophysiology Society; SCAI, Society for Cardiovascular Angiography and Interventions; and, SOLAECE, Sociedad Latinoamericana de Estimulación Cardíaca y Electrofisiología.

## 1.5. Abbreviations

Abbreviation	Meaning/Phrase
ACS	acute coronary syndromes
AED	automated external defibrillator
AMI	acute myocardial infarction
BNP	B-type natriuretic peptide
CABG	coronary artery bypass graft
CKD	chronic kidney disease
CPR	cardiopulmonary resuscitation
CRT	cardiac resynchronization therapy
CT	computed tomography
ECG	electrocardiogram
ERC	evidence review committee
ESRD	end-stage renal disease
GDMT	guideline-directed management and therapy
HCM	hypertrophic cardiomyopathy
HF	heart failure
HFP EF	heart failure with preserved ejection fraction
HFREF	heart failure with reduced ejection fraction
ICD	implantable cardioverter-defibrillator
LV	left ventricular
LVAD	left ventricular assist device
LVEF	left ventricular ejection fraction
MI	myocardial infarction
NICM	nonischemic cardiomyopathy
NSVT	nonsustained ventricular tachycardia
PET	positron emission tomography
PCI	percutaneous coronary intervention
PVC	premature ventricular complex
QoL	quality of life
RCT	randomized controlled trial
RV	right ventricular
RVOT	right ventricular outflow tract
SCA	sudden cardiac arrest
SCD	sudden cardiac death
SVT	supraventricular tachycardia
TOF	tetralogy of Fallot
VA	ventricular arrhythmia
VT	ventricular tachycardia

## 2. EPIDEMIOLOGY

### 2.1. General Concepts

Table 5

VA include a spectrum that ranges from premature ventricular complex (PVC) to ventricular fibrillation (VF), with a clinical presentation that ranges from a total lack of symptoms to cardiac arrest. Most life-threatening VA are associated with ischemic heart disease, particularly in older patients (S2.2.2-1). The risks of VA and SCD vary in

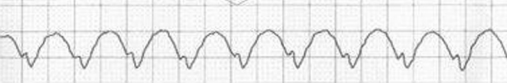
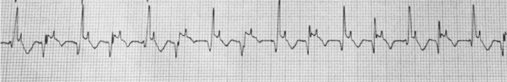
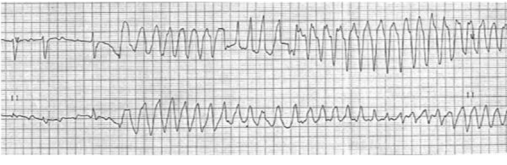
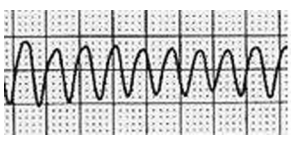
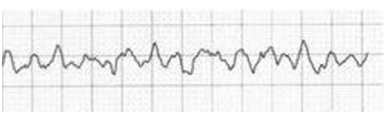
specific populations with different underlying cardiac conditions, and with specific family history and genetic variants, and this variation has important implications for studying and applying therapies.

#### 2.1.1. Premature Ventricular Complexes and Nonsustained VT

PVCs are common and increase in frequency with age. Although PVCs were found in a healthy military population in only 0.6% of those <20 years of age and 2.7% of those >50 years of age (S2.2.2-5) on 12-lead ECGs, longer term monitoring shows PVCs in about 50% of all people with or without heart disease (S2.2.2-6). The presence of PVCs on 2 minutes of monitoring of middle-aged patients in the ARIC (Atherosclerosis Risk In Communities) study was associated with increased risk of both ischemic heart disease events and mortality, with or without prevalent ischemic heart disease (S2.2.2-7,S2.2.2-8). In the general population, frequent PVCs, which are defined as the presence of at least 1 PVC on a 12-lead ECG or >30 PVCs per hour, are associated with increased cardiovascular risk and increased mortality (S2.2.2-9). In a study from Taiwan of patients without sustained VT or structural heart disease who had 24-hour Holter monitoring for clinical evaluation, multifocal PVCs were associated with increased risk of death and nonfatal cardiovascular adverse outcomes (S2.2.2-10). In the same population, nonsustained ventricular tachycardia (NSVT) was independently associated with increased risk of death and other cardiovascular adverse outcomes, including stroke (S2.2.2-11). An association of PVCs with increased risk of stroke was also seen in the ARIC population (S2.2.2-8).

Because some studies have shown an association of PVCs with adverse outcomes, the detection of PVCs, particularly if multifocal and frequent, is generally considered a risk factor for adverse cardiovascular outcomes, and such patients are generally evaluated to ensure they do not have underlying conditions (e.g., ischemic heart disease, left ventricular [LV] dysfunction) that warrant further treatment to reduce risk. PVC and NSVT in patients with cardiovascular disease are common and have been associated with adverse outcomes (S2.2.2-12,S2.2.2-13). In CAST (Cardiac Arrhythmia Suppression Trials), treatment of patients with post-myocardial infarction (MI) who took antiarrhythmic medications (e.g., flecainide, encainide, moricizine) increased the risk of death despite suppression of VA (S2.2.2-14,S2.2.2-15). Treatment of PVCs with antiarrhythmic medications has not been shown to reduce mortality and, in the post-MI population, treatment with class I sodium channel-blocking medications (e.g., quinidine, flecainide) increases the risk of death (S2.2.2-15,S2.2.2-16). Likewise, in patients with a reduced LVEF class I, sodium channel-blocking medications and d-sotalol increase the risk of death (S2.2.2-16,S2.2.2-17). Beta blockers,

**TABLE 5 Table of Definitions of Commonly Used Terms in this Document**

Term	Definition or Description
Ventricular tachycardia (S2.2.2-2)	<p>Cardiac arrhythmia of <math>\geq 3</math> consecutive complexes originating in the ventricles at a rate <math>&gt;100</math> bpm (cycle length: <math>&lt;600</math> ms). Types of VT:</p> <ul style="list-style-type: none"><li>■ Sustained: VT <math>&gt;30</math> s or requiring termination due to hemodynamic compromise in <math>&lt;30</math> s.</li><li>■ Nonsustained/unsustained: <math>\geq 3</math> beats, terminating spontaneously.</li><li>■ Monomorphic: Stable single QRS morphology from beat to beat.</li><li>■ Polymorphic: Changing or multiform QRS morphology from beat to beat.</li><li>■ Bidirectional: VT with a beat-to-beat alternation in the QRS frontal plane axis, often seen in the setting of digitalis toxicity or catecholaminergic polymorphic VT</li></ul> <p>Monomorphic VT</p> 
	<p>Polymorphic VT</p> 
	<p>Bidirectional VT</p> 
Torsades de pointes (S2.2.2-2)	<p>Torsades de pointes is polymorphic VT that occurs in the setting of a long-QT interval and is characterized by a waxing and waning QRS amplitude. It often has a long-short initiating sequence with a long coupling interval to the first VT beat and may present with salvos of NSVT. The twisting of the points, although characteristic, may not always be seen, especially if the episode is nonsustained or if only a limited number of leads are available. Torsades de pointes can result from bradycardia including high-grade AV block that leads to a long-short sequence initiating torsades de pointes.</p> 
Ventricular flutter (S2.2.2-2)	<p>A regular VA <math>\approx 300</math> bpm (cycle length: 200 ms) with a sinusoidal, monomorphic appearance; no isoelectric interval between successive QRS complexes.</p> 
Ventricular fibrillation (S2.2.2-2)	<p>Rapid, grossly irregular electrical activity with marked variability in electrocardiographic waveform, ventricular rate usually <math>&gt;300</math> bpm (cycle length: <math>&lt;200</math> ms).</p> 

*continued on the next page*

**TABLE 5** **Continued**

Term	Definition or Description
Sudden cardiac arrest (S2.2.2-2)	SCA is the sudden cessation of cardiac activity such that the victim becomes unresponsive, with either persisting gasping respirations or absence of any respiratory movements, and no signs of circulation as manifest by the absence of a perceptible pulse. An arrest is presumed to be of cardiac etiology unless it is known or likely to have been caused by trauma, drowning, respiratory failure or asphyxia, electrocution, drug overdose, or any other noncardiac cause.
Sudden cardiac death (S2.2.2-2)	Sudden and unexpected death occurring within an hour of the onset of symptoms, or occurring in patients found dead within 24 h of being asymptomatic and presumably due to a cardiac arrhythmia or hemodynamic catastrophe.
VT/VF storm (S2.2.2-3)	VT/VF storm (electrical storm or arrhythmic storm) refers to a state of cardiac electrical instability that is defined by ≥3 episodes of sustained VT, VF, or appropriate shocks from an ICD within 24 h.
Primary prevention ICD (S2.2.2-2)	ICD placement with the intention of preventing SCD in a patient who has not had sustained VT or SCA but who is at an increased risk for these events.
Secondary prevention ICD (S2.2.2-2)	ICD placement in a patient with prior SCA, sustained VT, or syncope caused by VA.
Structural heart disease*	This term encompasses IHD, all types of cardiomyopathy, valvular heart disease, and adult congenital heart disease.
Cardiac channelopathy (S2.2.2-4)	Arrhythmogenic disease due to a genetic abnormality that results in dysfunction of a cardiac ion channel (e.g., long-QT syndrome, catecholaminergic polymorphic VT).

\*The definition of this term may differ across publications. Refer to the entry for the definition used in this document.

AV indicates atrioventricular; ICD, implantable cardioverter-defibrillator; IHD, ischemic heart disease; NSVT, nonsustained ventricular tachycardia; SCA, sudden cardiac arrest; SCD, sudden cardiac death; VA, ventricular arrhythmia; VF, ventricular fibrillation; and VT, ventricular tachycardia.

nondihydropyridines calcium channel blockers, and some antiarrhythmic medications may relieve symptoms of palpitations (S2.2.2-18).

PVCs that occur during an exercise test are associated with a higher risk of death (S2.2.2-19). In 1 study, PVCs that occur during recovery are a stronger predictor of death than PVCs occurring only during exercise (S2.2.2-20). However, PVCs are common in trained athletes who have palpitations, in whom there does not appear to be increased risk of death based on studies of small numbers of athletes, at least in those without other cardiovascular abnormalities (S2.2.2-21,S2.2.2-22). Complex PVCs may not represent a benign finding in endurance athletes. An electrophysiological study may be needed to assess patients' arrhythmogenic risk (S2.2.2-22). Very frequent PVCs, >10,000 to 20,000 a day, can be associated with depressed LV function in some patients that is reversible with control of the PVCs, and has been referred to as PVC-induced cardiomyopathy (S2.2.2-23,S2.2.2-24). (See also Section 8.5. PVC-Induced Cardiomyopathy.) Very rarely, idiopathic PVCs from the outflow tract may trigger malignant VA in patients without structural heart disease (S2.2.2-25,S2.2.2-26).

### 2.1.2. VT and VF During ACS

Approximately half of patients with out-of-hospital cardiac arrest with the first rhythm identified as VF and who survive to hospital admission have evidence of acute MI (AMI) (S2.2.2-27). Of all out-of-hospital cardiac arrests, >50% will have significant coronary artery lesions on acute coronary angiography (S2.2.2-27). Of patients hospitalized with AMI, 5% to 10% have VF or sustained VT prior to hospital presentation, and another 5% will have VF or sustained VT after hospital arrival, most within 48

hours of admission. A study of patients with non-ST-elevation ACS who underwent cardiac catheterization within 48 hours found VT/VF in 7.6% of patients, with 60% of those events within 48 hours of admission (S2.2.2-28). Accelerated idioventricular rhythm is a common arrhythmia in patients with acute MI, including patients with ST-segment elevation MI undergoing primary percutaneous coronary intervention (PCI). Accelerated idioventricular rhythm is more closely related to the extent of infarction than to reperfusion itself (S2.2.2-29).

Sustained VA that occurs in the setting of an ACS is more often polymorphic VT or VF than monomorphic VT. Risk factors for VT/VF include prior history of hypertension, prior MI, ST-segment changes at presentation, and chronic obstructive pulmonary disease (S2.2.2-30). A nationwide Danish study found that 11.6% of patients with ST-segment elevation MI who underwent PCI had VF prior to the PCI, and that VF was associated with alcohol consumption, preinfarction angina, anterior infarct location, and complete coronary occlusion at the time of coronary angiography (S2.2.2-31). In a select group of patients undergoing primary PCI in a clinical trial, 5.7% developed sustained VT or VF, with two thirds of these events occurring prior to the end of the catheterization, and 90% within 48 hours from the procedure. VT or VF after primary PCI was associated with lower blood pressure, higher heart rate, poor coronary flow at the end of the procedure, and incomplete resolution of ST elevation (S2.2.2-32). Importantly, and in contrast to some earlier studies, VT or VF at any time was associated with a substantially higher risk of death within 90 days. Late VT or VF (after 48 hours of hospital presentation) was associated with a higher risk of death than early VT or VF (within 48 hours of hospital presentation) (S2.2.2-33).

### 2.1.3. Sustained VT and VF Not Associated With ACS

Patients with structural heart disease are at an increased risk for sustained VT and VF. Sustained VT that is not associated with an ACS is often monomorphic as it is usually due to scar-related reentry, but it may degenerate to VF ([S2.2.2-34](#)). The risk and predictors of VT in patients with structural heart disease depend on the type, severity, and duration of structural heart disease, increasing with the severity of ventricular dysfunction and the presence of symptomatic HF. Monomorphic VT occurring in the absence of structural heart disease is commonly referred to as idiopathic VT and is often due to an automatic focus in a characteristic location, giving rise to typical electrocardiographic appearances. Polymorphic VT and VF occurring in the absence of structural heart disease are rare and may be due to a cardiac channelopathy ([S2.2.2-35](#),[S2.2.2-36](#)), medication-induced long QT syndrome ([S2.2.2-36](#)), or they may be idiopathic ([S2.2.2-37](#),[S2.2.2-38](#)).

## 2.2. Sudden Cardiac Death

### 2.2.1. Incidence of SCD

SCA and its most common consequence, SCD, constitute major public health problems, accounting for approximately 50% of all cardiovascular deaths ([S2.2.2-1](#),[S2.2.2-39](#)), with at least 25% being first symptomatic cardiac events ([S2.2.2-1](#),[S2.2.2-40](#),[S2.2.2-41](#)). In addition, analyses of the magnitude of SCD are limited, in part because of the broad range of estimates of the risk based on different epidemiological methods ([S2.2.2-42](#)). During the past 20 to 30 years, SCD accounted for approximately 230,000 to 350,000 deaths per year in the United States, with a range of <170,000 to >450,000, depending on epidemiological methods, data sources, and inclusion criteria ([S2.2.2-41](#),[S2.2.2-43](#)). The lowest of these extremes came from national extrapolation of data from specific local programs, while the highest rates included noncardiac causes of sudden death such as pulmonary embolism or intracranial bleeding. The mid-range numbers were largely based on death certificate studies that required a code inclusive of ischemic heart disease.

The 2017 update of cardiovascular statistics from the AHA estimated the total annual burden of out-of-hospital cardiac arrest at 356,500 ([S2.2.2-44](#)). An additional 209,000 in-hospital cardiac arrests occur annually ([S2.2.2-45](#)). Among the out-of-hospital cardiac arrest group, approximately 357,000 events trigger emergency rescue response, with 97% occurring in adults >18 years of age.

The survival statistics for out-of-hospital cardiac arrest remain disappointing, with an estimated 10% overall survival rate ([S2.2.2-44](#)). Among the subgroup of 70% of out-of-hospital cardiac arrests that occur in the home, survival is 6%. The best reported outcomes are from locations with highly developed and publicly visible

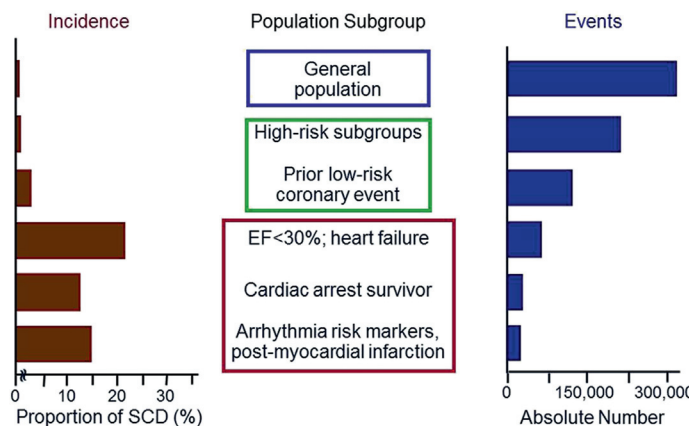
emergency rescue response, along with the combination of public location of cardiac arrest, bystander witnesses willing to provide CPR, first responders arriving quickly, shockable rhythm at initial contact, availability of automated external defibrillators (AEDs), and possibly a benefit from telecommunication-directed CPR ([S2.2.2-46](#),[S2.2.2-47](#)). Survival to hospital discharge after in-hospital cardiac arrests is estimated to be 24% ([S2.2.2-48](#)). In all settings, survival statistics appear to be better when rhythms recorded by responders are shockable (VF, pulseless VT), compared with pulseless electrical activity or asystole ([S2.2.2-49](#)). Although the apparent increase in the incidence of pulseless electrical activity or asystole could be due to the later arrival of medical care, the decrease in the incidence of shockable rhythm has also been attributed, in part, to improvements in diagnosis and treatment of structural heart disease ([S2.2.2-40](#)).

### 2.2.2. Population Subgroups and Risk Prediction

Risk prediction for SCA and SCD is complex. Risk analysis is divided into 2 general categories: population risk prediction and individual risk prediction ([S2.2.2-41](#),[S2.2.2-50](#)). Conventional epidemiological markers provide insight into probabilities for the development of ischemic heart disease within a general class of subjects, but adequately tested and validated profiles for SCA risk stratification of individuals in the general population do not presently exist. The challenge of defining SCA risk in individuals derives from a population model characterized by large numbers of events diluted into a very large denominator ([Figure 1](#)). The overall population can be subgrouped into categories based on integration of age, presence and extent of disease, and identification of small, high-risk subgroups within the large denominator general population.

Increasing age is a strong predictor of risk for SCA, but it is not linear. Risk in the general population, over time, beginning at 35 years of age has been estimated at 1 per 1000 population per year, increasing from a risk <1000 at the younger end of that spectrum to a higher risk in the elderly ([S2.2.2-41](#)). However, an analysis of lifetime risk of SCD, derived from the Framingham data, suggested that the incidence of SCD decreases in later years, especially in people >75 years of age ([S2.2.2-51](#)). The data also suggested that SCD is uniformly more common in men than in women at all age groups. In contrast, the population of children, adolescents, and young adults has an overall annual risk of 1 per 100,000, and there is somewhat a higher risk of SCD at the younger end of that age range ([S2.2.2-41](#)). An age-associated transition range, from the mid-20s to 35 to 40 years of age, is characterized by a steep increase in risk from that of the adolescent group to the middle-aged group, corresponding to the emergence of ischemic heart disease.

**FIGURE 1A SCD Incidence and Total Events (S2.2.2-1)**



EF indicates ejection fraction; and SCD, sudden cardiac death.

Although ischemic heart disease remains the most common underlying substrate associated with SCD, the incidence of ischemic heart disease-related SCD appears to be decreasing (S2.2.2-52), with various forms of cardiomyopathy associated with myocardial fibrosis and LV hypertrophy increasing (S2.2.2-53). In addition, a trend over time has suggested that out-of-hospital cardiac arrest patients who are admitted alive to a hospital are becoming more likely to have high-risk clinical profiles, as opposed to manifest disease (S2.2.2-54). The younger population—children, adolescents, and young adults—is affected by a series of disorders that manifest earlier in life, including the genetic structural disorders and cardiac channelopathies, myocarditis, congenital heart disease, and other rare disorders (S2.2.2-43). During the transition range, from the mid-20s to the mid-30s, causes of SCA and SCD include a lower proportion of inherited diseases and increasing proportion of ischemic heart disease (>40% of cases) (S2.2.2-43).

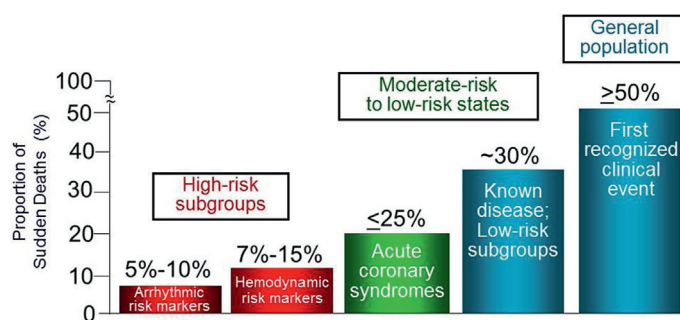
Despite the small progress that has been made in risk prediction of SCA and SCD, the greatest challenge is to identify the relatively small, high-risk subgroups concealed within the large general population who have no identified disease but are at risk of SCA as their first cardiac event (Figure 1) (S2.2.2-50).

### 3. MECHANISMS OF VA

#### 3.1. Cellular Mechanisms and Substrates

Mechanisms of VA include enhanced normal automaticity, abnormal automaticity, triggered activity induced by early or late afterdepolarizations, and reentry (S3.4-1–S3.4-3). Reentry requires a trigger to initiate the arrhythmia and a substrate to sustain it. The trigger may be a PVC, which may be due to automaticity. The substrate may be structural remodeling secondary to an underlying disease process, and often includes a scar secondary to a prior MI or surgical repair, or patchy

**FIGURE 1B SCD and Clinical Subsets (S2.2.2-1)**



SCD indicates sudden cardiac death.

fibrosis in the setting of cardiomyopathy or hypertrophy. Changes in ion channel or transporter function and/or expression and cell to cell coupling secondary to the underlying pathology may alter the initiation or propagation of the cardiac action potential. The electrophysiological substrate is dynamically influenced by a variety of factors including cardiac metabolism, electrolytes, signaling pathways and autonomic effects. Enhanced automaticity or abnormal automaticity causing VA may arise from subordinate pacemaker cells in the His-Purkinje system or ventricular myocardium.

### 3.2. Automaticity

Normal automaticity results from phase 4 spontaneous depolarization of the transmembrane action potential arising from a normal resting potential, reaching threshold and initiating an action potential ([S3.4-1,S3.4-3](#)). An initiating current ( $I_f$ ) is responsible for spontaneous phase 4 depolarization in the sinus node. The rate is determined by the integration of the maximum diastolic potential at the end of repolarization, the slope of phase 4 depolarization, and the threshold potential. In contrast, abnormal automaticity arises from a partially depolarized membrane potential that is usually close to the activation potential for calcium channels in the cell membrane ([S3.4-1,S3.4-3](#)). In the acute phase of an MI or during transient ischemia, increased extracellular potassium causes partial depolarization of the resting membrane potential creating injury currents between the infarcted/ischemic tissue and healthy myocardium. These injury currents may initiate spontaneous activity. In ischemia, abnormal automaticity may occur in both ventricular myocytes and Purkinje fibers, and may also enhance normal automaticity in Purkinje fibers in the ischemic zone.

### 3.3. Triggered Activity

Early afterdepolarizations occur during late phase 2 or early phase 3 of the action potential ([S3.4-3–S3.4-5](#)), usually in the setting of action potential prolongation due to an increase in inward currents (the late sodium current, the inward calcium current or the sodium calcium exchange current) or a decrease in repolarizing potassium currents. Under these conditions, early afterdepolarizations may be initiated when reactivation of the inward L-type calcium channel occurs before the membrane has returned to a more negative potential than that required for calcium channel reactivation. Spontaneous calcium release from the sarcoplasmic reticulum may also result in activation of a depolarizing sodium/calcium exchange current. Early afterdepolarizations are the trigger for torsades de pointes VT associated with QT prolongation either induced by medications or other acquired factors or due to mutations of ion channels causing the long QT syndrome. In these cases, it is possible that the early

afterdepolarization/triggered activity sequence is the trigger that culminates in polymorphic VT/VF.

Delayed afterdepolarizations occur after complete membrane repolarization and develop under conditions of intracellular calcium overload. Factors contributing to elevated intracellular calcium load include tachycardia, catecholamines, hypokalemia, digoxin toxicity, cardiac hypertrophy, and HF ([S3.4-6,S3.4-7](#)). Elevated sarcoplasmic calcium content or increased sensitivity of the ryanodine receptor can initiate spontaneous calcium release, which activates a transient inward current driven predominantly by the sodium-calcium exchange current. If the membrane depolarization is sufficiently large, the inward sodium current is activated resulting in a triggered action potential. Delayed afterdepolarizations are the underlying mechanism for VT in the setting of digoxin toxicity, catecholaminergic polymorphic VT, and idiopathic outflow tract VA. Delayed afterdepolarizations are also considered to be an important trigger of VA in the setting of HF. Purkinje cells are more susceptible to spontaneous sarcoplasmic reticulum calcium release than ventricular myocytes suggesting that delayed afterdepolarizations may be an important mechanism for some Purkinje fiber-related VA ([S3.4-3,S3.4-8,S3.4-9](#)).

### 3.4. Reentry

Reentry is the underlying mechanism for most sustained VA in the presence of structural heart disease ([S3.4-1–S3.4-3,S3.4-10–S3.4-12](#)). Reentry may occur around a fixed anatomical obstacle, such as scar after an MI or surgically repaired congenital heart disease. In this setting, an excitable gap separates the excitation wavefront from its tail of refractoriness. The existence of structural reentrant substrates provide the rationale for VT ablation in scar-related VTs ([S3.4-11,S3.4-12](#)).

Functional reentry around areas of functional block without anatomical obstacles can also occur. Two main models of functional reentry have been proposed ([S3.4-2, S3.4-3](#)). The leading circle model has a functionally refractory core and no excitable gap. Spiral wave reentry is driven by a rotor with a curved wavefront and wavetail pivoting around an excitable but unexcited core. There remains much debate about the precise mechanism(s) of VF (rotor versus multiple wavelet reentry). Both mechanisms may be operational in different phases of VF ([S3.4-10](#)).

Phase 2 reentry may occur due to heterogeneity of ventricular repolarization. Electrotonic currents may flow from endocardial sites with longer action potential durations to the epicardium with shorter action potential durations which can result in reexcitation when these sites have recovered from refractoriness. This is believed to be one potential mechanism of VT/VF in Brugada syndrome ([S3.4-3](#)) and may also be operative during ischemia.

## 4. GENERAL EVALUATION OF PATIENTS WITH DOCUMENTED OR SUSPECTED VA

### 4.1. History and Physical Examination

#### Recommendation for Syncope\*

Referenced studies that support the recommendation are summarized in [Online Data Supplement 1](#).

COR	LOE	RECOMMENDATION
I	B-NR	1. Patients presenting with syncope for which VA is documented, or thought to be a likely cause, should be hospitalized for evaluation, monitoring, and management ( <a href="#">S4.1-1–S4.1-4</a> ).

\*This section covers practices that are well accepted, and a new recommendation was determined to only be warranted for syncope.

**Table 6**

#### Synopsis

VA can produce a wide spectrum of symptoms, and the severity of symptoms does not necessarily reflect the extent of structural heart disease or the potential risk of

SCD. Symptoms of VA include palpitations, either skipped or extra beats or sustained palpitations, shortness of breath, chest pain, dizziness, near syncope, and syncope ([S4.1-5,S4.1-6](#)). Palpitations may correlate with VA but are frequently reported during normal rhythm ([S4.1-7](#)). The

**TABLE 6** Important Considerations in the Evaluation of Patients With Known or Suspected VA

Component	Assessment and Findings Relevant for VA and/or SCD Risk
History	<ol style="list-style-type: none"><li>1. Symptoms/events related to arrhythmia: Palpitations, lightheadedness, syncope, dyspnea, chest pain, cardiac arrest</li><li>2. Symptoms related to underlying heart disease: Dyspnea at rest or on exertion, orthopnea, paroxysmal nocturnal dyspnea, chest pain, edema</li><li>3. Precipitating factors: Exercise, emotional stress</li><li>4. Known heart disease: Coronary, valvular (e.g., mitral valve prolapse), congenital heart disease, other</li><li>5. Risk factors for heart disease: Hypertension, diabetes mellitus, hyperlipidemia, and smoking</li><li>6. Medications:<ul style="list-style-type: none"><li>■ Antiarrhythmic medications</li><li>■ Other medications with potential for QT prolongation and torsades de pointes</li><li>■ Medications with potential to provoke or aggravate VA<ul style="list-style-type: none"><li>&gt; Stimulants including cocaine and amphetamines</li><li>&gt; Supplements including anabolic steroids</li></ul></li><li>■ Medication-medication interaction that could cause QT prolongation and torsades de pointes</li></ul></li><li>7. Past medical history:<ul style="list-style-type: none"><li>■ Thyroid disease</li><li>■ Acute kidney injury, chronic kidney disease, or electrolyte abnormalities</li><li>■ Stroke or embolic events</li><li>■ Lung disease</li><li>■ Epilepsy (arrhythmic syncope can be misdiagnosed as epilepsy)</li><li>■ Alcohol or illicit drug use</li><li>■ Use of over-the-counter medications that could cause QT prolongation and torsades de pointes</li><li>■ Unexplained motor vehicle crashes</li></ul></li></ol>
Family History	<ol style="list-style-type: none"><li>1. SCD, SCA, or unexplained drowning in a first-degree relative</li><li>2. SIDS or repetitive spontaneous pregnancy losses given their potential association with cardiac channelopathies</li><li>3. Heart disease<ul style="list-style-type: none"><li>■ IHD</li><li>■ Cardiomyopathy: Hypertrophic, dilated, ARVC</li><li>■ Congenital heart disease</li><li>■ Cardiac channelopathies: Long QT, Brugada, Short QT, CPVT</li><li>■ Arrhythmias</li><li>■ Conduction disorders, pacemakers/ICDs</li></ul></li><li>4. Neuromuscular disease associated with cardiomyopathies<ul style="list-style-type: none"><li>■ Muscular dystrophy</li></ul></li><li>5. Epilepsy</li></ol>
Examination	<ol style="list-style-type: none"><li>1. Heart rate and regularity, blood pressure</li><li>2. Jugular venous pressure</li><li>3. Murmurs</li><li>4. Pulses and bruits</li><li>5. Edema</li><li>6. Sternotomy scars</li></ol>

ARVC indicates arrhythmogenic right ventricular cardiomyopathy; CPVT, catecholaminergic polymorphic ventricular tachycardia; IHD, ischemic heart disease; SCA, sudden cardiac arrest; SCD, sudden cardiac death; SIDS, sudden infant death syndrome; and VA, ventricular arrhythmia.

differential diagnosis of exercise intolerance, chest pain, dyspnea, presyncope, and syncope includes VA but also includes other etiologies. Nonetheless, more dramatic symptoms, particularly in patients with known or discovered structural or electrical heart disease should prompt focused investigation for possible association with VA (**Table 6**).

The elucidation of precipitating factors, such as exertional or emotional stress, concurrent medications or illness, and alleviating factors is important. The presence of a family history of SCD, ischemic heart disease, valvular heart disease, nonischemic cardiomyopathy (NICM), or HF raises concern for the presence of one of these disorders associated with VA. Obtaining a complete medication history is important. Various antiarrhythmic and other medications can cause QT prolongation and torsades de pointes ([www.crediblemeds.org](http://www.crediblemeds.org)) (**S4.1-8**); some medications can also induce Brugada type I electrocardiographic pattern and VF ([www.brugadadrugs.org](http://www.brugadadrugs.org)) (**S4.1-9,S4.1-10**).

Patients with bigeminy and trigeminy can present with effective bradycardia, an apical-radial pulse deficit and relative hypertension with a wide pulse pressure. Effective bradycardia from PVCs can result in inaccurate estimation of the heart rate. Although premature beats on auscultation of the heart can be detected, the physical examination is focused largely on finding evidence of structural heart disease. Carotid bruits or diminished peripheral pulses may be indicators of atherosclerotic disease associated with ischemic heart disease. Jugular venous distention, rales, gallops, and peripheral edema provide evidence of HF. Auscultation may reveal cardiac murmurs consistent with valvular heart disease, such as aortic stenosis or mitral regurgitation, and may be associated with HF and VA. A midsystolic click may indicate mitral valve prolapse that can be associated with VA (**S4.1-11–S4.1-13**). Many VA are asymptomatic and detected only on an ECG or telemetry. Such cases highlight the need to search for evidence of underlying heart disease.

## Recommendation-Specific Supportive Text

- Rapid, sustained VT may result in syncope secondary to marked reduction in cardiac output, followed by spontaneous recovery if VT terminates, or SCA if VT persists and is not treated promptly. Syncope or SCA may be the first manifestation of structural or electrical heart disease (**S4.1-14**), and some SCA victims have preceding “sentinel” syncope episodes (**S4.1-15**). Syncope, or its forewarnings of dizziness, lightheadedness, or near-syncope, may constitute a risk factor for SCA and SCD (**S4.1-2**). The initial evaluation at any age focuses on detection or exclusion of heart disease. Syncope during exercise should prompt thorough evaluation to rule out cardiac causes. Cardiac evaluation with echocardiography, ambulatory monitoring, and exercise testing may be warranted depending on the clinical information elicited (**S4.1-3,S4.1-4**). Cardiac causes of syncope include sustained VT, high-grade atrioventricular block or severe sinus bradycardia or prolonged sinus pauses, supraventricular tachycardia (SVT), malfunction of pacemakers, VA from cardiac channelopathies or structural heart disease syndromes, such as hypertrophic cardiomyopathy (HCM) or congenital heart disease (**S4.1-3,S4.1-4,S4.1-16**). Cardiac channelopathies and HCM are particularly important to consider in adolescents and young adults. Arrhythmic causes of syncope are often associated with very short periods of premonitory symptoms, or palpitations, and known preexisting heart disease, especially a history of a low LVEF or HF (**S4.1-1**). Among nonarrhythmic cardiac causes, considerations should include myocardial ischemia, severe aortic stenosis, HCM, HF, and prosthetic valve malfunction, pulmonary embolism, medications, and illicit drug use (**S4.1-3**).

### 4.2. Noninvasive Evaluation

#### 4.2.1. 12-lead ECG and Exercise Testing

#### Recommendations for 12-lead ECG and Exercise Testing

References studies that support the recommendations are summarized in [Online Data Supplement 2](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with sustained, hemodynamically stable, wide complex tachycardia, a 12-lead ECG during tachycardia should be obtained ( <b>S4.2.1-1–S4.2.1-3</b> ).
I	B-NR	2. In patients with VA symptoms associated with exertion, suspected ischemic heart disease, or catecholaminergic polymorphic ventricular tachycardia, exercise treadmill testing is useful to assess for exercise-induced VA ( <b>S4.2.1-4,S4.2.1-5</b> ).
I	B-NR	3. In patients with suspected or documented VA, a 12-lead ECG should be obtained in sinus rhythm to look for evidence of heart disease ( <b>S4.2.1-6</b> ).

### Recommendation-Specific Supportive Text

1. A 12-lead ECG during tachycardia is the first diagnostic test that should be done in any patient found to be in a stable wide QRS complex tachycardia on a monitor. VT is the diagnosis in most adults with wide complex tachycardia and underlying structural heart disease ([S4.2.1-3](#)). Criteria that support a diagnosis of VT include AV dissociation, a QRS complex >0.14 s, monophasic R wave in aVR, specific QRS morphologies (e.g., positively or negatively concordant QRS complexes in the precordial leads), the absence of an RS complex in all precordial leads and an RS interval >100 ms in at least 1 precordial lead ([S4.2.1-2](#)). Exceptions occur, particularly in patients with advanced heart disease and with the use of certain antiarrhythmic medications ([S4.2.1-1](#)). For patients with preexisting bundle branch block, comparison of the QRS morphology during sinus rhythm with that during wide complex tachycardia is often relevant.
2. For exertion-related arrhythmic symptoms, exercise in a monitored setting may reproduce the symptoms and/or the related arrhythmia, allowing for diagnosis. Exercise testing is particularly important when catecholaminergic polymorphic ventricular tachycardia is a

possibility. However, exertion-related symptoms and findings may not be reliably reproducible with exercise testing, and long-term electrocardiographic monitoring with external or implantable recorders may be necessary.

3. A 12-lead ECG may indicate the presence of structural heart disease such as prior MI or chamber enlargement that would increase the likelihood that a patient's symptoms might be due to VA, or it may provide evidence of the underlying substrate for documented VA. An ECG may also reveal evidence of inherited arrhythmia disorders, such as long QT syndrome, Brugada syndrome, and arrhythmogenic right ventricular cardiomyopathy. In patients with structural heart disease, QRS duration and the presence of conduction abnormalities provide prognostic information ([S4.2.1-7](#)–[S4.2.1-14](#)). Data on the use of microvolt T wave alternans and the signal averaged ECG are inconclusive, as such these tests are not routinely used in clinical practice ([S4.2.1-15](#)–[S4.2.1-19](#)); the one exception is the potential use of signal averaged ECG in patients with arrhythmogenic right ventricular cardiomyopathy (see [Section 7.3](#)).

### 4.2.2. Ambulatory Electrocardiography

#### Recommendation for Ambulatory Electrocardiography

Referenced studies that support the recommendation are summarized in [Online Data Supplement 3 and 4](#).

COR	LOE	RECOMMENDATION
I	B-NR	1. Ambulatory electrocardiographic monitoring is useful to evaluate whether symptoms, including palpitations, presyncope, or syncope, are caused by VA ( <a href="#">S4.2.2-1</a> – <a href="#">S4.2.2-4</a> ).

### Recommendation-Specific Supportive Text

1. Ambulatory electrocardiographic monitoring is often used to assess the effectiveness of treatments to suppress arrhythmias, but more robust data are needed on the clinical use of this practice. Continuous or intermittent ambulatory electrocardiographic recording with a Holter monitor or an event recorder is helpful in diagnosing suspected arrhythmias, establishing their frequency, relating them to symptoms, and assessing the response to therapy. Although the yield of these tests is relatively low, VT is occasionally documented ([S4.2.2-4](#)). A 24-hour continuous Holter recording is appropriate when symptoms occur at least once a day or when quantitation of PVCs/NSVT is desired to assess possible VA-related depressed ventricular function. For sporadic symptoms, event or “looping” monitors are more appropriate because they can be activated over extended periods of time and

increase diagnostic yield ([S4.2.2-2](#),[S4.2.2-3](#)). Adhesive patch electrocardiographic monitors can record for weeks and allow for continuous short-term 1-lead monitoring and patient activation for symptoms. Studies have shown satisfactory patient compliance, and arrhythmia detection; however, with some monitors, detected arrhythmias are not discovered until the patch is returned for analysis ([S4.2.2-1](#),[S4.2.2-4](#)). Serial evaluations with exercise testing and/or 24-hour ambulatory monitoring are also used to assess rhythm burden and response of VA to therapy. Notably, implantable monitors are covered in [Section 4.2.3](#). Importantly, when the suspicion of VA in a patient is high, outpatient ambulatory monitoring is inappropriate as prompt diagnosis and prevention of VA are warranted. It is important to accurately correlate the symptoms with the arrhythmias detected by ambulatory ECG monitoring.

#### 4.2.3. Implanted Cardiac Monitors

##### Recommendation for Implanted Cardiac Monitors

Referenced studies that support the recommendation are summarized in [Online Data Supplement 5](#).

COR	LOE	RECOMMENDATION
IIa	B-R	1. In patients with sporadic symptoms (including syncope) suspected to be related to VA, implanted cardiac monitors can be useful (S4.2.3-1–S4.2.3-4).

##### Recommendation-Specific Supportive Text

1. Implanted cardiac monitors provide continuous rhythm monitoring and stored recordings of electrograms based on patient activation or preset parameters, allowing a prolonged monitoring period of a few years. These devices require a minor invasive procedure with local anesthesia for implantation. In patients with sporadic symptoms, including syncope, implantable recorders are useful in diagnosing serious tachyarrhythmias (including VA) and bradyarrhythmias (S4.2.3-2–S4.2.3-4). They are generally reserved for patients in whom other ambulatory monitoring is nonrevealing due to the infrequency of events. A 25% added yield in diagnosis has been described after an unrevealing external ambulatory monitor (S4.2.3-5). In a study of patients with syncope, the implantable monitor had a greater diagnostic yield than “conventional” testing with external monitoring, tilt table testing and electrophysiological study (S4.2.3-2). A systematic review in patients with syncope concluded that use of these devices provide a higher rate of diagnosis and a trend toward reduction in syncope relapse after diagnosis, as compared with conventional management (S4.2.3-3). A prospective study of patients after MI, with LVEF <40%, demonstrated NSVT (>16 beats long) in 13%, VT (>30 s) in 3% and VF in 3% of patients (S4.2.3-1). It is important to accurately correlate the symptoms with the arrhythmias detected by implanted cardiac monitors.

#### 4.2.4. Noninvasive Cardiac Imaging

##### Recommendations for Noninvasive Cardiac Imaging

Referenced studies that support the recommendations are summarized in [Online Data Supplement 6](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with known or suspected VA that may be associated with underlying structural heart disease or a risk of SCA, echocardiography is recommended for evaluation of cardiac structure and function (S4.2.4-1,S4.2.4-2).
IIa	C-EO	2. In patients presenting with VA who are suspected of having structural heart disease, cardiac magnetic resonance imaging (MRI) or computed tomography (CT) can be useful to detect and characterize underlying structural heart disease.

including anomalous coronary origins. Cardiac MRI can be useful in the evaluation for myocardial scar and infiltrative processes evident as late gadolinium enhancement ([S4.2.4-5–S4.2.4-9](#)). Cardiac MRI also provides high-quality assessment of LV and RV

function, size, and degree of fibrosis and is particularly useful in arrhythmogenic right ventricular cardiomyopathy and HCM.

#### 4.2.5. Biomarkers

##### Recommendation for Biomarkers

Referenced studies that support the recommendation are summarized in [Online Data Supplement 7](#).

COR	LOE	RECOMMENDATION
IIa	B-NR	<ol style="list-style-type: none"><li>1. In patients with structural heart disease, measurement of natriuretic peptides (BNP or N-terminal pro-BNP) can be useful by adding prognostic information to standard risk factors for predicting SCD or SCA (<a href="#">S4.2.5-1–S4.2.5-4</a>).</li></ol>

##### Recommendation-Specific Supportive Text

1. Elevated levels of natriuretic peptides—B-type natriuretic peptide (BNP) or N-terminal pro-BNP—are associated with increased risk of SCA and appropriate ICD therapies, even after adjustment of LVEF and other risk factors ([S4.2.5-1–S4.2.5-4](#)). These biomarkers are also predictive of nonsudden cardiovascular mortality and thus are not specific to SCD risk alone. Natriuretic peptides have also been evaluated for predicting SCD in the general population ([S4.2.5-5,S4.2.5-6](#)). In the Nurses' Health Study, an elevated N-terminal pro-BNP was an independent risk marker for SCD in presumably healthy women ([S4.2.5-5](#)). In an older adult population, higher baseline levels of N-terminal pro-BNP

were associated with SCD over a 16-year follow-up period ([S4.2.5-6](#)). These biomarkers may also have a potential role in facilitating the identification of individuals at increased risk of SCD and VA in the general population, particularly in those at intermediate or high risk of ischemic heart disease, but further studies are needed. Use of biomarkers has not been shown to be useful for selecting patients for ICDs. A study of 4431 patients found high-sensitivity troponin to be only weakly predictive of SCD ([S4.2.5-7](#)). However, there are no data on whether high-sensitivity troponin can improve the current SCD prediction algorithms.

#### 4.2.6. Genetic Considerations in Arrhythmia Syndromes

##### Recommendation for Genetic Counselling\*

COR	LOE	RECOMMENDATION
I	C-EO	<ol style="list-style-type: none"><li>1. In patients and family members in whom genetic testing for risk stratification for SCA or SCD is recommended, genetic counseling is beneficial.</li></ol>

\*Please refer to section 7.9 for disease-specific recommendations.

##### Synopsis

The diagnosis of most inherited arrhythmia syndromes is based on clinical features and family history. The availability of genetic testing for inherited arrhythmia syndromes can: 1) provide opportunity to confirm a suspected clinical diagnosis and sometimes provide prognostic information for the proband and 2) offer cascade screening of potentially affected family members when a disease-causing mutation is identified in the proband. The yield of genetic testing varies by disease. The verification of pathogenicity of suspected mutations is an evolving field, and exome sequencing has identified an increasing number of variants of uncertain significance in the general

population ([S4.2.6-1–S4.2.6-5](#)). Genotyping can have therapeutic implications for some arrhythmogenic phenotypes such as long QT syndrome and Fabry's disease ([S4.2.6-6–S4.2.6-9](#)), where a monogenic pathogenic mutation has been clearly identified, the risk to mutation positive individuals has been extensively studied, and effective therapy relevant to the mutation can be instituted. In other diseases, such as Brugada syndrome, the role of a clear monogenic disease-causing mutation is less certain, and the genotype does not provide therapeutic or prognostic information for the proband ([S4.2.6-5,S4.2.6-10–S4.2.6-12](#)). In arrhythmogenic right ventricular cardiomyopathy, some desmosomal mutation positive

individuals do not develop disease, indicating that additional mutations and environmental interactions likely influence the clinical development of disease ([S4.2.6-13–S4.2.6-16](#)). Importantly, the absence of an identified disease-causing genetic mutation does not exclude the presence of disease, and as such, ongoing monitoring and decision-making are done based on the clinical phenotype. Genotyping is frequently most useful when a pathogenic mutation is identified in the proband, such that screening can be applied to relatives who are in a preclinical phase, allowing institution of lifestyle changes, therapy, or ongoing monitoring for those who are gene mutation positive ([S4.2.6-7](#)). Refer to [Section 7.9](#) for disease-specific recommendations.

In young patients (<40 years of age) without structural heart disease who have unexplained cardiac arrest, unexplained near drowning, or recurrent exertional syncope, genetic testing may be important to identify an inherited arrhythmia syndrome as a likely cause ([S4.2.6-17–S4.2.6-23](#)).

#### Recommendation-Specific Supportive Text

1. The decision to proceed with genetic testing requires discussion regarding the clinical use of genetic

information to be obtained for both the proband and family members, as well as consideration of the important psychological, financial, employment, disability, and life insurance implications of positive genotyping ([S4.2.6-17,S4.2.6-18,S4.2.6-20,S4.2.6-24](#)). Balancing privacy of health care information for the proband with the “right to know” for family members, and the ability to provide appropriate communication of information to all potentially affected family members can be challenging on many levels, including family dynamics, geographic proximity, and access to health care ([S4.2.6-25](#)). For these reasons, genetic counseling generally occurs before proceeding with genetic testing, and, from a patient’s perspective, is optimally provided by genetic counselors, if available, in collaboration with physicians ([S4.2.6-26,S4.2.6-27](#)). A combined approach of genetic counseling with medical guidance may appropriately balance the decision as to whether genetic testing would be beneficial on an individual basis.

#### 4.3. Invasive Testing

##### 4.3.1. Invasive Cardiac Imaging: Cardiac Catheterization or CT Angiography

#### Recommendation for Invasive Imaging: Cardiac Catheterization

COR	LOE	RECOMMENDATION
I	C-EO	1. In patients who have recovered from unexplained SCA, CT or invasive coronary angiography is useful to confirm the presence or absence of ischemic heart disease and guide decisions for myocardial revascularization.

#### Recommendation-Specific Supportive Text

1. Although randomized studies are unavailable, coronary angiography has an important role in establishing or excluding the presence of significant obstructive ischemic heart disease in patients with SCA or those with life-threatening VA ([S4.3.1-1–S4.3.1-4](#)). Recurrent polymorphic VT or VF can be due to ongoing myocardial ischemia that resolves with coronary revascularization. Presence of ST-elevation on resuscitation or early postresuscitation ECG suggests ischemia and

potential ACS warranting urgent angiography and revascularization ([S4.3.1-5](#)). ST-elevation can also result from coronary spasm or DC shocks. The absence of ST-elevation after cardiac arrest does not exclude obstructive or thrombotic coronary lesions. A coronary angiogram may not be warranted if a nonischemic cause of SCA is established. Coronary and CT angiography also have an important role excluding the presence of anomalous origin of the coronary arteries that may cause SCD.

#### 4.3.2. Electrophysiological Study for VA

##### Recommendations for Electrophysiological Study

References that support the recommendations are summarized in [Online Data Supplement 8 and 9](#).

COR	LOE	RECOMMENDATIONS
IIa	B-R	1. In patients with ischemic cardiomyopathy, NICM, or adult congenital heart disease who have syncope or other VA symptoms and who do not meet indications for a primary prevention ICD, an electrophysiological study can be useful for assessing the risk of sustained VT ( <a href="#">S4.3.2-1–S4.3.2-7</a> ).
III: No Benefit	B-R	2. In patients who meet criteria for ICD implantation, an electrophysiological study for the sole reason of inducing VA is not indicated for risk stratification ( <a href="#">S4.3.2-8–S4.3.2-11</a> ).
III: No Benefit	B-NR	3. An electrophysiological study is not recommended for risk stratification for VA in the setting of long QT syndrome, catecholaminergic polymorphic ventricular tachycardia, short QT syndrome, or early repolarization syndromes ( <a href="#">S4.3.2-12–S4.3.2-16</a> ).

##### Synopsis

Electrophysiological study can be used to induce sustained VA in patients with known or suspected VA. With the advent of the ICD and its proven benefit in the primary and secondary prevention of SCD, there are fewer indications for programmed stimulation to provoke VA. Patients with HF and LVEF  $\leq 35\%$  generally will have an indication for an ICD and specific induction of VT/VF before implantation is not necessary. Patients with LVEF  $>35\%$  and unexplained syncope or near-syncope may benefit from an electrophysiological study to determine if VT/VF is the cause of symptoms and to guide further therapy. Induction of VT/VF is often attempted before catheter ablation of the arrhythmia substrate to guide the procedure and to determine the success of the intervention after ablation is performed. An electrophysiological study can be used to determine the mechanism of a wide complex tachycardia. See [Sections 7.3, 7.4, 7.6, 7.9.1.3, and 10.8](#) for recommendations regarding electrophysiological study for specific disease states.

##### Recommendation-Specific Supportive Text

1. A study of electrophysiological testing in patients with symptomatic NICM found inducible VT/VF in 28% of patients which was associated with a higher rate of ICD events during follow-up ([S4.3.2-17](#)). In a prospective cohort of 180 patients with ischemic or NICM and syncope, induction of VT or VF at electrophysiological study correlated with cardiac mortality only in patients with ischemic heart disease. In patients with NICM, cardiac mortality correlated with LVEF but not with inducibility on electrophysiological study ([S4.3.2-18](#)).
2. In patients who meet criteria for ICD implantation (i.e., HF and LVEF  $\leq 35\%$ ), data do not support the routine use of electrophysiological study solely for risk

stratification, as such patients have been shown to derive survival benefit from the ICD ([S4.3.2-8–S4.3.2-11](#)). An electrophysiological study may be helpful, however, in selected patients suspected to have pre-excitation or supraventricular arrhythmias as the cause of symptoms or wide complex tachycardias that warrant definitive diagnosis and management. SVT leading to VT/VF or aberrantly conducted SVT may also be suspected in younger patients or those with a preserved LVEF. Induction of SVT and ablation may then be curative, with no need for an ICD. In such cases, failure to induce VT/VF after elimination of the substrate for SVT would be expected.

3. Risk stratification for channelopathies is generally made on the basis of symptoms, the ECG ([S4.3.2-13, S4.3.2-19–S4.3.2-24](#)), exercise treadmill testing ([S4.3.2-25–S4.3.2-27](#)), and the results of genetic testing ([S4.3.2-28–S4.3.2-32](#)). The electrophysiological study (i.e., programmed ventricular stimulation) does not have prognostic value for risk stratification in patients with these cardiac channelopathies ([S4.3.2-12–S4.3.2-15](#)).

## 5. THERAPIES FOR TREATMENT OR PREVENTION OF VA

### 5.1. Medication Therapy

With the exception of beta blockers (e.g., metoprolol succinate, carvedilol), there is no evidence from RCTs that antiarrhythmic medications for VA improve survival when given for the primary or secondary prevention of SCD. However, the use of these medications is essential in some patients to control arrhythmias and improve symptoms. Medication use for VA is discussed, and any recommendations are listed, in subsequent sections. Further, medication-induced proarrhythmia is addressed in [Section 10.7](#).

Antiarrhythmic medications are often categorized by the Vaughan Williams 4-level schema (class I: fast sodium channel blockers; class II: beta blockers; class III: repolarization potassium current blockers; class IV: non-dihydropyridines calcium channel blockers) (S5.1.5.2-1). This system does not address the complexities in antiarrhythmic medications, since nearly every agent has multiple effects. Table 7 shows uses, electrophysiological effects, pharmacological effects, and common adverse effects of antiarrhythmic medications.

#### 5.1.1. Medications With Prominent Sodium Channel Blockade

Except in specific circumstances, sodium channel blockers (Vaughn-Williams class I agents) have a limited role in the prevention of VT/SCD; this is based on a lack of survival benefit and increased mortality observed during chronic therapy in patients with ischemic heart disease (see Section 10.7). Specific circumstances where sodium channel blockers have been used to treat VT/SCA include: intravenous lidocaine for patients with refractory VT/cardiac arrest (especially witnessed) (S5.1.5.2-3); oral mexiletine for congenital long QT syndrome (S5.1.5.2-4); quinidine for patients with Brugada syndrome; and flecainide for patients with catecholaminergic polymorphic ventricular tachycardia (S5.1.5.2-5). These medications could also be used in ICD patients with drug- and ablation-refractory VT.

One newer medication of potential benefit, based on very limited data, is ranolazine. This medication, developed and FDA-approved as an antianginal agent, provides relatively specific late sodium channel current blockade in addition to less potent blockade of the phase 3 repolarizing potassium current; that is, the rapid delayed rectifier potassium current; IKr. The potential for clinical antiarrhythmic efficacy is supported by basic studies and experimental models (S5.1.5.2-6). Clinical data are scant. In a study of 12 patients, ranolazine reduced ICD shocks in otherwise medication-resistant VT/VF in 11 patients (S5.1.5.2-7). In MERLIN TIMI-36 (Metabolic Efficiency With Ranolazine for Less Ischemia in Non-ST-Elevation Acute Coronary Syndromes-Thrombolysis In Myocardial Infarction 36), ranolazine did not reduce SCD but did reduce VT in the first few days after a non-ST-segment elevation ACS (S5.1.5.2-8). In 1 RCT, high-risk ICD patients with ischemic or NICM were randomly assigned to ranolazine 1000 mg twice a day versus placebo (S5.1.5.2-9). High risk was defined as: 1) having a primary prevention ICD without a history of documented VT/VF and with one of the following conditions: BUN  $\geq 26$  mg/dL, QRS  $>120$  msec, atrial fibrillation, or NSVT or  $>500$  VPBs on 24-hour Holter recording; 2) having a primary prevention ICD with a history of documented VT/VF appropriately treated with ICD therapy or untreated NSVT; or 3) having a secondary prevention ICD after documented VT/VF or

cardiac arrest. Ranolazine did not significantly reduce the primary endpoint of VT/VF requiring appropriate ICD therapy or death. In a prespecified secondary analysis, ranolazine was associated with a significant reduction in VT events treated with anti-tachycardia pacing (S5.1.5.2-9).

#### 5.1.2. Beta Blockers

Because of their excellent safety profile and effectiveness in treating VA and reducing the risk of SCD, beta blockers are often first-line antiarrhythmic therapy (S5.1.5.2-10, S5.1.5.2-11). Their antiarrhythmic efficacy is related to the effects of adrenergic-receptor blockade on sympathetically mediated triggering mechanisms, slowing of the sinus rate, and possibly inhibition of excess calcium release by the ryanodine receptor (S5.1.5.2-12).

Beta blockers reduce all-cause mortality and SCD in patients with HF with reduced EF (HFrEF) (S5.1.5.2-13–S5.1.5.2-15). Although beta blockers have long been proven to reduce mortality after MI (S5.1.5.2-16), registry data confirm that early beta blocker use in patients with MI and risk factors for shock ( $>70$  years of age, symptoms  $<12$  hours [ST-elevation MI patients], systolic blood pressure  $<120$  mm Hg, and heart rate  $>110$  beat/min on presentation) is associated with an increased risk of shock or death (S5.1.5.2-17). In the setting of polymorphic VT after MI, beta blockers reduce mortality (S5.1.5.2-18). Beta blockers suppress VA in some patients with structurally normal hearts (S5.1.5.2-19). When used in combination with membrane-stabilizing antiarrhythmic medications, beta blockers can enhance antiarrhythmic efficacy (S5.1.5.2-20). Beta blockers (e.g., nadolol, propranolol) are also first-line therapy for some cardiac channelopathies (e.g., long QT syndrome, catecholaminergic polymorphic ventricular tachycardia).

#### 5.1.3. Amiodarone and Sotalol

Amiodarone possesses a wide spectrum of actions that include blockade of beta receptors and sodium, calcium and potassium currents (i.e., a multichannel blocker). Its overall long-term effect on survival is controversial, with most studies showing no clear advantage over placebo. A few studies and a meta-analysis of several large studies have shown a reduction in SCD using amiodarone in patients with LV dysfunction due to prior MI and NICM (S5.1.5.2-21–S5.1.5.2-23), but SCD-HeFT showed no survival benefit from amiodarone compared with placebo (S5.1.5.2-24). A secondary analysis of the SCD-HeFT showed increased risk of mortality with amiodarone in patients with New York Heart Association (NYHA) class III symptoms (S5.1.5.2-25). A systematic review of the literature in high-risk patients (LVEF  $<40\%$ , with or without coronary disease), concluded that, for primary prevention, amiodarone, compared with no treatment or

**TABLE 7** Pharmacological Characteristics of Available Antiarrhythmic Medications for Treating VA

Antiarrhythmic Medication (Class) and Dose	Uses in VA/SCA	Target	Electrophysiological Effects	Pharmacological Characteristics	Common Adverse Effects
Acebutolol PO 200-1200 mg daily or up to 600 mg bid	VT, PVCs	Beta 1, Mild intrinsic sympathomimetic activity	Sinus rate slowed AV nodal refractoriness increased	Active metabolite $t_{1/2}$ : 8-13 h pProlonged with renal impairment Metab: H Excr: F 60%, U 40%	Cardiac: Bradycardia, hypotension, HF, AVB Other: Dizziness, fatigue, anxiety, impotence, hyper/hypoesthesia
Amiodarone (III) IV: 300 mg bolus for VF/pulseless VT arrest; 150-mg bolus for stable VT; 1 mg/min x 6 h, then 0.5 mg/min x 18 h PO: 400 mg* q 8 to 12 h for 1-2 wk, then 300-400 mg daily; reduce dose to 200 mg daily if possible	VT, VF, PVC,	$I_{Na}$ , $I_{Ca}$ , $I_{Kr}$ , $I_{K1}$ , $I_{Ks}$ , $I_{to}$ , Beta receptor, Alpha receptor nuclear T3 receptor	Sinus rate slowed QRS prolonged QTc prolonged AV nodal refractoriness increased; increased DFT	$t_{1/2}$ : 26-107 d Metab: H Excr: F	Cardiac: Hypotension, bradycardia, AVB, TdP, slows VT below programmed ICD detection rate, increases defibrillation threshold Other: Corneal microdeposits, thyroid abnormalities, ataxia, nausea, emesis, constipation, photosensitivity, skin discoloration, ataxia, dizziness, peripheral neuropathy, tremor, hepatitis, cirrhosis, pulmonary fibrosis or pneumonitis
Atenolol (II) PO: 25-100 mg qd or bid	VT, PVC, ARVC, LQTS	Beta 1	Sinus rate slowed AV nodal refractoriness increased	$t_{1/2}$ : 6-7 h (prolonged with renal impairment) Metab: H Excr: F 50%, U 40%	Cardiac: Bradycardia, hypotension, HF, AVB Other: Dizziness, fatigue, depression, impotence
Bisoprolol (II) PO: 2.5-10 mg once daily	VT, PVC	Beta 1 receptor	Sinus rate slowed AV nodal refractoriness increased	$t_{1/2}$ : 9-12 h Metab: H Excr: U	Cardiac: Chest pain, bradycardia, AVB Other: Fatigue, insomnia, diarrhea
Carvedilol (II) PO: 3.125-25 mg q 12 h	VT, PVC	Beta 1 and 2 receptors, Alpha	Sinus rate slowed AV nodal refractoriness increased	$t_{1/2}$ : 7-10 h Metab: H Excr: F	Cardiac: Bradycardia, hypotension, AVB, edema, syncope Other: Hyperglycemia, dizziness, fatigue, diarrhea
Diltiazem (IV) IV: 5-10 mg qd: 15-30 min Extended release: PO: 120-360 mg/day	VT specifically RVOT, idiopathic LVT	$I_{Ca-L}$	Sinus rate slowed PR prolonged AV nodal conduction slowed	$t_{1/2}$ : Injection 2-5 h, immediate release 4.5-12 h, extended release 12 h, and severe hepatic impairment 14-16 h Metab: H Excr: U	Cardiac: Hypotension, edema, HF, AVB, bradycardia, exacerbation of HFrEF Other: Headache, rash, constipation
Esmolol (II) IV: 0.5 mg/kg bolus, 0.05 mg/kg/min	VT	Beta 1 receptor	Sinus rate slowed AV nodal refractoriness increased	$t_{1/2}$ : 9 min Metab: RBC esterases Excr: U	Cardiac: Bradycardia, hypotension, HF, AVB Other: Dizziness, nausea
Flecainide (IC) PO: 50-200 mg q 12 h	VT, PVC (in the absence of structural heart disease). Has a role in treating patients with CPVT	$I_{Na}$ , $I_{Kr}$ , $I_{Kur}$	PR prolonged QRS prolonged; increased DFT	$t_{1/2}$ : 7-22 h Metab: H Excr: U	Cardiac: Sinus node dysfunction, AVB, drug-induced Brugada syndrome, monomorphic VT in patients with a myocardial scar, exacerbation of HFrEF Other: Dizziness, tremor, vision disturbance, dyspnea, nausea
Lidocaine (IB) IV: 1 mg/kg bolus, 1-3 mg/min 1-1.5 mg/kg. Repeat 0.5-0.75 mg/kg bolus every 5-10 min (max cumulative dose 3 mg/kg). Maintenance infusion is 1-4 mg/min although one could start at 0.5 mg/min	VT, VF	$I_{Na}$	No marked effect on most intervals; QTc can slightly shorten	Initial $t_{1/2}$ 7-30 min; terminal 90-120 min. Prolonged in HF, liver disease, shock, severe renal disease Metab: H Excr: U	Cardiac: Bradycardia, hemodynamic collapse, AVB, sinus arrest Other: Delirium, psychosis, seizure, nausea, tinnitus, dyspnea, bronchospasm

(continued on the next page)

**TABLE 7** Continued

Antiarrhythmic Medication (Class) and Dose	Uses in VA/SCA	Target	Electrophysiological Effects	Pharmacological Characteristics	Common Adverse Effects
Metoprolol (II) IV: 5 mg q 5 min up to 3 doses PO: 25-100 mg Extended release qd or q 12 h	VT, PVC	Beta 1 receptor	Sinus rate slowed AV nodal refractoriness increased	t <sub>1/2</sub> : 3-4 h Metab: H Excr: U	Cardiac: Bradycardia, hypotension, AVB Other: Dizziness, fatigue, diarrhea, depression, dyspnea
Mexiletine (IB) PO: 150-300 mg q 8 h or q 12 h	T, VF, PVC, has a role in patients with LQT3	I <sub>Na</sub>	No marked effect on most intervals; QTc can slightly shorten	t <sub>1/2</sub> : 10-14 h Metab: H Excr: U	Cardiac: HF, AVB Other: Lightheaded, tremor, ataxia, paresthesias, nausea, blood dyscrasias
Nadolol (II) PO: 40-320 mg daily	VT, PVC, LQTS, CPVT	Beta 1 and 2 receptors	Sinus rate slowed AV nodal refractoriness increased	t <sub>1/2</sub> : 20-24 h Metab: none Excr: U	Cardiac: Bradycardia, hypotension, HF, AVB Other: Edema, dizziness, cold extremities, bronchospasm
Procainamide (IA) IV: loading dose 10-17 mg/kg at 20-50 mg/min Maintenance dose: 1-4 mg/min PO (SR preparation): 500-1250 mg q 6 h	VT	I <sub>Na</sub> , I <sub>Kr</sub>	QRS prolonged QTc prolonged; increased DFT	Metab: H t <sub>1/2</sub> : 2-5 h; NAPA 6-8 h t <sub>1/2</sub> prolonged in renal dysfunction. Anephric: proc 11 h and NAPA 42 h Excr: U	Cardiac: TdP; AVB, hypotension and exacerbation of HF/EF Other: Lupus symptoms, diarrhea, nausea, blood dyscrasias
Propafenone (IC) PO: Immediate release 150-300 mg q 8 h Extended release 225-425 mg q 12 h	VT, PVC (in the absence of structural heart disease)	I <sub>Na</sub> , I <sub>Kr</sub> , I <sub>Kur</sub> , Beta receptor, Alpha receptor	PR prolonged QRS prolonged; increased DFT	t <sub>1/2</sub> : 2-10 h or 10-32 h t <sub>1/2</sub> extensive metabolizers 2-10 h; poor metabolizers 10-32 h. Metab: H Excr: U	Cardiac: HF, AVB, drug-induced Brugada syndrome Other: Dizziness, fatigue, nausea, diarrhea, xerostomia, tremor, blurred vision
Propranolol (II) IV: 1-3 mg q 5 min to a total of 5 mg PO: Immediate release 10-40 mg q 6 h; Extended release 60-160 mg q 12 h	VT, PVC, LQTS	Beta 1 and 2 receptors, I <sub>Na</sub>	Sinus rate slowed AV nodal refractoriness increased	t <sub>1/2</sub> : Immediate release 3-6 h Extended release 8-10 h Metab: H Excr: U	Cardiac: Bradycardia, hypotension, HF, AVB Other: Sleep disorder, dizziness, nightmares, hyperglycemia, diarrhea, bronchospasm
Quinidine (IA) PO: sulfate salt 200-600 mg q 6 h to q 12 h gluconate salt 324-648 mg q 8 h to q 12 h IV: loading dose: 800 mg in 50 mL infused at 50 mg/min	T, VF, (including short QT syndrome, Brugada)	I <sub>Na</sub> , I <sub>to</sub> , I <sub>Kr</sub> , M, Alpha receptor	QRS prolonged QTc prolonged; increased DFT	t <sub>1/2</sub> : 6-8 h longer in HF, liver cirrhosis, and with older age Metab: H Excr: U	Cardiac: Syncope, TdP, AVB Other: Dizziness, diarrhea, nausea, esophagitis, emesis, tinnitus, blurred vision, rash, weakness, tremor; blood dyscrasias
Ranolazine (not classified) PO: 500-1000 mg q 12 h	VT	I <sub>Na</sub> , I <sub>Kr</sub>	Sinus rate slowed Tc prolonged	t <sub>1/2</sub> : 7 h Metab: H Excr: U 75%, F 25%	Cardiac: Bradycardia, hypotension Other: Headache, dizziness, syncope, nausea, dyspnea
Sotalol (III) IV: 75 mg q 12 h PO: 80-120 mg q 12 h, may increase dose every 3 d; max 320 mg/d	VT, VF, PVC	I <sub>Kr</sub> , Beta 1 and 2 receptor	Sinus rate slowed QTc prolonged AV nodal refractoriness increased; decreased DFT	t <sub>1/2</sub> : 12 h Metab: none Excr: U	Cardiac: Bradycardia, hypotension, HF, syncope, TdP Other: Fatigue, dizziness, weakness, dyspnea, bronchitis, depression, nausea, diarrhea
Verapamil (IV) IV: 2.5-5 mg q 15-30 min Sustained release PO: 240-480 mg/d	VT (specifically RVOT, verapamil-sensitive idiopathic LVT)	I <sub>Ca-L</sub>	Sinus rate slowed PR prolonged AV nodal conduction slowed	t <sub>1/2</sub> : 3-7 h Metab: H Excr: U	Cardiac: Hypotension, edema, HF, AVB, bradycardia, exacerbation of HF/EF Other: Headache, rash, gingival hyperplasia, constipation, dyspepsia

\*Although up to 800 mg every 8 h might be used, higher doses of amiodarone are associated with a higher risk of adverse events. Modified from Shleifer JW, et al. (S5.1.5.2-2).

Alpha indicates alpha-adrenergic receptor; ARVC, arrhythmogenic right ventricular cardiomyopathy; AV, atrioventricular; AVB, atrioventricular block; Beta, beta-adrenergic receptor; HF, heart failure; CPVT, catecholaminergic polymorphic ventricular tachycardia; DFT, defibrillation threshold; F, feces; H, hepatic; I<sub>Ca</sub>, L-type calcium channel current; I<sub>K1</sub>, inward rectifier potassium channel; I<sub>KACH</sub>, muscarinic receptor-gated potassium channel; I<sub>KATP</sub>, adenosine-activated potassium channel; I<sub>Kr</sub>, rapid delayed rectifier potassium current; I<sub>Ks</sub>, slow delayed rectifier potassium current; I<sub>Kur</sub>, ultra-rapid delayed rectifier potassium current; I<sub>Na</sub>, fast inward sodium current; I<sub>to</sub>, transient outward potassium current; LQTS, long-QT syndrome; LVT, left ventricular tachycardia; M, muscarinic; Metab, metabolism; NAPA, n-acetyl procainamide; PVC, premature ventricular complex; QTc, corrected QT interval; t<sub>1/2</sub>, half-life; RVOT, right ventricular outflow tract; T3, triiodothyronine; TdP, torsades de pointes; U, urine; VT, ventricular tachycardia; and VF, ventricular fibrillation.

placebo, decreased the risk of SCD (Risk ratio: 0.76; 95% CI: 0.66-0.88) and all-cause mortality (Risk ratio: 0.88; 95% CI: 0.78-1.00), but the quality of the supporting evidence was very low ([S5.1.5.2-26](#)). For secondary prevention of SCD, the same systematic review identified neither risk nor benefit with amiodarone ([S5.1.5.2-26](#)). Compared with beta-blocker therapy and other antiarrhythmic medications (including sotalol), amiodarone appears to reduce the risk of SCD and all-cause mortality ([S5.1.5.2-26](#)). Intravenous amiodarone has a role in reducing recurrent VF/VF during resuscitation ([S5.1.5.2-3,S5.1.5.2-27-S5.1.5.2-29](#)).

Chronic administration of amiodarone is associated with complex medication interactions and a host of adverse effects involving the lung, liver, thyroid, skin, and nervous system. As a general rule, the longer the therapy and the higher dose of amiodarone, the greater the likelihood of adverse effects that will require discontinuance of the medication ([S5.1.5.2-26](#)). For this reason, chronic treatment of young patients with amiodarone should be reserved as a bridge to more definitive treatment options such as catheter ablation. Baseline evaluation of patients may include ECG, liver function tests, thyroid function tests, chest x-ray, and pulmonary function tests (including diffusing capacity of the lungs for carbon monoxide). Monitoring for toxicity generally includes periodic history and physical examination, as well as evaluation of the ECG, chest x-ray, and thyroid, liver, and lung function. High-resolution chest CT is generally reserved for suspected pulmonary toxicity ([S5.1.5.2-30](#)).

Although sotalol has some efficacy in suppressing VA, it has significant proarrhythmic effects and has not been shown to improve survival ([S5.1.5.2-31](#)). D-sotalol was shown in the SWORD (Survival With Oral d-Sotalol) trial to increase the risk of death in patients with heart failure ([S5.1.5.2-32](#)). Unlike amiodarone and many other antiarrhythmic agents, sotalol appears to reduce the defibrillation threshold ([S5.1.5.2-33](#)). Also, sotalol may lead to HF decompensation, and so its use in patients with an LVEF <20% is generally avoided.

#### 5.1.4. Calcium Channel Blockers

For the treatment of most VA, nondihydropyridines calcium channel blockers have no role. In fact, intravenous verapamil given for sustained VT has been associated with hemodynamic collapse, especially in patients with prior MI ([S5.1.5.2-34,S5.1.5.2-35](#)). For patients with a structurally normal hearts, verapamil or diltiazem can suppress some outflow tract origin ([S5.1.5.2-35-S5.1.5.2-39](#)). Oral and intravenous verapamil are effective in treating idiopathic interfascicular reentrant LVT ([S5.1.5.2-38](#)). Calcium channel blockers should not be given to patients with VT in the setting of HFrEF.

#### 5.1.5. Nonantiarrhythmic Medications and Therapies

##### 5.1.5.1. Electrolytes

Administration of potassium and magnesium has been proposed as helpful adjuncts in the prevention of VA ([S5.1.5.2-40,S5.1.5.2-41](#)). Hypokalemia and hypomagnesemia are common consequences of diuretic therapy in HF, both have been associated with VA during an acute MI ([S5.1.5.2-41,S5.1.5.2-42](#)), and can increase the risk of torsades de pointes in patients on medications or with conditions known to prolong the QT interval ([S5.1.5.2-43](#)). In fact, in patients with torsades de pointes, intravenous magnesium is first-line therapy ([S5.1.5.2-44](#)). In patients who are deficient in both magnesium and potassium, magnesium should be repleted to facilitate replacement of the potassium ([S5.1.5.2-45](#)). In the case of potassium, some recommend keeping the potassium level between 4.5 mmol/L and 5 mmol/L to prevent VA and SCD ([S5.1.5.2-46,S5.1.5.2-47](#)). A large observational study of patients with an acute MI found that the lowest rates of death were seen in patients with serum potassium concentrations between 3.5 mmol/L and <4.5 mmol/L ([S5.1.5.2-48](#)). Interestingly, the rates of VA did not rise unless the potassium was <3 mmol/L or ≥5 mmol/L. Likewise, a large randomized, double-blind trial of intravenous magnesium in the post-MI period demonstrated no benefit in 30-day mortality ([S5.1.5.2-40](#)). It remains quite reasonable to monitor potassium and magnesium during aggressive diuresis and in the post-MI period.

##### 5.1.5.2. n-3 Fatty Acids and Lipids

Both n-3 poly-unsaturated fatty acids and statin therapies may have a role in the prevention of SCD, thought to be due to a stabilization of the bilipid myocyte membrane involved in maintaining electrolyte gradients ([S5.1.5.2-49](#)).

Early data were promising regarding the effects of n-3 polyunsaturated fatty acids on the reduction of cardiovascular events and SCD. In 2006, a large meta-analysis of 19 observational and RCTs demonstrated a significant association between the consumption of n-3 polyunsaturated fatty acids and prevention of SCD ([S5.1.5.2-50](#)). The randomized GISSI (Gruppo Italiano per lo Studio della Sopravvivenza nell'Infarto)-Prevenzione trial in people with recent MI, found that fish oil 1 g/d reduced mortality, due to fewer SCD ([S5.1.5.2-51](#)). However, subsequent RCTs have not replicated these benefits and have shown n-3 polyunsaturated fatty acids to be ineffective ([S5.1.5.2-52-S5.1.5.2-56](#)). Because studies showed a consistent lack of harm from n-3 polyunsaturated fatty acids, patients can be reassured of their safety. Longer-term data will hopefully clarify the conflicting results.

In contrast, statin medications clearly reduce mortality and appear to reduce the risk of SCD related to ischemic heart disease ([S5.1.5.2-57](#)). The predominant mechanism

remains uncertain. Prevention of coronary plaque rupture or a direct cardioprotective effect reducing VA has been suggested. Experimental ischemia/reperfusion models demonstrate a cardioprotective effect of statins, and a large observational analysis observed this effect in humans ([S5.1.5.2-42](#),[S5.1.5.2-56](#)–[S5.1.5.2-58](#)). This was explored further in HF in several secondary analyses of patients on statins in ICD prevention trials, including the MADIT-CRT (Multicenter Automatic Defibrillator Implantation Trial-Cardiac Resynchronization Therapy), SCD-HeFT, AVID (Antiarrhythmics versus Implantable Defibrillators) ([S5.1.5.2-59](#)), and DEFINITE (DEFibrillators In Non-Ischemic Cardiomyopathy Treatment Evaluation)

trials that showed less SCD risk among the patients on statins ([S5.1.5.2-58](#),[S5.1.5.2-60](#)–[S5.1.5.2-62](#)). However, this general effect in HF was not confirmed in 2 prospective RCTs of rosuvastatin in HF; the CORONA (Controlled Rosuvastatin Multinational Trial in Heart Failure) and GISSI-HF (Gruppo Italiano per lo Studio della Sopravvivenza nell’Insufficienza Cardiaca-Heart Failure) ([S5.1.5.2-63](#),[S5.1.5.2-64](#)). It appears that the beneficial effects of statins are confined to the population with or at risk for atherosclerotic cardiovascular disease and/or ischemia, and not HF generally.

## 5.2. Preventing SCD With HF Medications

### Recommendation for Pharmacological Prevention of SCD

References that support the recommendation are summarized in [Online Data Supplement 10](#).

COR	LOE	RECOMMENDATION
I	A	1. In patients with HFrEF (LVEF ≤40%), treatment with a beta blocker, a mineralocorticoid receptor antagonist and either an angiotensin-converting enzyme inhibitor, an angiotensin-receptor blocker, or an angiotensin receptor-neprilysin inhibitor is recommended to reduce SCD and all-cause mortality ( <a href="#">S5.2-1</a> – <a href="#">S5.2-8</a> ).

### Recommendation-Specific Supportive Text

1. For patients with HF and depressed LV function, appropriate medical therapy is important to reduce SCD. These therapies have various beneficial effects on arrhythmia mechanisms. Beta blockers reduce myocardial oxygen demand and electrical excitability, and counter arrhythmogenic effects of sympathetic stimulation. Angiotensin-converting enzyme inhibitors and angiotensin-receptor blockers decrease preload and afterload, decreasing myocardial oxygen demand, blocking the formation of angiotensin II, and slowing the progression of ventricular remodeling and fibrosis. Mineralocorticoid receptor antagonists limit potassium loss, decrease fibrosis, and increase the myocardial uptake of norepinephrine ([S5.2-7](#)).

RCTs in patients with HFrEF have consistently demonstrated that chronic therapy with beta blockers reduces all-cause mortality, VA, and SCD ([S5.2-2](#),[S5.2-4](#), [S5.2-5](#),[S5.2-9](#)). Three beta blockers (i.e., bisoprolol, carvedilol, sustained-release metoprolol succinate) have been proven to reduce mortality in patients with current or prior symptoms of HFrEF without beta-blocker contraindications. Angiotensin-converting enzyme inhibition also reduces mortality and SCD ([S5.2-3](#)). Angiotensin-receptor blockers added to angiotensin-converting enzyme inhibitor showed additional benefit to angiotensin-converting enzyme inhibitors in some ([S5.2-10](#)) but not other RCTs ([S5.2-8](#),[S5.2-11](#)). Therapy with the mineralocorticoid-receptor antagonists, spironolactone and eplerenone,

have also demonstrated reductions in both all-cause mortality and SCD ([S5.2-6](#),[S5.2-12](#),[S5.2-13](#)). Recent studies of the angiotensin receptor-neprilysin inhibitor (sacubitril/valsartan) versus angiotensin-converting enzyme inhibitor demonstrated a reduction in SCD and cardiac mortality ([S5.2-14](#)).

## 5.3. Defibrillators for Treatment of VA and SCD

See [Sections 7, 10.2, 10.3, 10.8](#), and 10.9.

Defibrillation is highly effective in terminating life-threatening VA. This therapy can be delivered by a transvenous ICD, a subcutaneous implantable cardioverter-defibrillator, a wearable cardioverter-defibrillator or an external defibrillator. These devices monitor the heart rhythm continuously and deliver therapy in response to a tachycardia that meets preprogrammed detection rates and arrhythmia duration. The vast majority of transvenous ICDs are implanted in the subclavicular area under fluoroscopy guidance. Subcutaneous implantable cardioverter-defibrillators are implanted in the left side of the chest over the sixth rib between the left midaxillary and left anterior axillary lines. ICDs with epicardial sensing and pacing leads are still being implanted in some patients especially those with certain forms of congenital heart disease.

The transvenous ICD has been in clinical use for >3 decades, and robust data from high-quality RCTs support its use in various patient populations including survivors of cardiac arrest, patients with VT and structural heart disease, and patients with significant LV dysfunction.

## 5.4. Catheter Ablation

### 5.4.1. General Considerations

Catheter ablation is an important treatment option for patients with VA when antiarrhythmic medications are ineffective, not tolerated, or not desired by the patient. Monomorphic VA usually have an origin or substrate that can be targeted for ablation. Ablation is an option for selected patients with polymorphic VT/VF only if an initiating PVC focus or substrate can be identified. The ablation strategy, risks and outcomes are related to the mechanism and location of the VA. Most VA originate close to the subendocardium and are approached through a transvenous (for the right ventricle) or transaortic/transeptal (for the left ventricle) catheterization. Some diseases give rise to VA from the subepicardium, which may be approached by epicardial mapping and ablation. Pericardial access is usually achieved by a percutaneous subxiphoid puncture. The catheter ablation procedure usually involves attempts to induce VT by programmed electrical stimulation to confirm the diagnosis and guide ablation. Problems limiting success include inability to induce an arrhythmia for mapping (common with idiopathic VA), or origin of the arrhythmia from an inaccessible location in the myocardium (common in some cardiomyopathies).

### 5.4.2. VA in Patients With No Apparent Structural Heart Disease

See [Section 8](#).

VA that are not associated with underlying structural heart disease or a genetic arrhythmia syndrome are commonly referred to as idiopathic. Most idiopathic VA are monomorphic and based on a focal mechanism of triggered activity or abnormal automaticity; a few are due to reentry. For patients who are symptomatic, and in whom antiarrhythmic medications are ineffective, not tolerated, or not desired by the patient, catheter ablation is a treatment option. The ablation strategy is to identify the site of origin manifested by the earliest site of electrical activation, or when this is not practical, by pace mapping. Catheter ablation of idiopathic VA is usually accomplished with endocardial catheterization, though

an epicardial approach through the coronary venous circulation or a subxiphoid pericardial puncture may occasionally be required. Ablation failure for idiopathic VA is often due to inability to provoke the arrhythmia to allow mapping in the electrophysiological laboratory or origin from an inaccessible region.

### 5.4.3. Scar-Related VT

See [Section 8](#).

For most patients with structural heart disease, sustained monomorphic VT is due to reentry through regions of surviving myocardial fibers associated with areas of fibrous scar. The ablation strategy for these reentry circuits is to identify and eliminate channels of surviving myocardium within the scar that are often associated with slow conduction facilitating reentry. For most VTs that are related to prior MI, the substrate is on the subendocardial surface of the left ventricle. In NICM, the reentrant circuits are more variable in location, often involve the epicardial surface of either ventricle and frequently extending into the midmyocardium where ablation may be difficult to achieve from either surface. In tetralogy of Fallot specific reentry paths have been defined ([S5.4.3-1](#)). Electroanatomical mapping that helps clarify the relation of electrophysiological abnormalities to cardiac anatomy is commonly employed. Areas of scar can be appreciated as regions of relatively low electrogram voltage. For scar-related VTs, hemodynamic intolerance often limits mapping during VT. Ablation is then often guided by substrate mapping, in which areas of scar and potential reentry circuit substrate are delineated in electroanatomic maps based on electrocardiographic and pacing characteristics assessed during hemodynamically stable sinus or paced rhythm. Catheter ablation of scar-related VT requires an advanced level of experience by the operator, electrophysiological laboratory staff, and anesthesiologists as well as availability of surgical backup and specialized mapping, imaging, and ablation equipment ([S5.4.3-2](#),[S5.4.3-3](#)).

## 5.5. Surgery and Revascularization Procedures in Patients With Ischemic Heart Disease

### Recommendations for Surgery and Revascularization Procedures in Patients With Ischemic Heart Disease

References that support the recommendations are summarized in [Online Data Supplement 11](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. Patients with sustained VA and survivors of SCA should be evaluated for ischemic heart disease, and should be revascularized as appropriate ( <a href="#">S5.5-1–S5.5-4</a> ).
I	C-EO	2. In patients with anomalous origin of a coronary artery suspected to be the cause of SCA, repair or revascularization is recommended.

### Recommendation-Specific Supportive Text

1. Myocardial ischemia is a cause of sustained polymorphic VT/VF, and revascularization is an effective treatment to prevent myocardial ischemia. For patients with life-threatening VA, observational studies show that patients undergoing coronary artery bypass graft (CABG) had substantially better survival after accounting for other predictors (S5.5-1,S5.5-5). The risk of SCD appears comparable for patients with complex ischemic heart disease randomized to treatment with PCI versus CABG (S5.5-6). For patients with low LVEF and ischemic heart disease amenable to CABG, the risk of SCD is lower with CABG than medical therapy (S5.5-2,S5.5-7). Observational studies show an association between a lower likelihood of death with revascularization for survivors of SCA and CABG (S5.5-3) or PCI (S5.5-4). Revascularization alone is usually insufficient to prevent recurrence of sustained monomorphic VT; further evaluation for inducible VT is generally considered if ventricular function is depressed and/or scar is present.

2. Anomalous aortic origin of the coronary arteries is detected in approximately 1% of patients undergoing routine coronary angiography, and <0.2% of children and adolescents undergoing echocardiography (S5.5-8). Although ischemic heart disease is detected in as many as 24% to 55% of SCD cases in young patients <35 years of age (S5.5-9,S5.5-10), anomalous aortic origin of the coronary arteries is an important cause of SCD in the young, reported in 10% to 17% of patients included in postmortem studies (S5.5-10,S5.5-11). Anomalous origin of the coronary arteries can be identified by echocardiography, invasive coronary angiography, CT angiography or cardiac MRI. In patients with SCA or life-threatening VA presumed related to ischemia caused by anomalous origin of a coronary artery, repair or revascularization is performed to alleviate ischemia and reduce the recurrence of VA (S5.5-6,S5.5-7,S5.5-12–S5.5-14).

#### 5.5.1. Surgery for Arrhythmia Management

### Recommendation for Surgery for Arrhythmia Management

References that support the recommendation are summarized in [Online Data Supplement 12](#).

COR	LOE	RECOMMENDATION
IIb	C-LD	1. In patients with monomorphic VT refractory to antiarrhythmic medications and attempts at catheter ablation, surgical ablation may be reasonable (S5.5.1-1–S5.5.1-7).

### Recommendation-Specific Supportive Text

1. Cardiac surgery as a standalone procedure for VT is rarely performed, but has a role in some highly symptomatic patients, when antiarrhythmic medications and catheter ablation fails or are not possible, particularly if the failure of ablation is due to an arrhythmia arising from an area that is inaccessible to catheter ablation, such as deep in the myocardium, beneath epicardial fat, or near the coronary arteries. Surgical ablation of tachycardia can also be performed at the time of other

cardiac surgical interventions, such as during surgical resection of large aneurysms due to prior MI in which the border zone is often a substrate for VT, or placement of an LV assist device (LVAD) (S5.5.1-5–S5.5.1-7). The procedure requires detailed characterization of the arrhythmia usually with preoperative imaging and mapping, therefore, surgical ablation is best undertaken at tertiary referral centers and with collaboration between experienced surgeons and electrophysiologists.

#### 5.6. Autonomic Modulation

### Recommendations for Autonomic Modulation

References that support the recommendations are summarized in [Online Data Supplement 13 and 14](#).

COR	LOE	RECOMMENDATIONS
IIa	C-LD	1. In patients with symptomatic, non-life-threatening VA, treatment with a beta blocker is reasonable (S5.6-1).
IIb	C-LD	2. In patients with VT/VF storm in whom a beta blocker, other antiarrhythmic medications, and catheter ablation are ineffective, not tolerated, or not possible, cardiac sympathetic denervation may be reasonable (S5.6-2–S5.6-4).

## Synopsis

Sympathetic activation is proarrhythmic and parasympathetic activation is generally antiarrhythmic in VT/VF. Modulating the autonomic nervous system for the purpose of preventing arrhythmias is an emerging therapeutic modality. For the prevention of VA, autonomic modulation can be done either through interruption of sympathetic outflow to the heart, pharmacological beta blockade, or through stimulation of the parasympathetic pathway (e.g., vagal nerve stimulators, spinal cord stimulators). Although autonomic modulation has proven efficacy for certain conditions such as long QT syndrome and catecholaminergic polymorphic ventricular tachycardia (see [Section 7.9](#)), evidence is limited for its applicability to the broader group of VA, but studies are ongoing. Currently, there are limited data on the role of vagal nerve stimulators and spinal cord stimulators for the prevention of VA/SCD in humans, and thus no formal recommendation could be supported ([S5.6-5](#)).

## Recommendation-Specific Supportive Text

- Many patients with non-life-threatening VA require only reassurance, but others have symptoms that warrant therapy. A small RCT of patients with symptomatic VA demonstrated a significant reduction in the arrhythmic burden with atenolol ([S5.6-1](#)).
- VT/VF storm causes significant morbidity and is associated with increased mortality. For VT/VF storm refractory to treatment (medications, catheter ablation), cardiac sympathetic denervation has been shown in several small, observational studies ([S5.6-3,S5.6-6](#)) and 1 RCT ([S5.6-4](#)) to reduce the arrhythmia burden. This has been shown for left or bilateral cardiac sympathetic denervation, and it has been suggested that bilateral cardiac sympathetic denervation may be superior ([S5.6-3](#)). Although data are limited, the significant morbidity and limited options in these patients make cardiac sympathetic denervation a reasonable option in selected patients.

## 6. ACUTE MANAGEMENT OF SPECIFIC VA

### Recommendations for Management of Cardiac Arrest

References that support the recommendations are summarized in [Online Data Supplement 15 and 16](#).

COR	LOE	RECOMMENDATIONS
I	A	<ol style="list-style-type: none"><li>CPR should be performed in patients in cardiac arrest, according to published basic and advanced cardiovascular life support algorithms (<a href="#">S6-1–S6-3</a>).</li></ol>
I	A	<ol style="list-style-type: none"><li>In patients with hemodynamically unstable VA that persist or recur after a maximal energy shock, intravenous amiodarone should be administered to attempt to achieve a stable rhythm after further defibrillation (<a href="#">S6-1,S6-4–S6-6</a>).</li></ol>
I	A	<ol style="list-style-type: none"><li>Patients presenting with VA with hemodynamic instability should undergo direct current cardioversion (<a href="#">S6-1–S6-3</a>).</li></ol>
I	B-NR	<ol style="list-style-type: none"><li>In patients with polymorphic VT or VF with ST-elevation MI, angiography with emergency revascularization is recommended (<a href="#">S6-7–S6-10</a>).</li></ol>
I	C-EO	<ol style="list-style-type: none"><li>Patients with a wide-QRS tachycardia should be presumed to have VT if the diagnosis is unclear.</li></ol>
IIa	A	<ol style="list-style-type: none"><li>In patients with hemodynamically stable VT, administration of intravenous procainamide can be useful to attempt to terminate VT (<a href="#">S6-11–S6-13</a>).</li></ol>
IIa	B-R	<ol style="list-style-type: none"><li>In patients with a witnessed cardiac arrest due to VF or polymorphic VT that is unresponsive to CPR, defibrillation, and vasopressor therapy, intravenous lidocaine can be beneficial (<a href="#">S6-1,S6-4,S6-5,S6-14, S6-15</a>).</li></ol>
IIa	B-R	<ol style="list-style-type: none"><li>In patients with polymorphic VT due to myocardial ischemia, intravenous beta blockers can be useful (<a href="#">S6-16,S6-17</a>).</li></ol>

(continued)

IIa	B-NR	9. In patients with a recent MI who have VT/VF that repeatedly recurs despite direct current cardioversion and antiarrhythmic medications (VT/VF storm), an intravenous beta blocker can be useful (S6-17,S6-18).
IIb	A	10. In patients in cardiac arrest, administration of epinephrine (1 mg every 3 to 5 minutes) during CPR may be reasonable (S6-1,S6-19–S6-24).
IIb	B-R	11. In patients with hemodynamically stable VT, administration of intravenous amiodarone or sotalol may be considered to attempt to terminate VT (S6-5,S6-13,S6-25,S6-26).
III: No Benefit	A	12. In patients with cardiac arrest, administration of high-dose epinephrine (>1 mg boluses) compared with standard doses is not beneficial (S6-19,S6-21).
III: No Benefit	A	13. In patients with refractory VF not related to torsades de pointes, administration of intravenous magnesium is not beneficial (S6-27,S6-28).
III: Harm	B-R	14. In patients with suspected AMI, prophylactic administration of lidocaine or high-dose amiodarone for the prevention of VT is potentially harmful (S6-16,S6-29).
III: Harm	C-LD	15. In patients with a wide QRS complex tachycardia of unknown origin, calcium channel blockers (e.g., verapamil and diltiazem) are potentially harmful (S6-30,S6-31).

**Figure 2****Recommendation-Specific Supportive Text**

1. The most common electrical mechanisms for cardiac arrest are VF and pulseless VT, but substantial numbers of cardiac arrests begin as severe bradyarrhythmias or asystole. Survival is better for patients presenting with VT or VF than for those with bradycardiac or asystolic mechanisms (S6-32). Rapid arrival of paramedical personnel is the major determinant of survival. A number of strategies for responding to unexpected cardiac arrest, including rapid defibrillation and initiation of CPR for a witnessed cardiac arrest, have improved survival probabilities for cardiac arrest victims (S6-2,S6-3). Nonetheless, the absolute number and proportion of survivors remain low, except in unique circumstances where there is an extraordinarily rapid response time to victims in VF or VT such as in monitored intensive care units, where survival is >90% (S6-33–S6-36). Survival decreases rapidly after the initial 2 minutes from the onset of cardiac arrest, so that by 4 to 5 minutes, survival may be ≤25%, and by 10 minutes it is 0% (S6-33,S6-35,S6-36). Advanced life support activities, other than those directly related to cardioversion and defibrillation for control of tachyarrhythmias, have led to the generation of comprehensive protocols to guide responders. These AHA documents cover the broad expanse of clinical circumstances and considerations of mechanisms (S6-1,S6-37).

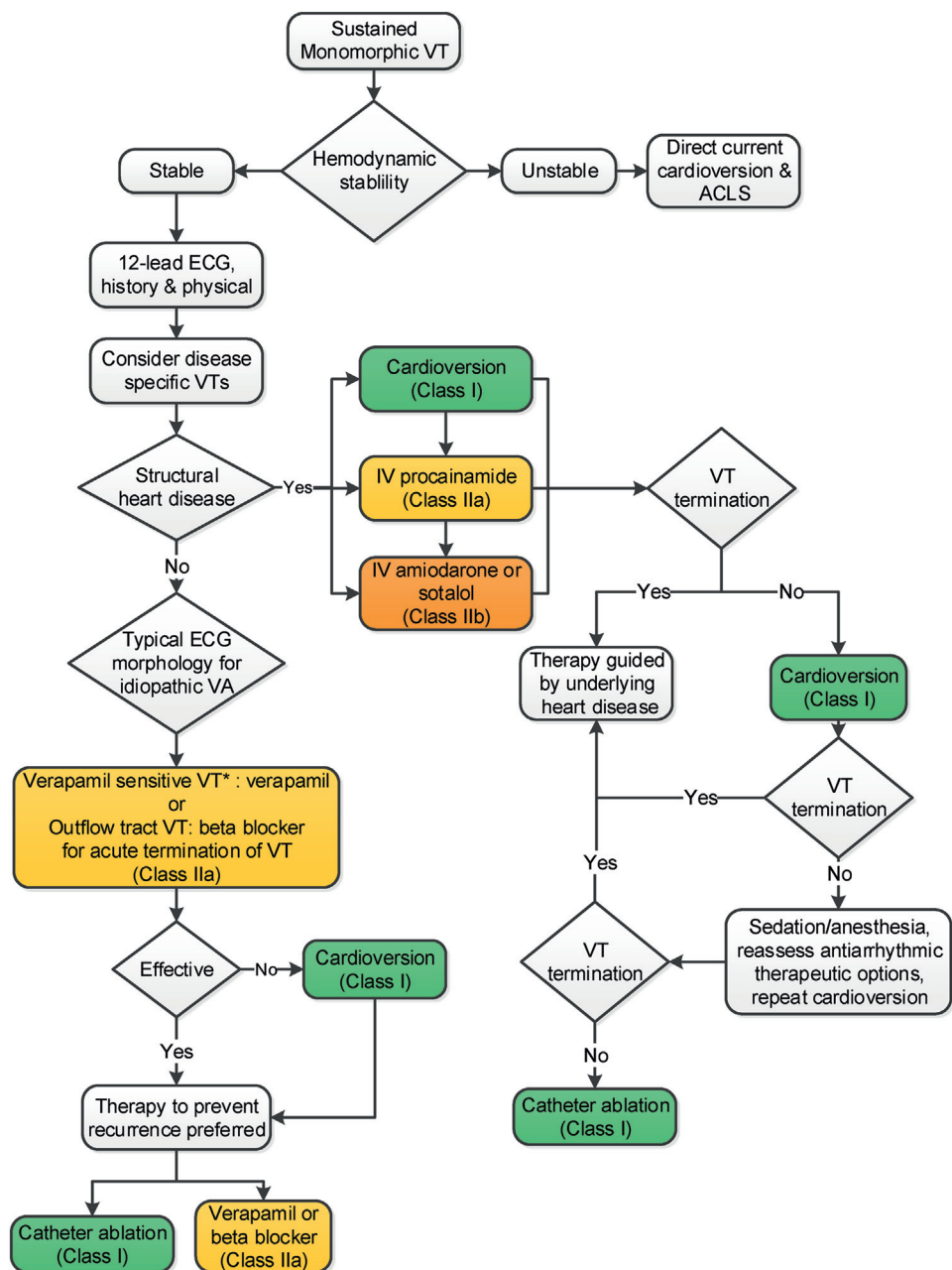
2. Paramedic administration of amiodarone after at least 3 failed shocks and administration of epinephrine improved hospital admission rates when compared with placebo (S6-6) or 1.5 mg/kg lidocaine (S6-1,S6-4) in RCTs in adults with out-of-hospital cardiac arrest due to refractory VF or polymorphic VT, although survival to hospital discharge and survival with favorable neurologic outcome were not improved with amiodarone or lidocaine (S6-5). However, in the subset of patients with witnessed cardiac arrest due to initial shock-refractory VF or pulseless VT, survival to hospital discharge after amiodarone administration was higher than with placebo (S6-5). The administration of procainamide in out-of-hospital cardiac arrest due to VF or pulseless VT has been associated with more shocks, more pharmacologic interventions, longer resuscitation times, and lower survival (S6-38).
3. VA with hemodynamic instability, including VF and pulseless monomorphic or polymorphic VT, causes loss of consciousness and leads to death if untreated. A short time to direct current cardioversion is the major determinant of survival, and defibrillation should be performed as quickly as possible. CPR is used until a perfusing rhythm is restored. If defibrillation is unsuccessful in returning spontaneous circulation, responders follow advanced cardiovascular life support activities (S6-1–S6-3).
4. Quickly identifying and treating patients with out-of-hospital cardiac arrest related to acute coronary occlusion is associated with improved survival and better functional recovery (S6-37). Coronary occlusion

as a cause of cardiac arrest is not reliably predicted by clinical and electrocardiographic findings (**S6-7**), and emergency coronary angiography should be considered (rather than later in the hospital stay or not at all) for unstable patients with a suspected cardiac etiology regardless of whether the patient is comatose or awake (**S6-9,S6-39**). In 1 observational study of patients resuscitated from SCA who did not have ST elevation and had angiography, one third were found to have a culprit lesion and coronary intervention appeared to be associated with a greater likelihood of favorable neurologic outcome (**S6-10**).

5. The initial management of any tachycardia should proceed according to published AHA advanced cardiovascular life support guidelines (**S6-40**). Immediate cardioversion should be performed for hemodynamic instability at presentation or if it develops subsequently. An ECG should be obtained for stable rhythms. Wide-complex tachycardias, defined by a QRS duration  $\geq 0.12$  s (**S6-37**), can be due to VT, SVT with aberrancy, preexcited tachycardia, or a paced rhythm such as pacemaker-mediated tachycardia. An irregular wide-complex tachycardia may be AF with aberrancy, preexcited AF (i.e., AF using an accessory pathway for anterograde conduction), atrial flutter, or VT (**S6-37**). A diagnosis should be established, and consultation with an arrhythmia expert considered (**S6-37**).
6. In 1 study, amiodarone was more effective than lidocaine in terminating incessant VT with improved survival at 24 hours (**S6-26**). For patients with recurrent, stable VT not in the setting of an AMI, intravenous procainamide has been shown to be superior to lidocaine for terminating the arrhythmia (**S6-11**). One randomized trial of 62 patients found procainamide superior to amiodarone for termination of stable VT (**S6-13**). Adverse events, including hypotension were more common with amiodarone, but the difference was not statistically significant. Procainamide and its metabolite n-acetylprocainamide have potassium channel blocking properties that may prolong the QT interval. In patients who already have QT prolongation, administration of procainamide may further prolong the QT interval and lead to torsades de pointes (**S6-11,S6-12,S6-26**).
7. Intravenous lidocaine is an alternative antiarrhythmic medication of long-standing and widespread familiarity. Compared with no antiarrhythmic medication, lidocaine did not consistently increase a return of spontaneous circulation after defibrillation and was not associated with improvement in survival to hospital discharge (**S6-4,S6-14,S6-41**). In prospective, blinded, RCTs, lidocaine was less effective than amiodarone in improving hospital admission rates

after out-of-hospital cardiac arrest due to shock-refractory VF or polymorphic VT; but there were no differences between the 2 medications in survival to hospital discharge (**S6-4,S6-5**). However, in the subset of patients with witnessed SCA due to initial shock-refractory VF or pulseless VT, a subgroup analysis showed that survival to hospital discharge with lidocaine was better than with placebo (**S6-5,S6-42**).

8. In a large meta-analysis of antiarrhythmic medications in the setting of AMI, beta blockers were associated with a significant reduction in mortality (**S6-16**). Beta blockers can be effective in suppressing recurrent VF in patients with recent MI, with an associated improvement in survival (**S6-17**).
9. In patients with recurrent VT/VF (VT/VF storm) in the setting of a recent MI that is refractory to amiodarone and/or lidocaine and repeated cardioversion, administration of a beta blocker has been shown to improve survival at 1 week. For those who did not survive, mortality was mostly due to recurrent VF. Survival at 1 year was also better in those treated with a beta blocker (**S6-17,S6-18**). Other measures to reduce sympathetic tone including sedation and general anesthesia are also often used.
10. Epinephrine produces beneficial effects in patients during cardiac arrest, primarily because of its alpha-adrenergic (i.e., vasoconstrictor) effects (**S6-1**). These alpha-adrenergic effects can increase coronary and cerebral perfusion pressure during CPR. The value and safety of the beta-adrenergic effects of epinephrine are controversial because they may increase myocardial work and reduce subendocardial perfusion (**S6-1**). One trial assessed short-term and longer-term outcomes when comparing standard-dose epinephrine to placebo (**S6-23**). Standard-dose epinephrine was defined as 1 mg given intravenously or intraosseously every 3 to 5 minutes. For both survival to discharge and survival to discharge with good neurologic outcome, there was no benefit with standard-dose epinephrine; however, the study was underpowered for analysis of either of these outcomes. There was, nevertheless, improved survival to hospital admission and improved return of spontaneous circulation with the use of standard-dose epinephrine. A number of trials have compared outcomes of standard-dose epinephrine with those of high-dose epinephrine. These trials did not demonstrate any benefit for high-dose epinephrine over standard-dose epinephrine in relation to survival to discharge with a good neurologic recovery, survival to discharge, or survival to hospital admission (**S6-1,S6-19,S6-21,S6-22**).
11. Amiodarone was more effective than lidocaine in terminating incessant VT with improved survival at 24

**FIGURE 2** Management of Sustained Monomorphic VT

Colors correspond to Class of Recommendation in Table 1. See Sections 7, 8.1.3, 8.2.3, and 10 for discussion. \*Known history of verapamil sensitive or classical electrocardiographic presentation. ACLS indicates advanced cardiovascular life support; ECG, electrocardiogram; VA, ventricular arrhythmia; and VT, ventricular tachycardia.

hours (S6-26). For patients with recurrent, stable VT not in the setting of an AMI, intravenous procainamide has been shown to be superior to lidocaine for terminating the arrhythmia (S6-11). One RCT in 62 patients found procainamide superior to amiodarone for termination of stable VT (S6-13). Adverse events,

including hypotension, were more common with amiodarone, but the difference was not statistically significant. Procainamide and its metabolite n-acetylprocainamide have potassium channel blocking properties that may prolong the QT interval. In patients who already have QT prolongation,

administration of procainamide may further prolong the QT interval and lead to torsades de pointes (S6-11). A single RCT of 33 patients comparing sotalol with lidocaine for treating patients with hemodynamically stable VT showed that VT was terminated in 69% of patients using sotalol and 18% using lidocaine (S6-25). Intravenous sotalol has been approved for use in the United States. Sotalol has potassium channel blocking properties that may prolong the QT interval. In patients who already have QT interval prolongation, administration of sotalol may further prolong the QT interval and lead to torsades de pointes (S6-25).

12. Epinephrine may increase coronary and cerebral perfusion pressure during CPR because of its vasoconstrictive effects. High doses of epinephrine (0.1 to 0.2 mg/kg IV, as opposed to a standard dose of 1 mg) have been studied in RCTs. In out-of-hospital cardiac arrest unresponsive to defibrillation, administration of high-dose epinephrine improved survival to hospital admission, but there was no difference compared to standard dose epinephrine in survival to hospital discharge (S6-19). There was also no improvement in long-term survival (S6-21). Of note, the administration of vasopressin is no longer recommended in the most recent advanced cardiovascular life support algorithms (S6-1).

13. Magnesium may suppress automaticity, suppress early and late after-depolarizations, and inhibit calcium flux into cardiomyocytes. It is effective in suppressing VA related to acquired long QT syndrome. However, 2 RCTs that investigated the use of intravenous magnesium in patients with cardiac arrest and refractory VF found no benefit (S6-27,S6-28). In a study of out-of-hospital cardiac arrest, administration of 2 to 4 g magnesium intravenously did not improve survival to hospital admission (S6-27). In a similar study involving inpatient cardiac arrest, magnesium did not improve return of spontaneous circulation, survival to 24 hours, or survival to hospital discharge (S6-28). There are

exceptions such as marked hypokalemia or medication-induced torsades de pointes in which administration of intravenous magnesium is warranted.

14. Several studies have tested the hypothesis that prophylactic administration of antiarrhythmic medications could reduce the incidence of post-MI VA and lead to better outcomes. One meta-analysis assessed studies in which beta blockers, class I antiarrhythmic agents such as lidocaine and procainamide, and amiodarone were given in the setting of AMI. The routine use of lidocaine and procainamide was associated with increased mortality, whereas beta blockers were associated with a significantly lower mortality rate (S6-16). Limited data with amiodarone appeared to be promising, but a subsequent RCT involving 1073 patients found that administration of high-dose amiodarone led to a higher mortality rate, although a moderate dose of amiodarone was not superior to placebo (S6-29).
15. With a stable, wide QRS complex tachycardia, differentiation between SVT with aberrancy and VT is often possible by review of the patient's history and the 12-lead ECG during tachycardia. Patients with wide QRS complex tachycardia and known structural heart disease should be presumed to have VT until proven otherwise. Administration of a calcium channel blocker such as verapamil to a patient with VT may result in severe hypotension or syncope (S6-31). The exception is verapamil-sensitive VT (interfascicular reentry) that occurs in a structurally normal heart; but this is often difficult to recognize on initial presentation (S6-30).

## 7. ONGOING MANAGEMENT OF VA AND SCD RISK RELATED TO SPECIFIC DISEASE STATES

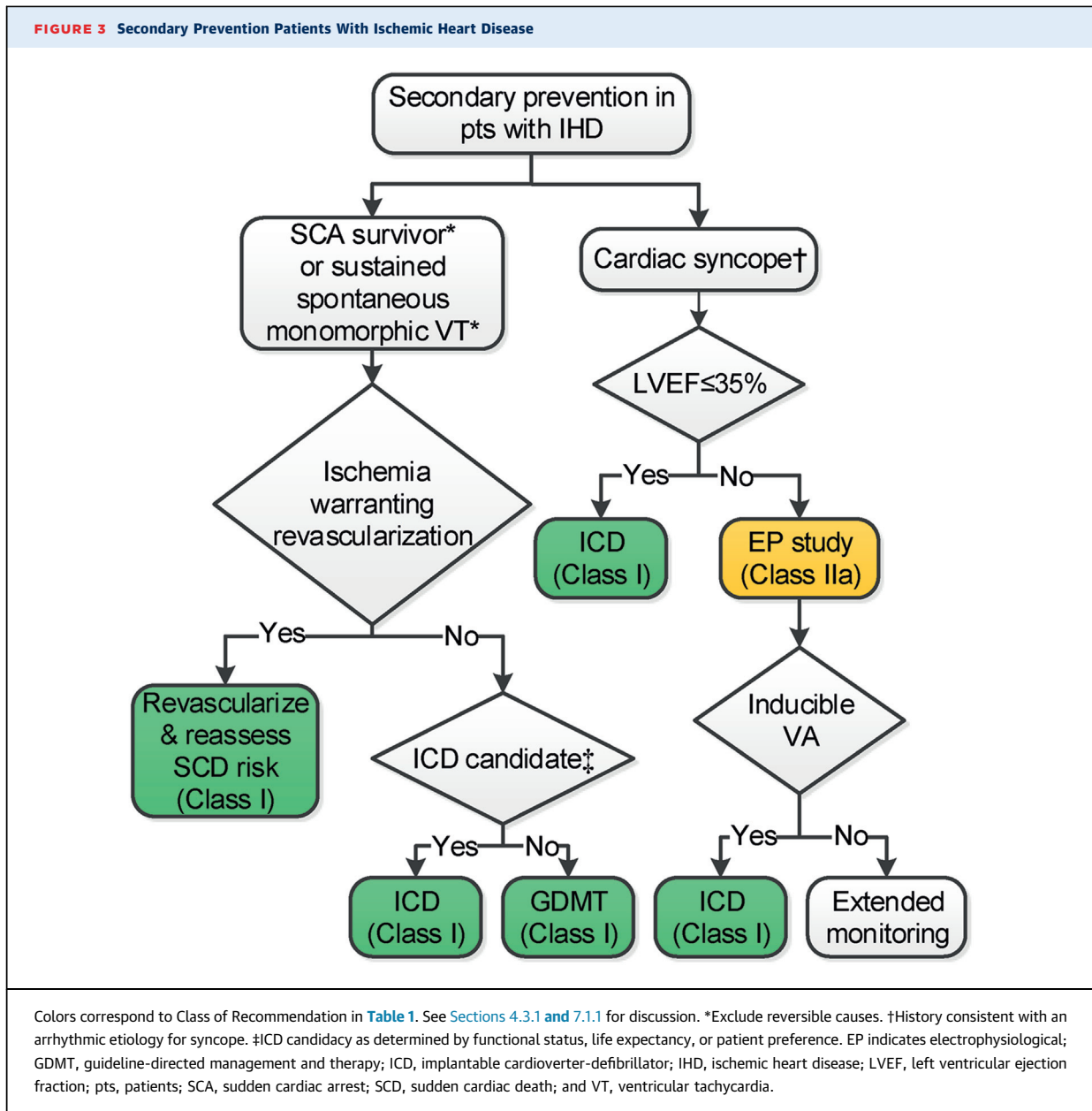
### 7.1. Ischemic Heart Disease

#### 7.1.1. Secondary Prevention of SCD in Patients With Ischemic Heart Disease

**Recommendations for Secondary Prevention of SCD in Patients With Ischemic Heart Disease**  
**References that support the recommendations are summarized in Online Data Supplement 17 and 18.**

COR	LOE	RECOMMENDATIONS
I	B-R B-NR	<p><b>Value Statement:</b> Intermediate Value (LOE: B-R)</p> <ol style="list-style-type: none"> <li>1. In patients with ischemic heart disease, who either survive SCA due to VT/VF or experience hemodynamically unstable VT (LOE: B-R) (S7.1.1-1–S7.1.1-4) or stable sustained VT (LOE: B-NR) (S7.1.1-5) not due to reversible causes, an ICD is recommended if meaningful survival greater than 1 year is expected.</li> <li>2. A transvenous ICD provides intermediate value in the secondary prevention of SCD particularly when the patient's risk of death due to a VA is deemed high and the risk of nonarrhythmic death (either cardiac or noncardiac) is deemed low based on the patient's burden of comorbidities and functional status (S7.1.1-6).</li> </ol>
I	B-NR	<ol style="list-style-type: none"> <li>3. In patients with ischemic heart disease and unexplained syncope who have inducible sustained monomorphic VT on electrophysiological study, an ICD is recommended if meaningful survival of greater than 1 year is expected (S7.1.1-7).</li> </ol>

Figure 3

**Recommendation-Specific Supportive Text**

- In the AVID trial (S7.1.1-1), the ICD improved overall survival compared with antiarrhythmic medication therapy (primarily amiodarone) in patients who survived SCD or with hemodynamically unstable VT, with a 2-year relative risk reduction in mortality of 27% and an absolute risk reduction of 7%. CIDS (Canadian Implantable Defibrillator Study) (S7.1.1-2), which was stopped early after the results of the AVID trial were released, showed a similar, but not statistically

significant, benefit of the ICD over antiarrhythmic medication therapy. A subsequent meta-analysis using data from 3 RCTs showed a statistically significant reduction in both arrhythmic and all-cause mortality with secondary prevention ICDs (S7.1.1-3).

In survivors of life-threatening VA that may be due to transient or reversible factors, such as AMI, proarrhythmic medication effects, or electrolyte disturbances, an ICD is not implanted if the cause may be correctable. This is a population of patients that still requires

thorough evaluation, treatment, and close follow-up and, as in the AVID registry, mortality was still high in the population that may have had a reversible cause for their arrest ([S7.1.1-8](#)). Small increases in troponin present a challenge in selecting patients for an ICD, as it often cannot be determined whether troponin elevation is due to ischemia from VT/VF and resuscitation, in which case an ICD is likely warranted, or an indication that ischemia caused the arrhythmia, in which case prevention of ischemia would be the therapeutic focus.

ICDs may improve the outcomes of patients with hemodynamically tolerated sustained VT and structural heart disease ([S7.1.1-5](#)); however, this has not proved in any RCT. VT ablation has been used as an alternative in selected patients with well-tolerated VT and appears to reduce recurrences, but the impact on long-term mortality is unknown; there is not yet sufficient evidence to recommend this approach as an alternative to ICD implantation ([S7.1.1-9,S7.1.1-10](#)).

2. Economic outcomes of ICD implantation for secondary prevention of SCD were assessed in the AVID and CIDS trials ([S7.1.1-11,S7.1.1-12](#)), as well as in a simulation model ([S7.1.1-13](#)) and an observational study of Medicare beneficiaries ([S7.1.1-14](#)). All studies compared ICD

recipients with non-ICD recipients, and all found that ICD recipients had longer overall survival and higher lifetime costs of medical care. All studies reported incremental cost-effectiveness ratios between \$64,000 and \$100,000 per year of life added by an ICD ([S7.1.1-11–S7.1.1-14](#)), which is in the range of intermediate value by the benchmarks applied in the ACC/AHA cost/value statement ([S7.1.1-15](#)).

3. VAs are an important cause of syncope or near syncope in patients with ischemic heart disease, particularly those with prior infarction. A study of 70 patients with unexplained syncope who underwent an electrophysiological study identified positive findings in 37 patients; 31 with VT. During 3 years of follow-up, patients with a positive electrophysiological study had higher rates of SCD and 3-year total mortality (61% versus 15%, respectively) than those with a negative electrophysiological study ([S7.1.1-7](#)). An ICD is warranted for patients with syncope and inducible sustained monomorphic VT even if they do not otherwise meet criteria for primary prevention ([Figure 4](#)).

### 7.1.1.1. Coronary Artery Spasm

#### Recommendations for Patients With Coronary Artery Spasm

References that support the recommendations are summarized in [Online Data Supplement 20](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with VA due to coronary artery spasm, treatment with maximally tolerated doses of a calcium channel blocker and smoking cessation are indicated to reduce recurrent ischemia and VA ( <a href="#">S7.1.1.1-1,S7.1.1.1-2</a> ).
IIa	B-NR	2. In patients resuscitated from SCA due to coronary artery spasm in whom medical therapy is ineffective or not tolerated, an ICD is reasonable if meaningful survival of greater than 1 year is expected ( <a href="#">S7.1.1.1-3–S7.1.1.1-6</a> ).
IIb	B-NR	3. In patients resuscitated from SCA due to coronary artery spasm, an ICD in addition to medical therapy may be reasonable if meaningful survival of greater than 1 year is expected ( <a href="#">S7.1.1.1-3–S7.1.1.1-6</a> ).

#### Recommendation-Specific Supportive Text

1. Coronary artery spasm results from vasomotor dysfunction and can occur in the presence or absence of atherosclerotic ischemic heart disease. Vasospasm episodes can lead to VA, syncope, and SCD. Treatment includes risk factor elimination including smoking cessation, and treatment with vasodilators including dihydropyridine calcium channel blockers with or without nitrates. A more detailed summary of treatments for coronary artery spasm can be found in other guideline documents ([S7.1.1.1-7,S7.1.1.1-8](#)).

2. Patients with coronary artery spasm who survive an SCA are a high-risk population ([S7.1.1.1-5](#)). Recurrent VA, even life-threatening, may be prevented if coronary artery spasm can be effectively addressed with risk factor modification, smoking cessation, and ongoing treatment with nitrates and dihydropyridine calcium channel blockers ([S7.1.1.1-9](#)). However, SCA or VA can recur despite medical therapy or if compliance is poor. Whether a wearable cardioverter-defibrillator may provide protection while medical therapy is being evaluated has not been assessed but is of interest

(S7.1.1.1-10). An ICD can terminate VT/VF initiated by spasm, potentially preventing SCD.

3. Patients with coronary vasospasm who survive an SCA are a high-risk population, and some support the use of an ICD in those patients based on the reported event rates from observational studies (S7.1.1.1-5) even before determining the patient's response to or compliance with medical therapy. Recurrent SCA can occur despite medical therapy. Regardless of the approach, risk factor modification (e.g., illicit drug use), smoking cessation, and ongoing treatment with dihydropyridine calcium channel blockers with or without nitrates represent essential treatments (S7.1.1.1-9).

#### 7.1.1.2. Post CABG VT/VF

The incidence of sustained VT or VF early after CABG is low, but these VAs are associated with high in-hospital mortality (S7.1.1.2-1). VF occurring very early (intraoperatively or within 24 hours postoperatively) may be due to the transient effects of reperfusion, electrolyte and acid base disturbances, and the use of inotropes. Patients who present with VF or polymorphic VT in the post-operative period more often have associated ischemia, while patients presenting with monomorphic VT usually have an old infarct and ventricular scar (S7.1.1.2-2). Polymorphic VT/VF occurring after CABG warrants a therapeutic approach targeting treatment of myocardial ischemia, including a possible need for assessment of graft patency, as well as identification and treatment of

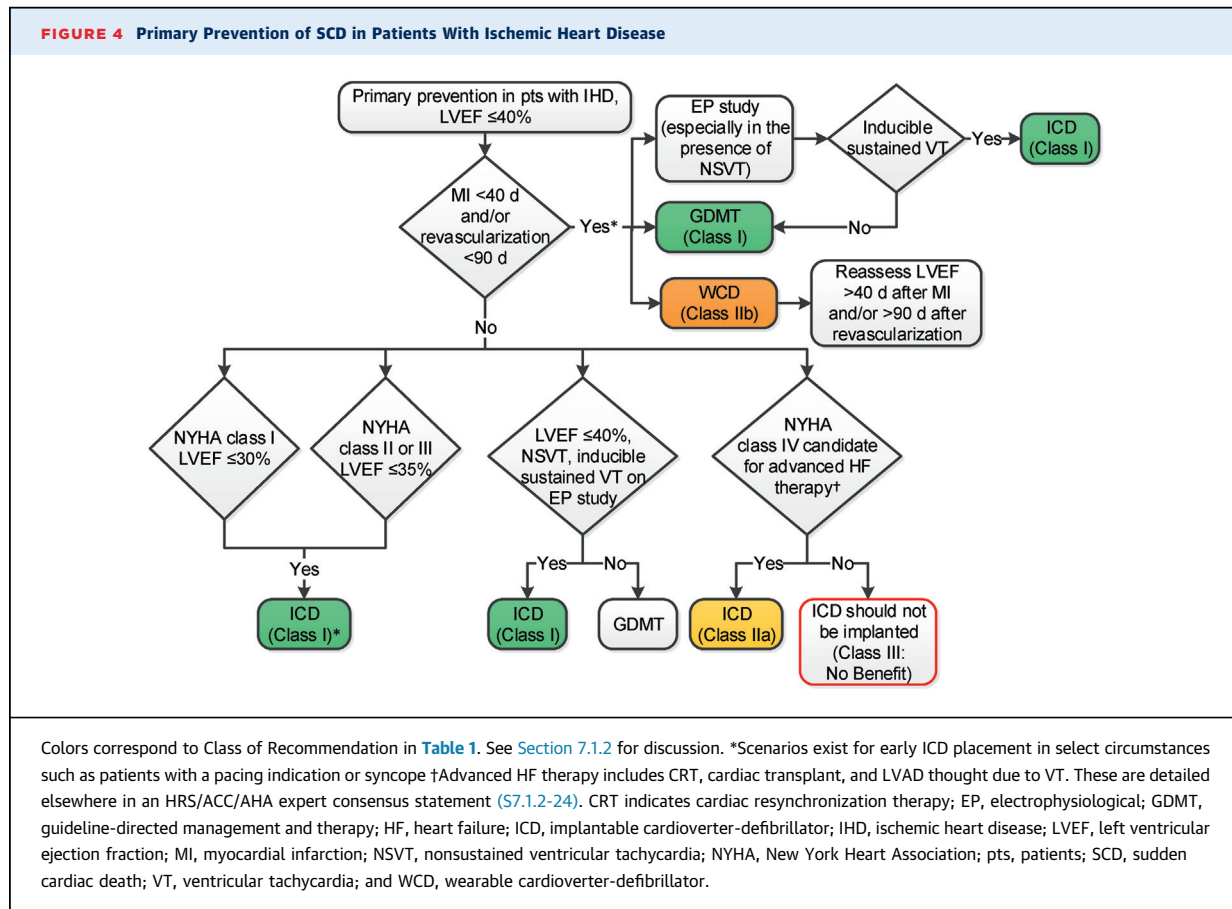
mechanical complications and acute electrolyte or acid base disturbances. Risk factors for occurrence of monomorphic VT early after CABG include prior MI, ventricular scar, LV dysfunction, and placement of a bypass graft across a noncollateralized occluded coronary vessel to a chronic infarct zone (S7.1.1.2-3). Unlike polymorphic VT and VF, sustained monomorphic VT is typically not due to acute ischemia. Many of these patients have inducible sustained VT at electrophysiological study. Management of symptomatic VA in the early period after CABG follows the recommendations for acute and ongoing management of VT detailed elsewhere in this document. In patients without sustained VT or VF but with LV dysfunction prior to undergoing CABG, implantation of an ICD did not improve survival (S7.1.1.2-4). For patients with LV dysfunction who are undergoing revascularization, there is a possibility that the LV function may improve, so many advocate for reassessment of the LV function 3 months after revascularization before a decision about ICD implantation is made (S7.1.1.2-5). For patients with a high burden of NSVT and reduced LVEF, an electrophysiological study may be helpful for risk stratification; those with inducible sustained VT may benefit from an ICD (S7.1.1.2-6). The wearable cardioverter-defibrillator may play a role in patients at risk of SCD in the early phase after revascularization to allow time for recovery of ventricular function (S7.1.1.2-7).

#### 7.1.2. Primary Prevention of SCD in Patients With Ischemic Heart Disease

##### Recommendations for Primary Prevention of SCD in Patients With Ischemic Heart Disease References that support the recommendations are summarized in Online Data Supplement 21.

COR	LOE	RECOMMENDATIONS
I	A	<ol style="list-style-type: none"> <li>1. In patients with LVEF of 35% or less that is due to ischemic heart disease who are at least 40 days' post-MI and at least 90 days postrevascularization, and with NYHA class II or III HF despite GDMT, an ICD is recommended if meaningful survival of greater than 1 year is expected (S7.1.2-1,S7.1.2-2).</li> </ol>
I	A	<ol style="list-style-type: none"> <li>2. In patients with LVEF of 30% or less that is due to ischemic heart disease who are at least 40 days' post-MI and at least 90 days postrevascularization, and with NYHA class I HF despite GDMT, an ICD is recommended if meaningful survival of greater than 1 year is expected (S7.1.2-2,S7.1.2-3).</li> </ol>
Value Statement: High Value (LOE: B-R)		<ol style="list-style-type: none"> <li>3. A transvenous ICD provides high value in the primary prevention of SCD particularly when the patient's risk of death due to a VA is deemed high and the risk of nonarrhythmic death (either cardiac or noncardiac) is deemed low based on the patient's burden of comorbidities and functional status (S7.1.2-4).</li> </ol>
I	B-R	<ol style="list-style-type: none"> <li>4. In patients with NSVT due to prior MI, LVEF of 40% or less and inducible sustained VT or VF at electrophysiological study, an ICD is recommended if meaningful survival of greater than 1 year is expected (S7.1.2-5).</li> </ol>
IIa	B-NR	<ol style="list-style-type: none"> <li>5. In nonhospitalized patients with NYHA class IV symptoms who are candidates for cardiac transplantation or an LVAD, an ICD is reasonable if meaningful survival of greater than 1 year is expected (S7.1.2-6–S7.1.2-9).</li> </ol>
III: No Benefit	C-EO	<ol style="list-style-type: none"> <li>6. An ICD is not indicated for NYHA class IV patients with medication-refractory HF who are not also candidates for cardiac transplantation, an LVAD, or a CRT defibrillator that incorporates both pacing and defibrillation capabilities.</li> </ol>

Figure 4



#### Recommendation-Specific Supportive Text

- The rationale for recommending that an ICD be offered to patients with NYHA class II or III HF, in addition to LVEF ≤35%, is based on the survival benefit observed in SCD-HeFT and MADIT-II (which used LVEF cutoff of below 35% and 30%, respectively). Selection for implantation of an ICD must be individualized. Patients with serious comorbidities associated with a survival of <1 year are generally not considered ICD candidates. The recommendation to wait at least 40 days after an MI before implanting a primary prevention ICD is based on the fact that such patients were excluded from MADIT-II and SCD-HeFT and 2 other RCTs showed no survival benefit from ICDs implanted early after an acute MI (S7.1.2-10,S7.1.2-11).
- In the MADIT-II trial (S7.1.2-2), which randomized patients with LVEF ≤30% and prior MI to an ICD or not, approximately one third of the patients had NYHA class I symptoms. A subgroup analysis supported benefit of the ICD on survival in this subgroup (S7.1.2-2).

- Economic outcomes of ICD implantation for primary prevention of SCD were assessed in 3 RCTs [MADIT-I (S7.1.2-12), MADIT-II (S7.1.2-13), and SCD-HeFT (S7.1.2-14)], 1 observational study (S7.1.2-15), and 4 simulation models (S7.1.2-16–S7.1.2-19), which all had generally consistent results. All studies 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 <\$50,000 per year of life added by an ICD, which provides high value according to the benchmarks adopted for the current guideline (S7.1.2-20). The value provided by an ICD was consistently high when life expectancy was projected to increase by >1.4 years (S7.1.2-18). In contrast, when survival was not increased by ICD implantation, as in the CABG-Patch trial (S7.1.2-18), the ICD did not provide value, because the higher costs were unaccompanied by a gain in life expectancy.
- MUSTT (Multicenter Unsustained Tachycardia Trial) demonstrated that patients with prior MI, NSVT, and

reduced LVEF with inducible VT at electrophysiological study have a higher overall mortality rate than similar patients without inducible sustained VT ([S7.1.2-21](#)). Patients who received an ICD after failing to have inducible VT suppressed by an antiarrhythmic medication had lower mortality rate than those who did not receive an ICD. Although the entry criteria into MUSTT required an LVEF of  $\leq 40\%$ , the average LVEF in enrolled patients was 30%, and ICD placement was not randomized but rather was selected by the treating physician for patients with VT that could not be suppressed with antiarrhythmic medication therapy. MUSTT allowed enrollment of patients who were  $\geq 4$  days after an acute MI or revascularization. The ICD was of no benefit in 2 other RCTs that examined the efficacy of the ICD in the acute phase of an MI ([S7.1.2-10](#), [S7.1.2-11](#)). In a single center observational study, an electrophysiological study was performed a median of 9 days after acute MI in 115 patients with LVEF  $<40\%$  and ICDs recommended for those with inducible VT. Median follow-up was 12 months. Sustained VT was induced in 27% of patients, and 22% of those who received ICDs had spontaneous VT terminated by the ICD during follow-up. None of the patients without inducible VT had VT or SCD during follow-up ([S7.1.2-22](#)).

5. In a retrospective analysis of the UNOS (United Network for Organ Sharing) registry that extended from 1999 to 2014, data on 32,599 patients showed that during a median follow-up of 154 days, 3,638 patients (11%) died while on the waitlist for cardiac transplantation (9% in the ICD group versus 15% in the non-ICD group;  $p < 0.0001$ ). The presence of an ICD at listing was associated with an adjusted 13% relative risk reduction in mortality. In the subgroup of patients with an LVAD ( $n = 9,478$ ), an ICD was associated with an adjusted 19% relative risk reduction in mortality ([S7.1.2-9](#)). In another study of 380 patients listed for heart transplantation between 2005 and 2009 at 1

tertiary heart transplant center, 122 patients received an ICD before or within 3 months after being listed for heart transplantation. Non-ICD patients were more likely to die while on the transplant list. In a multi-variable model, the ICD was not associated with improved survival; however, that analysis was limited by the small sample size ([S7.1.2-8](#)). Another small study ( $n = 79$ ) conducted at 1 institution suggested that ICDs reduce the risk of SCD in patients with LVEF  $\leq 30\%$  who are awaiting heart transplantation; however, this study was limited by the small number of patients ([S7.1.2-6](#)). In a retrospective multicenter study of 1,089 patients listed for heart transplantation, 550 patients (51%) had an ICD. In 216 patients, the ICD was for primary prevention of SCD and, in 334 patients, the ICD was for secondary prevention. The remaining 539 patients did not receive an ICD. During a median time on the waiting list of 8 months, the ICD was associated with a reduction in all-cause mortality in the primary and secondary prevention cohorts (estimated 1-year:  $88 \pm 3\%$  versus  $77 \pm 3\%$  versus  $67 \pm 3\%$ ;  $p = 0.0001$ ). This relationship between the ICD and improved survival persisted even after adjusting for potential confounders ([S7.1.2-7](#)).

- There are insufficient data from RCTs regarding the value of the ICD in patients with NYHA class IV HF. Ambulatory class IV patients with HF were included in the COMPANION (Comparison of Medical Therapy, Pacing, and Defibrillation in Heart Failure) trial, which showed an overall improved functional status and survival with a CRT defibrillator ([S7.1.2-23](#)). Unless such a patient is a candidate for CRT or advanced HF therapies such as heart transplantation or an LVAD, an ICD is not expected to meaningfully prolong survival ([S7.1.2-23](#)).

### 7.1.3. Treatment and Prevention of Recurrent VA in Patients With Ischemic Heart Disease

#### Recommendations for Treatment of Recurrent VA in Patients With Ischemic Heart Disease

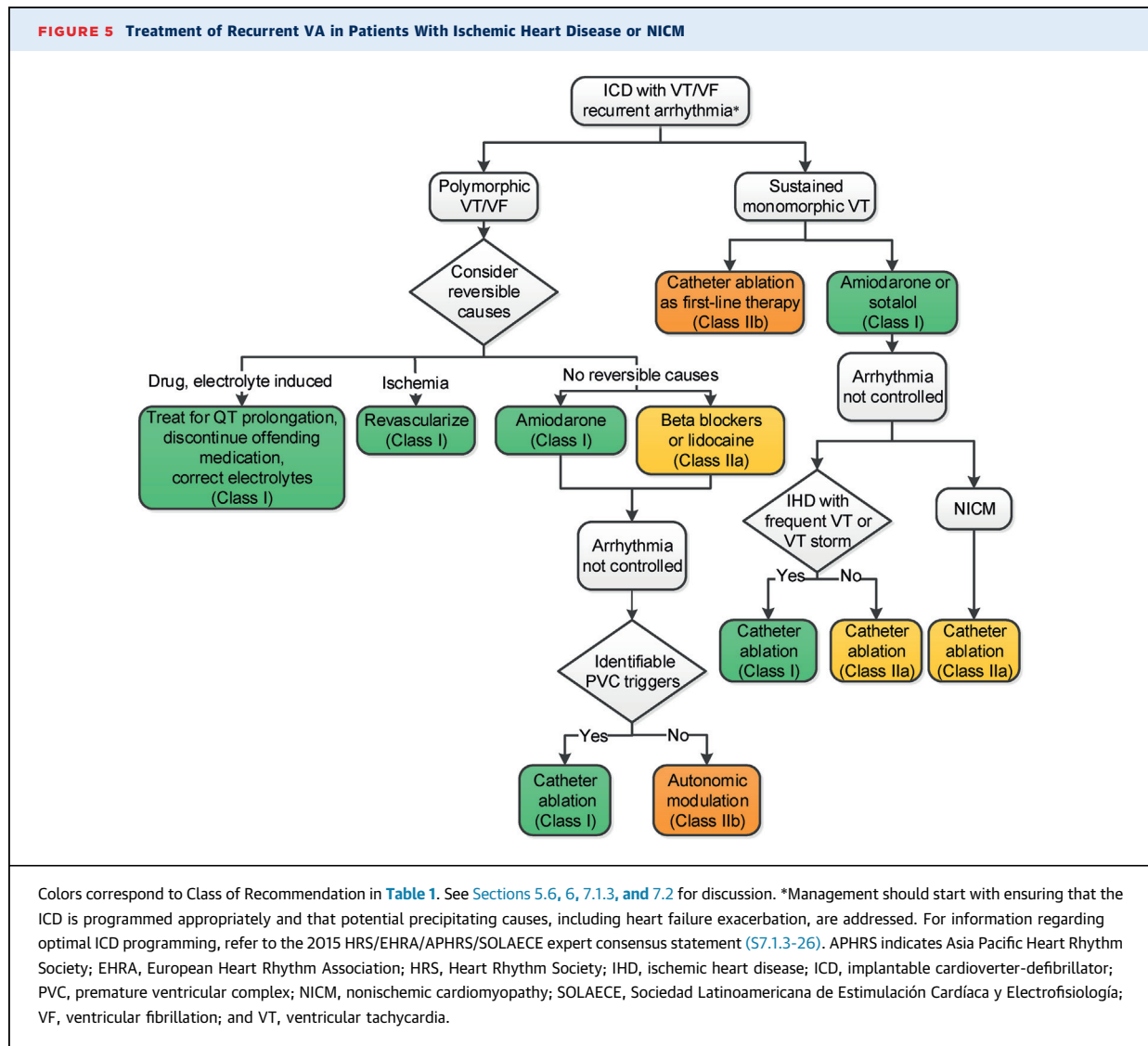
References that support the recommendations are summarized in [Online Data Supplement 22 and 23](#).

COR	LOE	RECOMMENDATIONS
I	B-R	<ol style="list-style-type: none"> <li>In patients with ischemic heart disease and recurrent VA, with significant symptoms or ICD shocks despite optimal device programming and ongoing treatment with a beta blocker, amiodarone or sotalol is useful to suppress recurrent VA (<a href="#">S7.1.3-1–S7.1.3-3</a>).</li> </ol>
I	B-R B-NR	<ol style="list-style-type: none"> <li>In patients with prior MI and recurrent episodes of symptomatic sustained VT, or who present with VT storm and have failed or are intolerant of amiodarone (LOE: B-R) (<a href="#">S7.1.3-4</a>) or other antiarrhythmic medications (LOE: B-NR) (<a href="#">S7.1.3-5–S7.1.3-9</a>), catheter ablation is recommended (<a href="#">S7.1.3-10–S7.1.3-12</a>).</li> </ol>

(continued)

IIb	C-LD	3. In patients with ischemic heart disease and ICD shocks for sustained monomorphic VT or symptomatic sustained monomorphic VT that is recurrent, or hemodynamically tolerated, catheter ablation as first-line therapy may be considered to reduce recurrent VA ( <a href="#">S7.1.3-10,S7.1.3-11</a> ).
III: Harm	B-R	4. In patients with prior MI, class IC antiarrhythmic medications (e.g., flecainide and propafenone) should not be used ( <a href="#">S7.1.3-13</a> ).
III: Harm	C-LD	5. In patients with incessant VT or VF, an ICD should not be implanted until sufficient control of the VA is achieved to prevent repeated ICD shocks ( <a href="#">S7.1.3-14</a> ).
III: No Benefit	C-LD	6. In patients with ischemic heart disease and sustained monomorphic VT, coronary revascularization alone is an ineffective therapy to prevent recurrent VT ( <a href="#">S7.1.3-15,S7.1.3-16</a> ).

Figure 5



### Recommendation-Specific Supportive Text

1. The most common antiarrhythmic medications used for suppression of VA include amiodarone and sotalol, while mexiletine, quinidine, and ranolazine are occasionally used ([S7.1.3-17](#),[S7.1.3-18](#)). Amiodarone appears to be more effective than sotalol and has a low rate of ventricular proarrhythmia, but has an increased risk of medication-related adverse effects that lead to its discontinuation in many patients within 18 to 24 months from initiation of therapy ([S7.1.3-1](#),[S7.1.3-19](#)). Data supporting effectiveness of sotalol for suppression of VA are conflicting, but given its more favorable adverse effect profile than amiodarone, it may be a better first-line antiarrhythmic medication in appropriate patients ([S7.1.3-1](#)–[S7.1.3-3](#)). However, sotalol is generally avoided in patients with a severely reduced LVEF <20% due to its negative inotropic effects and the risk of torsades de pointes. In a double-blind placebo-controlled study of 674 patients with HF and ≥10 PVCs/h and an LVEF ≤40% randomly assigned to receive amiodarone (336 patients) or placebo (338 patients), there was no significant difference in overall mortality or SCD between the 2 arms. There was a trend toward a reduction in overall mortality among the patients with NICM who received amiodarone ( $p=0.07$ ) ([S7.1.3-20](#)).
2. Patients with prior MI may present with frequent episodes of sustained monomorphic VT or recurrent VF episodes that are initiated by PVCs arising from Purkinje Fibers in the peri-infarct zone. VA storms are associated with increased mortality ([S7.1.3-12](#)). The arrhythmia substrate is usually in the sub-endocardium. The randomized VANISH (Ventricular Tachycardia Ablation versus Escalated Antiarrhythmic Drug Therapy in Ischemic Heart Disease) trial ([S7.1.3-4](#)) compared escalating antiarrhythmic medication therapy versus catheter ablation for patients with prior MI and recurrent sustained monomorphic VT despite antiarrhythmic medications. The primary outcome, a composite of death, VT storm, or ICD shocks occurred in 59.1% in the ablation group and in 68.5% in the escalated-therapy group. There was no difference in mortality between the groups. Recurrent ICD shocks and VT storm and treatment-related adverse events were lower in the ablation group. In a subgroup analysis, patients having VT on amiodarone had better outcomes with ablation compared with increasing amiodarone or adding mexiletine to amiodarone. For patients receiving medications other than amiodarone, catheter ablation did not reduce the risk of ICD shocks or VT storm compared with switching to amiodarone. Although recurrent VT after catheter ablation is associated with increased mortality ([S7.1.3-9](#)), whether

mortality is reduced by catheter ablation has not been established. Procedural complications occur in approximately 6% of patients, most of which are related to vascular access but stroke, tamponade, and atrioventricular block can occur. Procedure mortality is <1% in experienced centers ([S7.1.3-4](#),[S7.1.3-9](#)).

Sustained monomorphic VT often occurs as occasional isolated episodes in patients with prior MI. Several nonrandomized studies have shown that catheter ablation reduces recurrent VT or ICD shocks ([S7.1.3-5](#),[S7.1.3-7](#),[S7.1.3-8](#)). A meta-analysis of 5 VT ablation studies ([S7.1.3-5](#)) reported that VT recurred in 35% of patients after catheter ablation compared with 55% on antiarrhythmic medications. In a multicenter study of catheter ablation ([S7.1.3-7](#)) for patients with ≥3 episodes of sustained VT in the prior 6 months, 53% were free from recurrent VT at 6 months follow-up; the median number of VT episodes was reduced from 11.5 to 0. Superiority of ablation over escalating medication therapy was shown in the composite endpoint of death, VT storm, or ICD shocks by the VANISH trial ([S7.1.3-4](#)).

3. Patients with prior MI who develop sustained monomorphic VT often have recurrent episodes. The VTACH (Ventricular Tachycardia Ablation in Addition to Implantable Defibrillators in Coronary Heart Disease) trial ([S7.1.3-11](#)) randomized patients undergoing ICD implantation for stable sustained monomorphic VT, who had not failed antiarrhythmic medication therapy, to catheter ablation versus ICD implantation alone. At 2 years, any VT had recurred in 53% of the ablation group and 71% of the control group. Ablation prolonged the time to recurrent VT from a median of 5.9 months to 18.6 months ([S7.1.3-11](#)). Several nonrandomized studies have shown that catheter ablation reduces the risk of recurrent VT or ICD shocks in patients with sustained VT related to prior MI ([S7.1.3-5](#),[S7.1.3-7](#),[S7.1.3-8](#)). In a multicenter study of catheter ablation ([S7.1.3-7](#)) for patients with ≥3 episodes of sustained VT in the prior 6 months, 53% were free from recurrent VT at 6 months follow-up; the median number of VT episodes was reduced from 11.5 to 0. A meta-analysis of 5 VT ablation studies ([S7.1.3-5](#)) reported that VT recurred in 35% of patients after catheter ablation compared with 55% on antiarrhythmic medications. Another study of 63 patients with recurrent VT after MI demonstrated acute success with catheter ablation in 83% of mappable VTs and 40% of nonmappable VTs ([S7.1.3-8](#)). Superiority of ablation over escalating medication therapy for patients with recurrent VT despite antiarrhythmic medications was shown by the VANISH trial ([S7.1.3-4](#)). See Section 5.6.

4. CAST ([S7.1.3-21](#)) demonstrated higher rates of mortality or nonfatal cardiac arrest in post-MI patients treated with encainide or flecainide when used to suppress PVCs and NSVT ([S7.1.3-13](#)). Propafenone is associated with increased mortality in SCA survivors compared with beta blockers, amiodarone, and the ICD ([S7.1.3-22](#)).
5. Implantation of an ICD prior to achieving suppression of frequent or incessant VA places the patient at high risk of repetitive shocks, which can be psychologically detrimental and has been associated with increased mortality ([S7.1.3-23](#),[S7.1.3-24](#)).
6. Sustained monomorphic VT in the setting of prior MI is typically due to scar-related reentry and is not due to

acute ischemia. Although it may be appropriate to recommend revascularization when another indication for revascularization exists, revascularization alone is unlikely to reduce the recurrence of monomorphic VT and specific therapies such as antiarrhythmic medications or ablation may be needed to prevent recurrence ([S7.1.3-16](#)). On the contrary, revascularization might be beneficial in patients with ischemic heart disease and VF, polymorphic VT, or exercise-induced arrhythmias associated with ischemia ([S7.1.3-25](#)).

## 7.2. Nonischemic Cardiomyopathy

### Recommendations for Patients With NICM

References that support the recommendations are summarized in [Online Data Supplement 24](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with suspected NICM from myocardial infiltrative processes, cardiac MRI with late gadolinium enhancement is useful for diagnosis ( <a href="#">S7.2-1–S7.2-3</a> ).
IIa	B-NR	2. In patients with suspected NICM, cardiac MRI with late gadolinium enhancement can be useful for assessing risk of SCA/SCD ( <a href="#">S7.2-1–S7.2-3</a> ).
IIa	C-EO	3. In patients with NICM who develop conduction disease or LV dysfunction at less than 40 years of age, or who have a family history of NICM or SCD in a first-degree relative (<50 years of age), genetic counseling and genetic testing are reasonable to detect a heritable disease that may clarify prognosis and facilitate cascade screening of relatives ( <a href="#">S7.2-4</a> , <a href="#">S7.2-5</a> ).

### Recommendation-Specific Supportive Text

1. Cardiac MRI allows for evaluation of structural heart disease and assessment of LV and RV function including quantification of LVEF, LV mass and volume, and valvular structure. Cardiac MRI can help in the evaluation for myocardial infiltrative processes and evidence of scar, indicated by delayed hyperenhancement, associated with VA ([S7.2-1](#)–[S7.2-4](#),[S7.2-6](#)).
2. The presence of delayed hyperenhancement has been associated with worse outcomes, including SCD ([S7.2-1](#)–[S7.2-3](#)).

3. It is important to consider genetic etiologies for NICM. Goals of genetic testing for NICM are to identify at-risk relatives who host a disease-causing mutation and to help clarify prognosis. *Lamin A/C* and *NKX 2.5* mutations ([S7.2-7](#)–[S7.2-12](#)) are associated with a particularly high risk of early conduction disease, arrhythmias, and SCD, and their identification often prompts consideration of early use of an ICD. It is unknown, however, whether early pharmacological treatment of mutation-positive, asymptomatic subjects can prevent or delay manifestation of the disease or whether genetic testing ultimately improves survival.

### 7.2.1. Secondary Prevention of SCD in Patients With NICM

#### Recommendations for Secondary Prevention of SCD in Patients With NICM

References that support the recommendations are summarized in [Online Data Supplement 25 and 26](#).

COR	LOE	RECOMMENDATIONS
I	B-R B-NR	<p>1. In patients with NICM who either survive SCA due to VT/VF or experience hemodynamically unstable VT (LOE: B-R) (<a href="#">S7.2.1-1–S7.2.1-4</a>) or stable sustained VT (LOE: B-NR) (<a href="#">S7.2.1-5</a>) not due to reversible causes, an ICD is recommended if meaningful survival greater than 1 year is expected.</p>
IIa	B-NR	<p>2. In patients with NICM who experience syncope presumed to be due to VA and who do not meet indications for a primary prevention ICD, an ICD or an electrophysiological study for risk stratification for SCD can be beneficial if meaningful survival greater than 1 year is expected (<a href="#">S7.2.1-6–S7.2.1-11</a>).</p>
IIb	B-R	<p>3. In patients with NICM who survive a cardiac arrest, have sustained VT, or have symptomatic VA who are ineligible for an ICD (due to a limited life-expectancy and/or functional status or lack of access to an ICD), amiodarone may be considered for prevention of SCD (<a href="#">S7.2.1-12,S7.2.1-13</a>).</p>

*Figure 6*

#### Recommendation-Specific Supportive Text

- Three prospective RCTs compared the ICD with pharmacological therapy in patients resuscitated from SCA due to VT/VF or hemodynamically significant VT ([S7.2.1-1,S7.2.1-2,S7.2.1-4](#)). The antiarrhythmic medications most commonly used were amiodarone, a beta blocker, or both, although in the CASH (Cardiac Arrest Study Hamburg) trial ([S7.2.1-4](#)), there was also a propafenone arm that was terminated early due to increased mortality. The 3 trials enrolled 1,963 patients, but only 292 (14.8%) had NICM. A meta-analysis in which data from AVID and CIDS were pooled found a nonsignificant 31% reduction in all-cause mortality relative to medical therapy in patients with NICM ([S7.2.1-3](#)). Although this analysis was underpowered, the observed mortality reduction was consistent with the observed benefit in the entire study population. In the AVID trial ([S7.2.1-1](#)), patients who were ineligible for the RCT were included in a registry, and sustained VT without serious symptoms or hemodynamic compromise was associated with a mortality rate similar to that of patients with unstable VT who were assigned to medical therapy. Therefore, stable VT is likely a marker for a substrate capable of producing subsequent lethal arrhythmias ([S7.2.1-5](#)).
- Small observational studies demonstrated high mortality and frequent appropriate ICD shocks in patients with syncope and NICM ([S7.2.1-7–S7.2.1-9](#)). The assumption that malignant VAs are the likely cause of syncope and that the ICD would be protective has recently been challenged. In a subgroup analysis of SCD-HeFT that included 472 patients, the ICD did not reduce either recurrent syncope or the increased risk of

mortality associated with syncope ([S7.2.1-10](#)). A subgroup analysis of the MADIT- RIT (Multicenter Automatic Defibrillator Implantation Trial - Reduce Inappropriate Therapy) trial found syncope to be arrhythmic only in 39% of patients ([S7.2.1-11](#)). These studies suggest that syncope in some HF patients may be an indicator of an end-stage cardiomyopathy associated with a poor prognosis ([S7.2.1-11](#)). In a substudy of DEFINITE, inducible sustained VT/VF was found in a minority of patients, but it was associated with appropriate ICD therapy ([S7.2.1-14](#)). Another study of electrophysiological testing in NICM found inducible VT/VF in 27.8% of patients, which was associated with future ICD events ([S7.2.1-15](#)). In a study of patients with NICM, cardiac mortality correlated with LVEF but not with inducibility on electrophysiological study ([S7.2.1-16](#)). Based on these data, many experts are uncomfortable withholding an ICD from patients with NICM who experience syncope potentially due to a VA even if the electrophysiological study shows no inducible sustained VT.

- Access to ICDs may be limited by financial, medical, or personal considerations. In addition, not all patients at high risk of SCD meet ICD indications, such as those with class IV HF without CRT possibility or with a life expectancy <1 year. A meta-analysis of RCTs, which examined the use of amiodarone for the prevention of SCD, included 15 studies with 8522 patients assigned to amiodarone or placebo/control ([S7.2.1-12](#)). Amiodarone reduced the risk of SCD by 29%; however, it did not reduce all-cause mortality and was associated with an increased risk of pulmonary and thyroid toxicity. In a subgroup analysis, the benefit of amiodarone appeared similar in patients with ischemic cardiomyopathy and those with NICM ([S7.2.1-12](#)). In a separate meta-analysis ([S7.2.1-13](#)), the evidence was insufficient to support

amiodarone's efficacy for reduction of SCD and all-cause mortality in survivors of cardiac arrest or those with syncope due to VA. A subgroup analysis of the VALIANT (Valsartan in Acute Myocardial Infarction) trial found that amiodarone was associated with increased mortality in patients with NYHA class III HF

(S7.2.1-17). These data call for a careful and nuanced approach to using amiodarone for the secondary prevention of SCD in patients with NICM.

## 7.2.2. Primary Prevention of SCD in Patients With NICM

### Recommendations for Primary Prevention of SCD in Patients With NICM

References that support the recommendations are summarized in [Online Data Supplement 27 and 28](#).

COR	LOE	RECOMMENDATIONS
I	A	1. In patients with NICM, HF with NYHA class II-III symptoms and an LVEF of 35% or less, despite GDMT, an ICD is recommended if meaningful survival of greater than 1 year is expected (S7.2.2-1–S7.2.2-6).
IIa	B-NR	2. In patients with NICM due to a <i>Lamin A/C</i> mutation who have 2 or more risk factors (NSVT, LVEF <45%, nonmissense mutation, and male sex), an ICD can be beneficial if meaningful survival of greater than 1 year is expected (S7.2.2-7–S7.2.2-10).
IIb	B-R	3. In patients with NICM, HF with NYHA class I symptoms and an LVEF of 35% or less, despite GDMT, an ICD may be considered if meaningful survival of greater than 1 year is expected (S7.2.2-5).
III: No Benefit	C-EO	4. In patients with medication-refractory NYHA class IV HF who are not also candidates for cardiac transplantation, an LVAD, or a CRT defibrillator that incorporates both pacing and defibrillation capabilities, an ICD should not be implanted.

### Figure 6 Recommendation-Specific Supportive Text

1. For all patients with NICM, it is imperative that patients be on GDMT for HF for at least 3 months before a primary prevention ICD is offered. Four prospective RCTs (S7.2.2-1, S7.2.2-2, S7.2.2-5, S7.2.2-6) initially evaluated ICDs for primary prevention of SCD in patients with NICM. Two (S7.2.2-2, S7.2.2-6) were small studies that were terminated early due to a low event rate. In DEFINITE (S7.2.2-5), an ICD reduced the risk of SCD, with a trend toward reduced all-cause mortality. SCD-HeFT included 792 NICM patients (S7.2.2-1). Total mortality at 5 years was 27% in the placebo group and 21% in the ICD group ( $p=0.06$ ). A pooled analysis of these studies demonstrated a significant 31% reduction in all-cause mortality for ICD relative to medical therapy (S7.2.2-4). The DANISH (Danish Study to Assess the Efficacy of ICDs in Patients with Non-ischemic Systolic Heart Failure on Mortality) trial (S7.2.2-11) raised questions about the role of primary prevention ICDs in patients with NICM. This trial randomized 1116 patients with NICM LVEF <35% and class II, III, or IV (if CRT was planned) HF to an ICD or no ICD. CRT (either ICD or pacemaker) was present in 58% of patients in the ICD and medical therapy arms. Therefore, the results of DANISH should not be generalized to patients with NICM who are ineligible for CRT. During a median follow-up of 5.6 years, ICD reduced SCD from 8.4% to

4.3%, but there was no difference in all-cause mortality (S7.2.2-11). Several meta-analyses have been published (S7.2.2-12, S7.2.2-13). One provided data on ICDs with and without CRT and showed survival benefit from the ICD (S7.2.2-13). The second used patient level data from 2 trials and adopted a more robust approach to reducing heterogeneity by excluding patients with CRT and those randomized to antiarrhythmic medications; a 25% relative risk reduction in mortality with an ICD was shown (S7.2.2-12).

2. Laminopathies are diseases caused by mutations mainly in the *Lamin A/C* gene that produce various inherited diseases including subtypes of muscular dystrophy and progeria. Isolated cardiac involvement is also observed and is an important cause of familial cardiomyopathy (S7.2.2-9). The disease is highly penetrant such that all affected individuals have evidence of disease by 60 years of age. Cardiac manifestations may include atrial fibrillation, conduction disturbances, VA, and NICM. A number of observational studies reported a high risk of SCD when cardiac involvement is present (S7.2.2-7–S7.2.2-10). One study reported SCD as the most frequent mode of death (46%) in both the isolated cardiac and the neuromuscular phenotypes of *Lamin* diseases (S7.2.2-9). In a cohort of 269 *LMNA* mutation positive individuals (S7.2.2-10), NSVT during ambulatory electrocardiographic monitoring, LVEF <45% at first evaluation, male sex, and nonmissense mutations

were independent risk factors for VA. Malignant VA were observed only in persons with  $\geq 2$  of these risk factors ([S7.2.2-10](#)). No studies have tested the effect of the ICD on long-term survival.

3. Patients with NICM and class I HF symptoms were not included in SCD-HeFT or DANISH ([S7.2.2-1,S7.2.2-11](#)). Although such patients were included in the DEFINITE trial, only 99 (21.6%) of 458 patients in the DEFINITE trial had class I HF ([S7.2.2-5](#)). Therefore, it is uncertain whether a primary prevention ICD in such patients improves survival.

4. There are insufficient data from RCTs regarding the value of the ICD in patients with NYHA class IV. Ambulatory class IV HF patients were included in the COMPANION trial that, overall, showed improved functional status and survival with a CRT defibrillator ([S7.2.2-3](#)). Unless such a patient is a candidate for CRT or advanced HF therapies such as heart transplantation or an LVAD, an ICD is not expected to meaningfully prolong survival ([S7.2.2-3](#)).

### 7.2.3. Treatment of Recurrent VA in Patients With NICM

#### Recommendations for Treatment of Recurrent VA in Patients With NICM

References that support the recommendations are summarized in [Online Data Supplement 29](#).

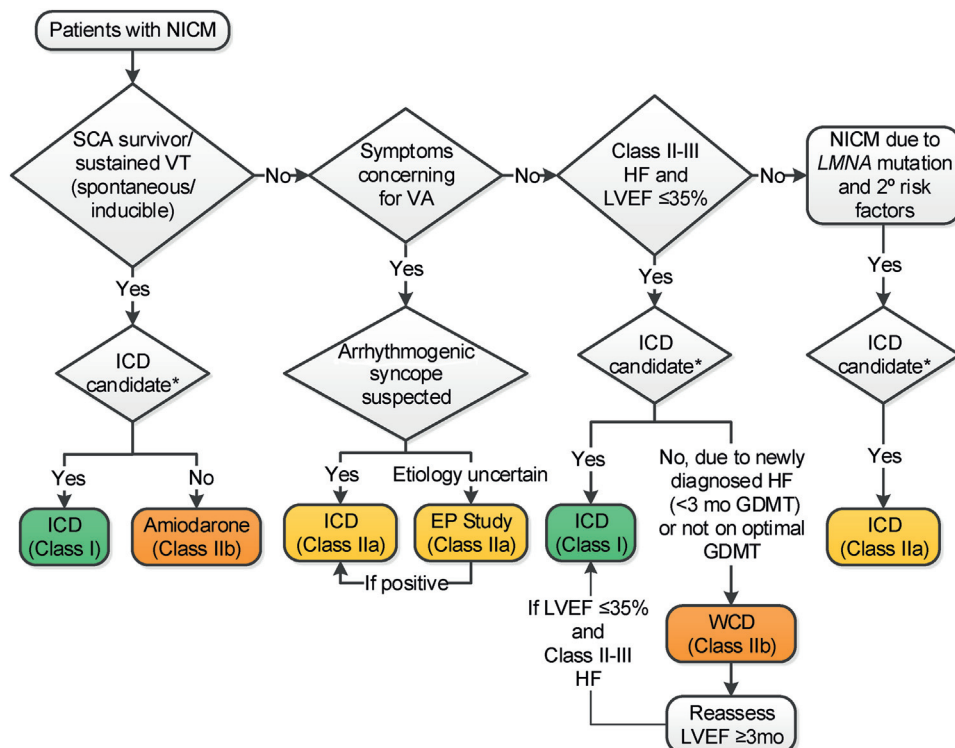
COR	LOE	RECOMMENDATIONS
IIa	B-R	<ol style="list-style-type: none"> <li>1. In patients with NICM and an ICD who experience spontaneous VA or recurrent appropriate shocks despite optimal device programming and treatment with a beta blocker, amiodarone or sotalol can be beneficial (<a href="#">S7.2.3-1</a>).</li> </ol>
IIa	B-NR	<ol style="list-style-type: none"> <li>2. In patients with NICM and recurrent sustained monomorphic VT who fail or are intolerant of antiarrhythmic medications, catheter ablation can be useful for reducing recurrent VT and ICD shocks (<a href="#">S7.2.3-2,S7.2.3-3</a>).</li> </ol>

#### Recommendation-Specific Supportive Text

1. ICDs reduce mortality from VA, yet ICD shocks are painful and associated with significant morbidity and poor QoL. Although ICDs are highly programmable and provide antitachycardia pacing therapy that can terminate most VT episodes without the need for a shock, prevention of shocks, both appropriate and inappropriate, remains an important concern. In the OPTIC (Optimal Pharmacological Therapy in Cardioverter Defibrillator Patients) study, 412 patients with documented VT and VF who received an ICD within 21 days of the documented arrhythmia ([S7.2.3-1](#)) were randomized to amiodarone plus beta blocker, sotalol alone, or beta blocker alone. Over 1 year, shocks occurred in 38.5% assigned to beta blocker alone, 24.3% assigned to sotalol, and 10.3% assigned to amiodarone plus beta blocker. The rates of study medication discontinuation at 1 year were 18.2% for amiodarone, 23.5% for sotalol, and 5.3% for beta blocker alone. Adverse pulmonary and thyroid events and symptomatic bradycardia were more common among patients randomized to amiodarone. Thus, amiodarone plus beta blocker were more effective than sotalol in preventing ICD shocks but at the expense of increased risk of medication-related adverse effects ([S7.2.3-1](#)). Sotalol should not be used in patients with an LVEF  $<20\%$  due to its negative inotropic effects.

2. Sustained monomorphic VT due to NICM is most often due to scar-related reentry. Cardiac MRI often indicates scar location, which tends to be basal along the mitral annulus or in the septum ([S7.2.3-4,S7.2.3-5](#)). The VT substrate can be subendocardial, subepicardial, or intramyocardial, and all locations may be affected and require endocardial and epicardial ablation. In the HELP-VT (Heart Center of Leipzig VT) study ([S7.2.3-2](#)), successful ablation of all VT morphologies was achieved in 66.7% of patients with NICM, compared with the 77.4% success rate in ischemic cardiomyopathy. An epicardial approach to ablation was required in 30.2% of NICM patients, compared with only 1.2% with ischemic cardiomyopathy. Epicardial ablation was an independent predictor of successful ablation. Acute and long-term success of ablation is lower for NICM, compared with post-MI patients. The long-term survival-free of VT recurrence after catheter ablation appears to be better for patients with ischemic than NICM (57% versus 40.5% at 1 year) ([S7.2.3-2](#)). Risks are similar to those observed for post-MI VT ablation, with additional risks of epicardial access and ablation when required. Although any NICM can produce scar-related VT, cardiac sarcoidosis (see [Section 7.6](#)) and Lamin mutations are particularly associated with sustained monomorphic VT ([S7.2.3-6](#)).

**FIGURE 6 Secondary and Primary Prevention of SCD in Patients With NICM**



Colors correspond to Class of Recommendation in **Table 1**. See **Section 7.2** for discussion. \*ICD candidacy as determined by functional status, life expectancy or patient preference. 2° indicates secondary; EP, electrophysiological; GDMT, guideline-directed management and therapy; HF, heart failure; ICD, implantable cardioverter-defibrillator; LVEF, left ventricular ejection fraction; NICM, nonischemic cardiomyopathy; SCA, sudden cardiac arrest; SCD, sudden cardiac death; VA, ventricular arrhythmia; and WCD, wearable cardiac-defibrillator.

### 7.3. Arrhythmogenic Right Ventricular Cardiomyopathy

#### Recommendations for Arrhythmogenic Right Ventricular Cardiomyopathy

References that support the recommendations are summarized in [Online Data Supplement 30](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	<ol style="list-style-type: none"> <li>In selected first-degree relatives of patients with arrhythmogenic right ventricular cardiomyopathy, clinical screening for the disease is recommended along with genetic counseling and genetic testing, if the proband has a disease causing mutation (<a href="#">S7.3-1–S7.3-4</a>).</li> </ol>
I	B-NR	<ol style="list-style-type: none"> <li>In patients with suspected arrhythmogenic right ventricular cardiomyopathy and VA or electrocardiographic abnormalities, cardiac MRI is useful for establishing a diagnosis and for risk stratification (<a href="#">S7.3-5–S7.3-8</a>).</li> </ol>
I	B-NR	<ol style="list-style-type: none"> <li>In patients with arrhythmogenic right ventricular cardiomyopathy and an additional marker of increased risk of SCD (resuscitated SCA, sustained VT, significant ventricular dysfunction with RVEF or LVEF ≤35%), an ICD is recommended if meaningful survival greater than 1 year is expected (<a href="#">S7.3-9–S7.3-13</a>).</li> </ol>

(continued)

I	B-NR	4. In patients with arrhythmogenic right ventricular cardiomyopathy and VA, a beta blocker is recommended (S7.3-11,S7.3-14,S7.3-15).
I	B-NR	5. In patients with a clinical diagnosis of arrhythmogenic right ventricular cardiomyopathy, avoiding intensive exercise is recommended (S7.3-11,S7.3-12,S7.3-16–S7.3-21).
IIa	B-NR	6. In patients with clinically diagnosed or suspected arrhythmogenic right ventricular cardiomyopathy, genetic counseling and genetic testing can be useful for diagnosis and for gene-specific targeted family screening (S7.3-1,S7.3-4,S7.3-22–S7.3-26).
IIa	B-NR	7. In patients with arrhythmogenic right ventricular cardiomyopathy and syncope presumed due to VA, an ICD can be useful if meaningful survival greater than 1 year is expected (S7.3-10,S7.3-11,S7.3-13).
IIa	B-NR	8. In patients with clinical evidence of arrhythmogenic right ventricular cardiomyopathy but not VA, a beta blocker can be useful (S7.3-14,S7.3-15).
IIa	B-NR	9. In patients with arrhythmogenic right ventricular cardiomyopathy and recurrent symptomatic sustained VT in whom a beta blocker is ineffective or not tolerated, catheter ablation with availability of a combined endocardial/epicardial approach can be beneficial (S7.3-27–S7.3-33).
IIa	B-NR	10. In patients with suspected arrhythmogenic right ventricular cardiomyopathy, a signal averaged ECG can be useful for diagnosis and risk stratification (S7.3-14,S7.3-34,S7.3-35).
IIb	B-NR	11. In asymptomatic patients with clinical evidence of arrhythmogenic right ventricular cardiomyopathy, an electrophysiological study may be considered for risk stratification (S7.3-9,S7.3-36).

### Synopsis

Arrhythmogenic right ventricular cardiomyopathy is an inherited cardiomyopathy that predominantly affects the right ventricle but can affect the left ventricle causing areas of myocardial replacement with fibrosis and adipose tissue that frequently causes VA and SCD.

### Recommendation-Specific Supportive Text

1. Selected first-degree relatives refers to relatives who are willing to undergo further testing and who could benefit from further screening and testing (and not the terminally ill patients or those who do not want to be screened and tested). Arrhythmogenic right ventricular cardiomyopathy is often due to a mutation involving a desmosomal protein, and it usually has autosomal dominant inheritance with variable penetrance. SCD can be the initial manifestation of arrhythmogenic right ventricular cardiomyopathy. Clinical screening with ECG, cardiac imaging, and ambulatory rhythm monitoring and/or exercise testing may identify family members at risk for arrhythmogenic right ventricular cardiomyopathy. Arrhythmogenic right ventricular cardiomyopathy is detected clinically in approximately 35% to 40% of first-degree relatives (S7.3-3,S7.3-4), most commonly in siblings or symptomatic first-degree relatives (S7.3-4). When a

proband is identified with a disease-causing mutation, targeted genotype screening can identify mutation positive relatives (S7.3-1), with approximately 35% of mutation positive individuals ultimately developing progressive disease expression (S7.3-1,S7.3-4). In studies of arrhythmogenic right ventricular cardiomyopathy mutation-positive individuals who do not initially manifest the disease, 8% to 16% have a major arrhythmic event over the next 7 to 39 years (S7.3-1,S7.3-4,S7.3-26). Early identification of affected or potentially affected family members can allow lifestyle modifications in sports participation and serial monitoring for development of electrocardiographic abnormalities, symptoms, ventricular dysfunction, or arrhythmia. As genetic testing for arrhythmogenic right ventricular cardiomyopathy has subtle complexities, the decision to proceed with family screening is facilitated by informed genetic counseling to discuss the cost of testing, the potential lack of a single gene as the determinant for disease expression, psychological implications of uncertain disease progression, and implications for lifestyle modification, screening, and potential treatment.

2. Cardiac MRI provides high-quality assessment of ventricular function, size, regional wall motion abnormalities, and extent of scar and fibrosis

(late gadolinium enhancement) that are seen in 30% to 95% of patients with the clinical diagnosis of arrhythmogenic right ventricular cardiomyopathy ([S7.3-5,S7.3-6,S7.3-37,S7.3-38](#)). Cardiac MRI detects biventricular involvement in 34% to 56% of patients, with isolated LV involvement noted in 4% to 9% of patients ([S7.3-37-S7.3-40](#)). Cardiac MRI should include assessment of late gadolinium enhancement with quantification of fibrosis. Application of the 2010 Task Force Criteria to cardiac MRI criteria for diagnosis of arrhythmogenic right ventricular cardiomyopathy has improved the specificity of this test ([S7.3-5, S7.3-8](#)). Electrocardiographic and Holter findings precede detectable cardiac MRI abnormalities in arrhythmogenic right ventricular cardiomyopathy mutation-positive individuals, with only 4% of patients with normal electrocardiographic and Holter results having cardiac MRI abnormalities, suggesting that evaluation of cardiac structure and function using cardiac MRI may be unnecessary in mutation-positive individuals who do not have electrical abnormalities ([S7.3-7](#)). The presence of both electrocardiographic abnormalities and abnormal cardiac MRI findings may identify patients at an increased risk for developing sustained VA ([S7.3-7,S7.3-38](#)). Areas of scar identified on cardiac MRI have correlated with the location of VT substrate identified by endocardial and epicardial mapping ([S7.3-38](#)). During early stages of disease, a baseline cardiac MRI may provide useful information along with electrocardiographic and rhythm abnormalities to monitor disease progression over time. Experience and expertise in interpretation of cardiac MRI are important ([S7.3-5,S7.3-8](#)).

3. Arrhythmogenic right ventricular cardiomyopathy is characterized by progressive ventricular myocyte loss with replacement by fatty or fibrous tissue, and is associated with progressive ventricular dysfunction that may involve both ventricles. VA, syncope, and SCD may occur at a relatively young age, particularly in the second and third decades of life and often occurring during physical activity ([S7.3-1,S7.3-16,S7.3-22, S7.3-41](#)). Sustained VT is an important predictor of SCA and SCD or appropriate ICD shocks in patients with arrhythmogenic right ventricular cardiomyopathy ([S7.3-10,S7.3-13](#)). In patients receiving an ICD for primary prevention, appropriate ICD shocks are reported in 24% to 48% of patients ([S7.3-9,S7.3-10, S7.3-12,S7.3-13](#)). As sustained VT in arrhythmogenic right ventricular cardiomyopathy patients is monomorphic in 55% to 90% of episodes based on ICD interrogation or electrophysiological studies ([S7.3-12,S7.3-36](#)), antitachycardia pacing algorithms are used to terminate VT.

4. Frequent PVCs, >760 to 1000 per 24 hours during ambulatory rhythm monitoring, correlate with arrhythmic risk ([S7.3-9,S7.3-23](#)). The presence of NSVT or sustained VT is an important predictor of adverse cardiac events ([S7.3-9,S7.3-12,S7.3-13,S7.3-42,S7.3-43](#)). The increased arrhythmia risk conferred by intense exercise is consistent with beta-adrenergic modulation of disease expression ([S7.3-17,S7.3-20,S7.3-21](#)). An observational registry reported that treatment with atenolol or amiodarone was associated with less clinically relevant VA, while sotalol was associated with no effect or increased arrhythmia ([S7.3-15](#)). Ambulatory monitoring to assess VA burden and adequacy of beta-blocker therapy is usually used ([S7.3-9, S7.3-14,S7.3-23,S7.3-42](#)).
5. Patients with arrhythmogenic right ventricular cardiomyopathy have a significantly increased risk of SCD during exertion ([S7.3-16,S7.3-17,S7.3-20,S7.3-21](#)). Vigorous exercise in patients with arrhythmogenic right ventricular cardiomyopathy has been shown to impair myocardial function by echocardiography and cardiac MRI ([S7.3-19](#)). Participation in high intensity/duration or endurance physical activity accelerates the penetrance/disease progression and arrhythmic risk for arrhythmogenic right ventricular cardiomyopathy patients and mutation positive individuals, as well as mutation positive family members ([S7.3-17, S7.3-19-S7.3-21](#)). Patients with arrhythmogenic right ventricular cardiomyopathy who participate in competitive sports are at increased risk for VT or SCD, compared with those who participate in recreational sports or are inactive ([S7.3-17-S7.3-19,S7.3-21](#)). Exercise influences disease progression in a linear manner; family members who limited activity to less than the AHA recommended minimum for activity guidelines (<650 metabolic equivalent hours per year [MET-Hr/year]) were less likely to develop VA or disease progression ([S7.3-21](#)). In a study of arrhythmogenic right ventricular cardiomyopathy probands and exercise, athletes (defined as subjects with ≥4 h vigorous exercise/week) were found to have reduced biventricular function compared with nonathletes in arrhythmogenic right ventricular cardiomyopathy patients and in mutation-positive family members ([S7.3-19](#)). Many advise limiting exercise intensity and duration to <650 MET-Hr/year, or 12.5 MET-Hr/week ([S7.3-21](#)).
6. The proband with arrhythmogenic right ventricular cardiomyopathy is usually diagnosed by the presence of clinical symptoms along with the presence of arrhythmogenic right ventricular cardiomyopathy Task Force criteria including: abnormalities on ECG, structural and functional changes of either ventricle, arrhythmias, and arrhythmogenic right ventricular

- cardiomyopathy in first-degree relatives ([S7.3-6](#)). A pathogenic genetic mutation was added to the major Task Force criteria in 2010 ([S7.3-44](#)). The yield of genetic testing in probands with suspected arrhythmogenic right ventricular cardiomyopathy is generally 30% to 54%, and is up to 58% among patients with a strong family history of SCD in multiple members ([S7.3-3,S7.3-25,S7.3-45](#)). A negative genetic test for arrhythmogenic right ventricular cardiomyopathy does not exclude the disease, and a positive genetic test currently does not guide therapy ([S7.3-22](#)). For the proband with a clinical diagnosis of arrhythmogenic right ventricular cardiomyopathy, identification of pathogenic mutations provides limited prognostic information relative to the risk of VT/VF ([S7.3-22,S7.3-26](#)) or development of HF ([S7.3-22](#)). In a large multicenter study, the presence of positive mutations among probands was not associated with a difference in mortality or cardiac transplantation ([S7.3-1](#)). However, the identification of a pathogenic mutation facilitates targeted genetic screening for that mutation in first-degree relatives, that may identify approximately 60% to 70% as gene positive ([S7.3-1](#)), highest among siblings, and those with symptoms ([S7.3-4](#)). Screening for the specific mutation can identify some gene positive family members prior to disease expression, while relieving others from the need for lifestyle changes and long-term monitoring ([S7.3-2,S7.3-3](#)).
7. Syncope is reported in 16% to 39% of arrhythmogenic right ventricular cardiomyopathy patients at the time of diagnosis ([S7.3-13,S7.3-14,S7.3-16,S7.3-41,S7.3-43](#)), is frequently exercise-related, and has been associated with high arrhythmic risk in some studies ([S7.3-10,S7.3-41](#)). Among patients with arrhythmogenic right ventricular cardiomyopathy and implanted ICDs, syncope was an important predictor of appropriate shocks in 1 study ([S7.3-10](#)), but not in other studies ([S7.3-9,S7.3-12,S7.3-13,S7.3-43](#)). Studies have not provided information about ventricular function or abnormalities on ECG in patients with syncope, limiting its assessment as an independent risk factor. Syncope may be a harbinger of progression of underlying disease and should be integrated into the decision-making process for ICD implantation with the patient.
8. Asymptomatic patients with arrhythmogenic right ventricular cardiomyopathy and no VA or ventricular dysfunction are generally observed without antiarrhythmic therapy other than beta-blocker therapy, with ongoing periodic reassessment for the development of arrhythmias or ventricular dysfunction ([S7.3-46,S7.3-47](#)). Atenolol was shown to reduce VA in 1 study ([S7.3-15](#)). Ambulatory monitoring and/or

exercise testing can be performed to assess adequacy of beta-blocking dosing.

9. Interrogation of ICDs shows that >90% of spontaneous sustained VTs in arrhythmogenic right ventricular cardiomyopathy are monomorphic ([S7.3-12](#)), while sustained monomorphic VT is inducible at electrophysiological study in 55% of patients ([S7.3-36](#)). VT is usually related to scar-related reentry, and the subepicardium usually has more extensive scar than the endocardium ([S7.3-27](#)). In experienced centers, use of epicardial mapping and ablation is associated with better outcomes ([S7.3-27,S7.3-28,S7.3-30,S7.3-31,S7.3-33](#)). Important complications including pericardial tamponade, MI, and death occur in 2.3% to 3.3% of ablation cases ([S7.3-27-S7.3-29](#)), emphasizing the need for performance in centers with specialized expertise in epicardial procedures. Ablation reduces the frequency of recurrent VT, although 27% to 55% of patients ([S7.3-27,S7.3-28](#)) have at least 1 recurrence; ablation of VT in arrhythmogenic right ventricular cardiomyopathy patients does not eliminate the need for an ICD in appropriate candidates. The potential risk of VT recurrence due to disease progression should be reviewed with patients when considering ablation. There are no randomized comparisons of antiarrhythmic therapy to suppress recurrent VT. Beta blockers, sotalol and amiodarone have been used ([S7.3-15](#)). In an observational series, sotalol suppressed inducible VT in 58% of patients with <10% of patients experiencing arrhythmia recurrence during follow-up ([S7.3-48](#)). Effectiveness of the different medications appears to be variable, and so more studies are needed.
10. In arrhythmogenic right ventricular cardiomyopathy, areas of fibrofatty scar in the RV free wall create areas of delayed ventricular activation causing fractionated deflections following the QRS, known as epsilon waves on the surface ECG (a major criterion) and late potentials in the signal averaged ECG (minor criterion) in the 2010 Task Force Criteria for diagnosis of arrhythmogenic right ventricular cardiomyopathy ([S7.3-6](#)). When the standard ECG QRS duration is  $\leq 110$  ms, criteria for abnormal signal-averaged ECG include any 1 of the following: filtered QRS duration  $\geq 114$  ms, duration of the terminal QRS  $< 40 \mu\text{V}$  exceeding 37 ms, or a root mean square voltage in the terminal 40 ms of  $\leq 20 \mu\text{V}$  ([S7.3-6](#)). Abnormal findings on signal averaged ECG correlated with disease severity on cardiac MRI ([S7.3-35](#)), and increased adverse events in males ([S7.3-34](#)). In an assessment of the diagnostic use of testing for arrhythmogenic right ventricular cardiomyopathy, signal averaged ECG was of greater value than cardiac MRI or biopsy ([S7.3-14](#)).

11. The value of an electrophysiological study is uncertain in asymptomatic arrhythmogenic right ventricular cardiomyopathy patients with preserved ventricular function in predicting subsequent risk for SCD. Studies of programmed ventricular stimulation in patients with definite or probable arrhythmogenic right ventricular cardiomyopathy include most symptomatic patients, making recommendations on asymptomatic patients difficult. Electrophysiological studies induce sustained VT in approximately 60% of patients (S7.3-10,S7.3-36); many of whom have had prior spontaneous episodes of sustained VT. In

patients with primary prevention ICDs, inducible sustained VT did not predict subsequent appropriate ICD shocks (S7.3-13). In 1 study including symptomatic patients, patients without inducible VT were less likely to receive appropriate ICD shocks (S7.3-9). In asymptomatic patients without evidence of VA on ambulatory monitoring, a negative electrophysiological study may have limited value in decision-making for an ICD.

#### 7.4. Hypertrophic Cardiomyopathy

##### Recommendations for HCM

References that support the recommendations are summarized in [Online Data Supplement 31](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with HCM, SCD risk stratification should be performed at the time of initial evaluation and periodically thereafter (S7.4-1–S7.4-8).
I	B-NR	2. In patients with HCM who have survived an SCA due to VT or VF, or have spontaneous sustained VT causing syncope or hemodynamic compromise, an ICD is recommended if meaningful survival greater than 1 year is expected (S7.4-1,S7.4-6,S7.4-9,S7.4-10).
I	B-NR	3. In first-degree relatives of patients with HCM, an ECG and echocardiogram should be performed (S7.4-11–S7.4-17).
I	B-NR	4. In first-degree relatives of patients with HCM due to a known causative mutation, genetic counseling and mutation-specific genetic testing are recommended (S7.4-13–S7.4-15,S7.4-18,S7.4-19).
IIa	B-NR	5. In patients with clinically suspected or diagnosed HCM, genetic counseling and genetic testing are reasonable (S7.4-13–S7.4-15,S7.4-18–S7.4-22).
IIa	B-NR C-LD C-LD	6. In patients with HCM and 1 or more of the following risk factors, an ICD is reasonable if meaningful survival of greater than 1 year is expected: a. Maximum LV wall thickness $\geq$ 30 mm (LOE: B-NR) (S7.4-2,S7.4-3,S7.4-23,S7.4-24). b. SCD in 1 or more first-degree relatives presumably caused by HCM (LOE: C-LD) (S7.4-25,S7.4-26). c. 1 or more episodes of unexplained syncope within the preceding 6 months (LOE: C-LD) (S7.4-8,S7.4-26).
IIa	B-NR C-LD	7. In patients with HCM who have spontaneous NSVT (LOE: C-LD) (S7.4-2,S7.4-26,S7.4-27) or an abnormal blood pressure response with exercise (LOE: B-NR) (S7.4-5,S7.4-28,S7.4-29), who also have additional SCD risk modifiers or high risk features, an ICD is reasonable if meaningful survival greater than 1 year is expected.
IIb	B-NR B-NR	8. In patients with HCM who have NSVT (LOE: B-NR) (S7.4-2,S7.4-26,S7.4-27) or an abnormal blood pressure response with exercise (LOE: B-NR) (S7.4-5,S7.4-28,S7.4-29) but do not have any other SCD risk modifiers, an ICD may be considered, but its benefit is uncertain.
IIb	C-LD	9. In patients with HCM and a history of sustained VT or VF, amiodarone may be considered when an ICD is not feasible or not preferred by the patient (S7.4-30,S7.4-31).
III: No Benefit	B-NR	10. In patients with HCM, an invasive electrophysiological study with programmed ventricular stimulation should not be performed for risk stratification (S7.4-32,S7.4-33).
III: No Benefit	B-NR	11. In patients with an identified HCM genotype in the absence of SCD risk factors, an ICD should not be implanted (S7.4-7,S7.4-34,S7.4-35).

**Table 8** and **Figure 7****TABLE 8 Major Clinical Features Associated With Increased Risk of SCD in Patients With HCM****Established risk factors\***

- Survival from a cardiac arrest due to VT or VF ([S7.4-1,S7.4-5,S7.4-6](#))
- Spontaneous sustained VT causing syncope or hemodynamic compromise ([S7.4-1,S7.4-5,S7.4-6](#))
- Family history of SCD associated with HCM ([S7.4-25,S7.4-26](#))
- LV wall thickness  $\geq 30$  mm ([S7.4-2,S7.4-3,S7.4-23,S7.4-24](#))
- Unexplained syncope within 6 mo ([S7.4-8,S7.4-26](#))
- NSVT  $\geq 3$  beats ([S7.4-2,S7.4-26,S7.4-27](#))
- Abnormal blood pressure response during exercise† ([S7.4-5,S7.4-28,S7.4-29](#))

**Potential risk modifiers‡**

- $<30$  y ([S7.4-5,S7.4-26](#))
- Delayed hyperenhancement on cardiac MRI ([S7.4-37–S7.4-39,S7.4-54](#))
- LVOT obstruction ([S7.4-2,S7.4-4](#))
- Syncope  $>5$  y ago ([S7.4-8,S7.4-26](#))

**High-risk subsets§**

- LV aneurysm ([S7.4-40,S7.4-55,S7.4-56](#))
- LVEF  $<50\%$  ([S7.4-52](#))

\*There is general agreement in the literature that these factors independently convey an increased risk for SCD in patients with HCM. †Decrease in blood pressure of 20 mm Hg or failure to increase systolic blood pressure  $>20$  mm Hg during exertion. ‡There is a lack of agreement in the literature that these modifiers independently convey an increased risk of SCD in patients with HCM; however, a risk modifier when combined with a risk factor often identifies a patient with HCM at increased risk for SCD beyond the risk conveyed by the risk factor alone. §A small subset of patients with an LVEF  $<50\%$  (end-stage disease) or an LV aneurysm warrant consideration for ICD implantation ([S7.4-52](#)).

HCM indicates hypertrophic cardiomyopathy; ICD, implantable cardioverter-defibrillator; LV, left ventricular; LVEF, left ventricular ejection fraction; LVOT, left ventricular outflow tract; NSVT, nonsustained ventricular tachycardia; SCD, sudden cardiac death; VT, ventricular tachycardia; and VF, ventricular fibrillation.

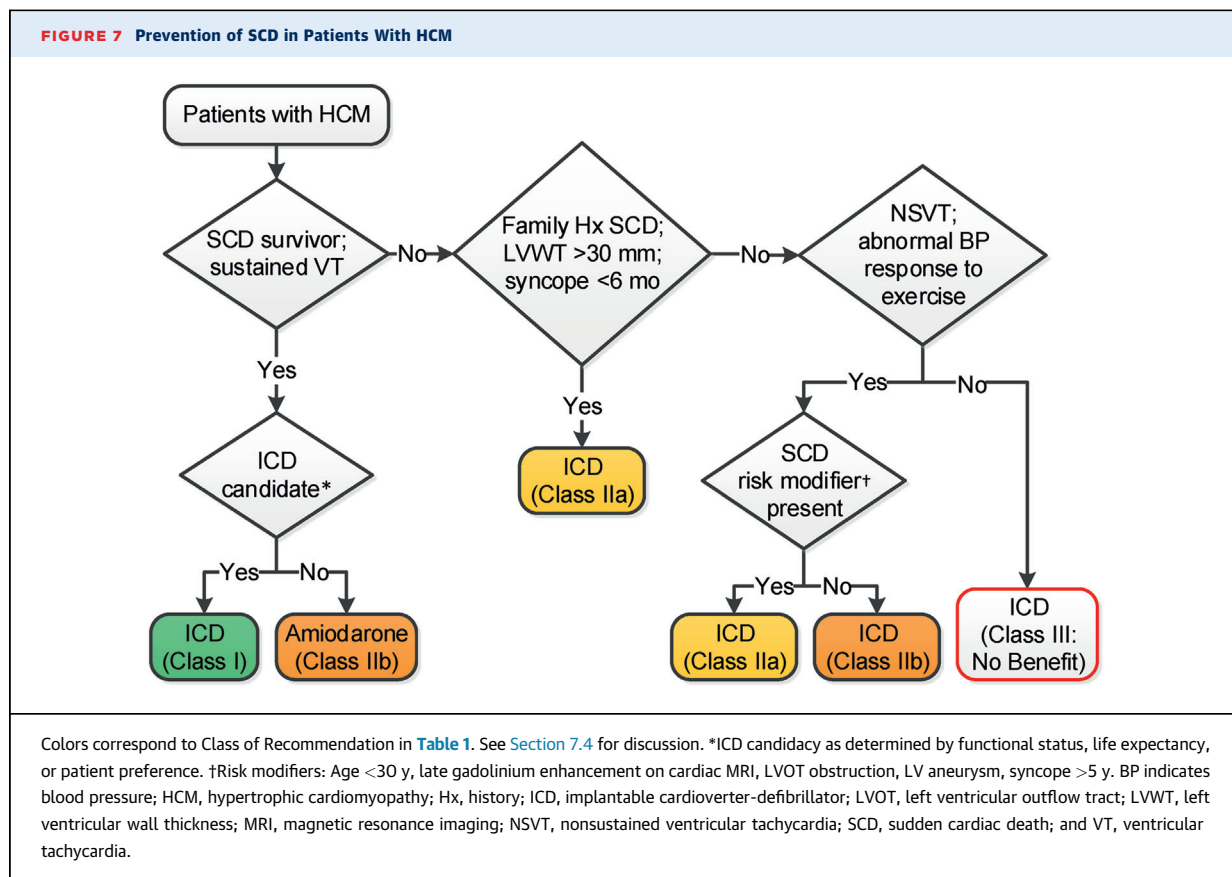
Refer to the ACCF/AHA HCM guideline for the definition of HCM ([S7.4-36](#)).

**Recommendation-Specific Supportive Text**

1. Patients with HCM have approximately a 1% risk of SCD per year ([S7.4-1,S7.4-6](#)). Selection of patients who are appropriate candidates for implantation of an ICD can be a difficult clinical decision because of the individuality of each patient and family, variable definitions of risk factors and risk modifiers, sparse clinical data, the relative infrequency of both HCM and SCD in most clinical practices, and the potential complications of living with an ICD. **Table 8** lists risk factors and risk modifiers associated with SCD in patients with HCM. ICD risk stratification should be performed every 1 to 3 years in patients with HCM. There is increasing evidence supporting the association of late gadolinium enhancement on cardiac MRI with the risk of sudden death and it is included as a risk modifier ([S7.4-37–S7.4-39](#)). LV aneurysm may be associated with a risk of sustained monomorphic VT ([S7.4-40](#)). Age is also an important consideration, as sudden death risk is greater in those  $<30$  years of age, and low in patients whose initial presentation is after the age of 60 years ([S7.4-5,S7.4-26](#)), ([S7.4-41](#)).
2. HCM is the most common cause of SCD in individuals  $<40$  years of age ([S7.4-26](#)). Individuals who have survived an episode of SCD, VF, or sustained VT resulting in syncope or hemodynamic compromise warrant ICD implantation ([S7.4-1,S7.4-6,S7.4-9,S7.4-10](#)). Although there are no RCTs assessing the use of the ICD

in patients with HCM who have survived SCD, 1 study reported that 54% of patients with an ICD placed for secondary prevention received appropriate ICD therapy during an average follow-up of 4.6 years ([S7.4-10](#)). Select patients with HCM may be candidates for implantation of the subcutaneous implantable cardioverter-defibrillator ([S7.4-42](#)); however, more data on this group are needed especially given their higher risk of T wave oversensing that may increase the risk of inappropriate ICD shocks.

3. Clinical and/or genetic screening of first- and second-degree family members of patients with HCM is important to identify those with unrecognized disease. Genetic counseling should precede genetic testing of family members to enhance their understanding of the usefulness and cost of testing ([S7.4-18,S7.4-20,S7.4-43](#)). On the basis of family history, clinical screening, and pedigree analyses, the pattern of inheritance is ascertained to identify and manage relatives at risk ([S7.4-13,S7.4-14,S7.4-18,S7.4-19,S7.4-43–S7.4-45](#)). Because familial HCM is a dominant disorder, the risk that an affected patient will transmit disease to each offspring is 50%. When a pathogenic mutation is identified in an index patient, the genetic status of each family member can be readily ascertained. Relatives with overt HCM will have the same pathogenic HCM mutation as the index patient. Pathogenic mutations may also be identified in other relatives with unknown clinical status. These mutation-positive individuals should be evaluated by physical examination, electrocardiography ([S7.4-11,S7.4-17](#)), and echocardiography



(S7.4-12,S7.4-16,S7.4-17) and, if HCM is identified, these individuals should undergo risk stratification. Gene-positive subjects without evidence of HCM may be at risk for future development of HCM and benefit from ongoing clinical evaluation (S7.4-15,S7.4-46,S7.4-47). If the proband's implicated mutation is the bona fide disease-causing mutation, then mutation-negative family members and their descendants are not at an increased risk for developing HCM and do not need further evaluation. However, such mutation-negative family members must have an echocardiogram to ensure genotype and phenotype concordance.

4. In a study of 1,053 unrelated patients with clinically manifest HCM, 359 patients (34%) were genotype positive for an HCM-associated mutation in  $\geq 1$  HCM-associated genes (S7.4-22). Whether the results of genetic testing in the proband improve outcomes is uncertain, but identification of a mutation can help inform screening of relatives.
5. Genetic counseling is important in patients with HCM, and genetic screening of relatives is also important unless there are no living first- or second-degree relatives. Most HCM is caused by an autosomal dominant mutation in genes that encode sarcomere proteins or sarcomere-associated proteins. Presence of a

pathogenic sarcomere protein gene mutation in patients with HCM identifies risk of LV dysfunction and adverse outcome irrespective of the myofilament involved (S7.4-13–S7.4-15,S7.4-18,S7.4-19,S7.4-22). A single mutation in 1 of the 2 alleles (or copies) of a gene is sufficient to cause HCM; however, 5% of patients with HCM have  $\geq 2$  mutations in the same gene or different genes, which can be a marker for worse outcomes (S7.4-13,S7.4-34,S7.4-48). When genetic testing reveals a mutation in the index patient, ascertainment of genetic status in first- and second-degree relatives can be predictive of risk for developing HCM (S7.4-14, S7.4-49). Relatives with overt HCM will have the same pathogenic HCM mutation as the index patient. 6. Several studies have described an independent relationship between hypertrophy and SCD when the magnitude of hypertrophy is  $\geq 30$  mm (S7.4-2,S7.4-3, S7.4-23,S7.4-24). Risk does not abruptly increase for patients with a  $\geq 30$  mm wall thickness, but it rather increases in a linear manner (S7.4-24) and appears to carry more prognostic significance in younger patients. A young adult with hypertrophy that approaches 30 mm may have similar or greater SCD risk than an older patient with maximum wall thickness  $\geq 30$  mm (S7.4-23,S7.4-50).

Patients with HCM are at an increased risk for SCD if they have a first-degree relative who experienced SCD presumably caused by HCM. Family history appears to be an independent predictor of SCD although the supportive studies are small and observational (S7.4-25, S7.4-26). Syncope can be neurally mediated or medication-related as well as due to VA and requires a careful evaluation before considering it a risk factor for SCD (S7.4-8,S7.4-26). In an analysis, syncope that was unexplained or thought not to be neurally mediated was associated with SCD risk only when it occurred within the past 6 months but not if the most episode occurred >5 years previously (S7.4-8).

7. Although sustained VT is clearly associated with SCD, the data for NSVT are less robust. Most studies do not support NSVT as an independent risk factor for SCD in patients with HCM (S7.4-2,S7.4-26,S7.4-27), but the risk increases if risk modifiers are present, especially in patients <30 years of age (S7.4-27). Up to one third of patients with HCM have an abnormal blood pressure response during exercise testing (defined variably as either a 20 mm Hg decrease in blood pressure or a failure to increase systolic blood pressure by at least 20 mm Hg during effort) (S7.4-28,S7.4-29). This finding has been postulated to be a risk factor for SCD; however, it is unclear how this relates to the increase in dynamic LV outflow tract obstruction that occurs with exertion, a hemodynamic condition that is readily modifiable with medication or mechanical procedures. The significance of an abnormal blood pressure response with exercise predicting SCD risk increases in the presence of risk modifiers (Table 8).
8. Most studies have found that NSVT alone has a low positive predictive value for SCD (S7.4-2,S7.4-26,S7.4-27); therefore, use of an ICD is more appropriate if risk modifiers are also present. An abnormal blood pressure response to exercise has also been associated with the risk of sudden death (S7.4-5,S7.4-28,S7.4-29), but it is unclear how this relates to the increase in dynamic LV outflow tract obstruction that occurs with effort, which is often treatable. The significance of an abnormal blood pressure response with exercise for

predicting SCD risk increases when risk modifiers are present (Table 8).

9. The ICD is recommended for the prevention of SCD in patients with HCM who have survived sustained VT or VF as antiarrhythmic medications have limited effectiveness (S7.4-31). Amiodarone has been associated with improved survival in observational studies and is an option for patients for whom an ICD is not feasible due to limited expectation for survival or patient preference (S7.4-30,S7.4-31).
10. Approximately one third of consecutive patients with HCM undergoing an electrophysiological study have polymorphic VT or VF induced by programmed ventricular stimulation, but the results of programmed stimulation do not predict SCD risk. Programmed ventricular stimulation in patients with HCM has low predictive value and a nontrivial risk of complications (S7.4-32,S7.4-33,S7.4-51). Electrophysiological studies can help to clarify the diagnosis of wide complex tachycardia or guide therapy for supraventricular tachycardia or bundle branch reentry.
11. SCD may cluster in certain families with HCM, and the possibility that specific sarcomere mutations may confer SCD risk has been hypothesized. However, subsequent studies of selected patients with HCM (S7.4-34,S7.4-35) were unable to establish a clinically useful relation between genotype and SCD risk. In some cases, the rate of adverse events (and prevalence of associated SCD risk factors) was lower in patients with mutations initially felt to be malignant than it was in those with mutations believed to be benign (S7.4-34, S7.4-35). Data from series of unselected consecutive outpatients suggest that most mutations are novel and limited to particular families (S7.4-34,S7.4-35). Therefore, routine mutation screening would appear to be of little prognostic value in HCM (S7.4-52). The short-term risk of sudden death in patients who are genotype positive but have no other manifestations of the disease appears to be low (S7.4-53). Therefore, an ICD is not indicated in these individuals.

## 7.5. Myocarditis

### Recommendations for Myocarditis

References that support the recommendations are summarized in Online Data Supplement 32.

COR	LOE	RECOMMENDATIONS
I	C-LD	<ol style="list-style-type: none"> <li>1. In patients with life-threatening VT or VF associated with confirmed or clinically suspected myocarditis, referral to centers with mechanical hemodynamic support and advanced arrhythmia management is recommended (S7.5-1).</li> </ol>
IIb	C-LD	<ol style="list-style-type: none"> <li>2. In patients with giant cell myocarditis with VF or hemodynamically unstable VT treated according to GDMT, an ICD and/or an antiarrhythmic medication may be considered if meaningful survival of greater than 1 year is expected (S7.5-2–S7.5-4).</li> </ol>

### Recommendation-Specific Supportive Text

1. Myocarditis is an inflammatory process often related to infection ([S7.5-1,S7.5-5–S7.5-9](#)). When patients are treated in centers with the availability of mechanical hemodynamic support procedures, cardiac catheterization, endomyocardial biopsy, advanced cardiac imaging procedures, and arrhythmia management including ICD implantation, outcomes appear improved ([S7.5-1](#)). The acute course of myocarditis varies ranging from an asymptomatic finding of transient ST-T changes noted on ECG to cardiogenic shock and recurrent VA ([S7.5-10–S7.5-12](#)). Acute management is largely supportive and can rapidly advance to requiring mechanical support ([S7.5-13,S7.5-14](#)). Cardiac arrhythmias range from conduction abnormalities to life-threatening VT and VF ([S7.5-15–S7.5-17](#)). Arrhythmias may require antiarrhythmic medications and/or device therapy ([S7.5-18](#)). Giant cell myocarditis is fairly uncommon, but it is of particular importance because it typically affects young individuals and is usually fatal if untreated ([S7.5-2–S7.5-4,S7.5-19](#)). VT may require antiarrhythmic medications such as amiodarone and/or an ICD that in some instances can be used as a bridge to more advanced HF therapies such as LVAD or trans-

plant. Myocarditis and SCD have been reported with HIV infection ([S7.5-20,S7.5-21](#)). Systemic lupus erythematosus can cause myocarditis but only rarely VT or VF ([S7.5-8,S7.5-22](#)). In patients with Chagas disease, acute myocarditis is rare but more than one third of affected patients develop late myocardial damage with progressive HF. Conduction defects with progression to complete heart block and VT or VF are common. Amiodarone appears to be effective in treating VA ([S7.5-23](#)). An ICD is frequently used in the late phase of myocarditis ([S7.5-24](#)), and radiofrequency catheter ablation has been successfully used to control recurrent VA in some patients ([S7.5-25](#)).

2. Giant cell myocarditis is fairly uncommon, but it is of particular importance as it typically affects young individuals and is usually fatal if untreated. The diagnosis is confirmed by endomyocardial biopsy. Patients may develop heart block, requiring a temporary or a permanent pacemakers. An ICD and antiarrhythmic medications, such as amiodarone are often used in the acute phase to treat VT or VF and reduce the risk of SCD ([S7.5-2–S7.5-4,S7.5-19,S7.5-26–S7.5-28](#)).

### 7.6. Cardiac Sarcoidosis

#### Recommendations for Cardiac Sarcoidosis

References that support the recommendations are summarized in [Online Data Supplement 33](#).

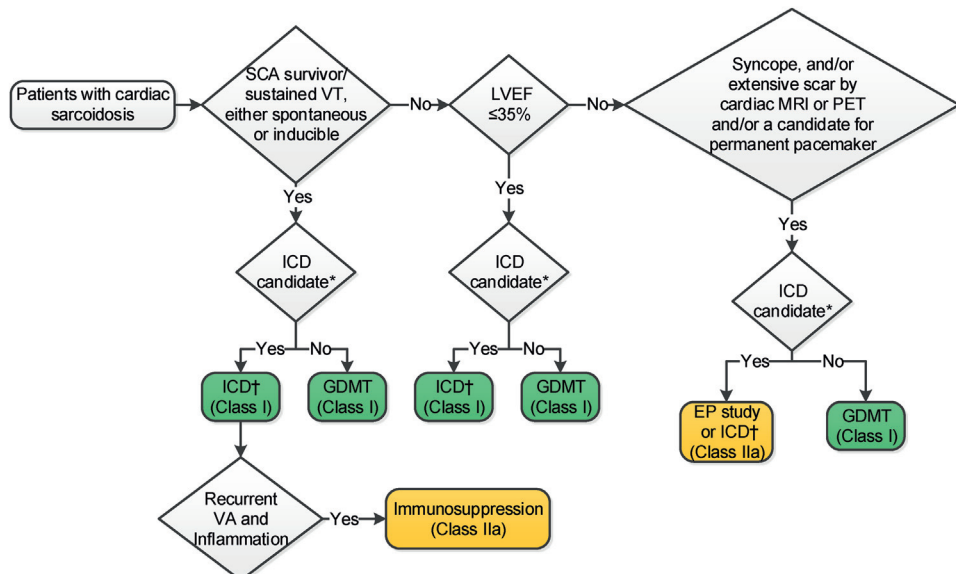
COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with cardiac sarcoidosis who have sustained VT or are survivors of SCA or have an LVEF of 35% or less, an ICD is recommended, if meaningful survival of greater than 1 year is expected ( <a href="#">S7.6-1–S7.6-5</a> ).
IIa	B-NR	2. In patients with cardiac sarcoidosis and LVEF greater than 35% who have syncope and/or evidence of myocardial scar by cardiac MRI or positron emission tomographic (PET) scan, and/or have an indication for permanent pacing, implantation of an ICD is reasonable, provided that meaningful survival of greater than 1 year is expected ( <a href="#">S7.6-6–S7.6-10</a> ).
IIa	C-LD	3. In patients with cardiac sarcoidosis and LVEF greater than 35%, it is reasonable to perform an electrophysiological study and to implant an ICD, if sustained VA is inducible, provided that meaningful survival of greater than 1 year is expected ( <a href="#">S7.6-11,S7.6-12</a> ).
IIa	C-LD	4. In patients with cardiac sarcoidosis who have an indication for permanent pacing, implantation of an ICD can be beneficial ( <a href="#">S7.6-13</a> ).
IIa	C-LD	5. In patients with cardiac sarcoidosis with frequent symptomatic VA and evidence of myocardial inflammation, immunosuppression in combination with antiarrhythmic medication therapy can be useful to reduce VA burden ( <a href="#">S7.6-14–S7.6-16</a> ).

**Figure 8**

### Recommendation-Specific Supportive Text

1. Sarcoidosis is a systemic granulomatous disease of unknown cause. Pulmonary involvement is most frequent but any organ can be affected. Cardiac

involvement, diagnosed by cardiac MRI or positron emission tomography (PET), has been reported in up to 55% of patients with extracardiac disease, while isolated cardiac sarcoidosis was seen in most patients diagnosed with cardiac sarcoidosis in 1 report ([S7.6-17](#)).

**FIGURE 8** Prevention of SCD in Patients With Cardiac Sarcoidosis

Colors correspond to Class of Recommendation in **Table 1**. See **Section 7.6** for discussion. \*ICD candidacy as determined by functional status, life expectancy, or patient preference. †For recurrent sustained monomorphic VT, refer to **Figure 2**. CEP indicates electrophysiological; GDMT, guideline-directed management and therapy; ICD, implantable cardiac-defibrillator; LVEF, left ventricular ejection fraction; MRI, magnetic resonance imaging; PET, positron emission tomography; SCA, sudden cardiac arrest; SCD, sudden cardiac death; VA, ventricular arrhythmia; and VT, ventricular tachycardia.

Cardiac manifestations include conduction abnormalities, VA, and depressed ventricular function with or without HF, and these contribute greatly to a higher mortality in cardiac sarcoidosis compared with sarcoidosis without cardiac involvement (**S7.6-2**). In a 25-year study of 110 patients with cardiac sarcoidosis in Finland with HF at presentation, marked LV dysfunction at diagnosis (LVEF <35%), and isolated cardiac sarcoidosis predicted an adverse outcome (**S7.6-1**). VA can also occur in patients with relatively normal LV function, some of whom have RV involvement that can mimic arrhythmogenic right ventricular cardiomyopathy. Several reports of patients with cardiac sarcoidosis and ICDs implanted for either primary or secondary prevention of SCD show a high frequency of appropriate ICD therapies (**S7.6-3–S7.6-5**), supporting use of ICDs for primary and secondary prevention of SCD according to the indications applied for other cardiomyopathies. The frequency of conduction abnormalities often warrants a device that provides bradycardia pacing as well.

2. Patients with cardiac sarcoidosis can experience VA and SCD, even if the LVEF is normal, and approaches to identification of patients at risk of SCD despite preserved LV function are not well defined. A number of

studies have evaluated the role of cardiac MRI for predicting VA and SCD. A meta-analysis (**S7.6-6**), which included 760 patients in 10 studies, found that late gadolinium enhancement was associated with increased all-cause mortality and more VA compared with those without late gadolinium enhancement. Applicability is limited by the lack of precise quantification of late gadolinium enhancement burden that may allow for more nuanced risk stratification. Some studies suggested that a threshold effect exists, with extensive LV and RV involvement being a particularly high-risk feature (**S7.6-7,S7.6-8**). However, late gadolinium enhancement can be present even if the LVEF is >50% and was associated with a risk of death or VT of 4.9% per year compared to 0.24% per year when late gadolinium enhancement was absent in 1 observational study (**S7.6-7**). PET for assessing inflammation and scar is also being increasingly used, but data are limited. In 1 report, the presence of inflammation and RV involvement on PET scanning was associated with increased risk of death or e.g. of relatives (**S7.6-10**). Electrophysiological studies in a series of 76 patients with evidence of cardiac sarcoid found that 11% had inducible VT. During a median follow-up of 5 years, 75% of patients with inducible VT had spontaneous VT

- or death compared with 1.5% of those who did not have inducible VT ([S7.6-18](#)).
3. Electrophysiological study has been proposed as a potential tool for risk stratification of VA and SCD in patients who had demonstrable evidence of cardiac sarcoidosis based on imaging studies or biopsy, but do not have documented arrhythmias or arrhythmic symptoms nor meet standard primary prevention criteria for ICD implantation.
- One study evaluated 76 patients with documented cardiac sarcoidosis by PET or cardiac MRI who underwent electrophysiological study ([S7.6-12](#)). Eight (11%) were inducible for sustained VAs and received an ICD, while the rest did not receive an ICD because they were not inducible. LVEF was lower in patients with inducible VA (36.4 + 4.2% versus 55.8 + 1.5%). Over a median follow-up of 5 years, 6 of 8 patients in the group with inducible VA had VA or died, compared with 1 death in the negative group ([S7.6-12](#)). An important caveat is that it remains unclear if electrophysiological study is more predictive than LVEF alone, because inducibility appears to reversely correlate with LVEF. Furthermore, in this study the average LVEF of the inducible patients declined further during the followup period ([S7.6-12](#)).
4. In addition to VA and LV dysfunction, conduction abnormalities, including heart block, can also be a common manifestation of cardiac sarcoidosis. Patients with documented VA and LV dysfunction are at increased risk of cardiac events including cardiac death. One study compared outcomes in 22 patients with high-degree atrioventricular block as the initial manifestation of cardiac sarcoidosis, to 31 patients who initially presented with VT and/or HF. After a median follow up of 34 months, the patients who presented with heart block had fewer HF hospitalization, yet fatal cardiac events, including sustained VAs, were similar to those with VT and/or HF, suggesting that the risk of fatal cardiac events is high regardless of the initial clinical presentation ([S7.6-13](#)). In the same study, administration of steroids led to some clinical improvement, with some patients recovering conduction, yet steroid effectiveness was not universal and did not seem to be protective against adverse cardiac events ([S7.6-13](#)).
5. Several studies have attempted to evaluate the role of immunosuppression for reducing VA in patients with cardiac sarcoidosis, but results have been inconsistent ([S7.6-14–S7.6-16](#)). Furthermore, a worsening of VA has been reported with immunosuppressive therapy (usually glucocorticoids) in a number of patients, including electrical storm developing in some within 12 months of initiating therapy ([S7.6-15](#)). One study reported a

decrease of arrhythmia burden with steroid therapy but only when given in the early stages of the disease; those with advanced LV dysfunction did not experience benefit ([S7.6-16](#)). A systematic combined treatment approach was successful in 63% of patient in a series in which medical therapy included both steroids and antiarrhythmic medications, followed by radiofrequency catheter ablation if needed ([S7.6-14](#)). Immunosuppressive therapy may serve a dual purpose beyond arrhythmia effects as it may help stabilize disease progression and prevent further deterioration of LV function, although this has yet to be demonstrated in RCTs. Steroids do not appear to reverse advanced ventricular dysfunction once present, which supports the importance of early diagnosis and intervention ([S7.6-1](#)). PET scanning for assessing inflammation and scar is being increasingly used in sarcoidosis as well, but data supporting its use for guiding therapy of arrhythmias are limited.

#### 7.6.1. Other Infiltrative Cardiomyopathies

Infiltrative cardiomyopathies are a heterogeneous group of uncommon systemic diseases with associated cardiac involvement. In some infiltrative cardiomyopathies, such as Fabry's disease, VAs are uncommon. Some, such as hemochromatosis, are highly treatable especially when diagnosed early. In all cases, treatment of the underlying condition must accompany management of cardiac arrhythmias. Most studies of infiltrative cardiomyopathies and arrhythmias are small and observational ([S7.6.1-1](#)) but, in general, unless contraindications are present, VAs should be treated as in any other cardiomyopathy. See [Section 7.6](#) for sarcoidosis. Until recently, cardiac amyloidosis was associated with a very poor prognosis with patients ultimately succumbing to progressive HF ([S7.6.1-2](#)). This perception is changing with advances in medical therapy for light-chain amyloidosis, which have led to improved outcomes ([S7.6.1-3](#)). Yet, decisions must be individualized because data remain too limited to allow formal recommendations as published reports on ICD effectiveness in amyloidosis are small, observational and with limited follow up ([S7.6.1-4](#)). Whether there is greater benefit to ICD placement in light chain amyloidosis versus transthyreitin-related amyloidosis remains uncertain, because most studies included mainly patients with amyloid light-chain amyloidosis for which the rate of VA may be greater and prognosis is generally worse. Whether ICDs are effective for primary prevention of SCD is uncertain, but many deaths in patients with cardiac amyloidosis do not appear to be preventable by an ICD ([S7.6.1-2](#)).

## 7.7. Heart Failure

### 7.7.1. HF With Reduced Ejection Fraction

#### Recommendation for HFrEF

References that support the recommendation are summarized in [Online Data Supplement 35](#).

COR	LOE	RECOMMENDATION
IIa	B-NR	1. In patients with HFrEF who are awaiting heart transplant and who otherwise would not qualify for an ICD (e.g., NYHA class IV and/or use of inotropes) with a plan to discharge home, an ICD is reasonable ( <a href="#">S7.7.1-1–S7.7.1-5</a> ).

#### Synopsis

Patients with HFrEF are at an increased risk for VA and SCD. The risk is increased irrespective of HFrEF etiology ([S7.7.1-6](#)). SCD makes up a greater proportion of deaths in patients with milder HF symptoms and lesser proportion in those with moderate/severe HF symptoms ([S7.7.1-7](#)). The reported incidence of SCD varies depending on the definition used and the population studied. Although many deaths, classified as sudden, are indeed due to lethal VA, others may be due to bradyarrhythmias, pulseless electrical activity, and sudden hemodynamic deterioration ([S7.7.1-7–S7.7.1-9](#)).

Medical therapy with neurohormonal agents decreases the risk of SCD by reducing both the incidence of VA and disease progression ([S7.7.1-7,S7.7.1-10–S7.7.1-12](#)). Despite GDMT for HFrEF, some patients remain at risk for SCD, and an ICD may be helpful. See [Sections 7.1 and 7.2](#) for the indications on ICDs in patients with reduced LVEF. CRT, in appropriate patients, has also been shown to reduce the incidence of SCD ([S7.7.1-13](#)).

The pathophysiology of SCD in HF is complex, resulting from interactions between both functional and structural changes that occur in patients with HFrEF that result in increased susceptibility to SCD ([S7.7.1-14](#)). Although many of the risk factors are shared among HFrEF patients, the reason that SCD strikes a particular individual is usually unknown; however, some individuals may have a genetic susceptibility ([S7.7.1-15](#)). Varying degrees of myocardial fibrosis, neurohormonal activation, and increased wall stress alter the electrophysiological properties with changes in cell coupling, ionic currents (electrical remodeling), and calcium handling that likely contribute to the development of lethal VA ([S7.7.1-16](#)). Contributing factors extrinsic to the heart include electrolyte abnormalities related to volume shifts and diuretic use, sympathetic activation, hemodynamic stress, and hypoxia.

#### Recommendation-Specific Supportive Text

- Many patients with advanced HF listed for heart transplant would not otherwise qualify for ICD given the severity of illness including NYHA class IV status and/or use of inotropic infusion. Although no randomized

data on ICD use in this population exist, data from observational and large registry studies of patients awaiting heart transplant suggest improved survival in patients with an ICD ([S7.7.1-1,S7.7.1-4,S7.7.1-5](#)). One alternative to ICD in this population is the wearable cardioverter-defibrillator ([S7.7.1-2,S7.7.1-3](#)). The recommendation in this section is relevant to those patients without an ICD where there is a plan to discharge the patient to home to await cardiac transplant and not, for example, to those patients who remain hospitalized with no intention to discharge home until transplant occurs. For those patients with an LVAD, the decision to place an ICD is generally independent of whether they are awaiting heart transplant but rather the indication in those patients is generally based on the need to treat VA ([S7.7.1-17](#)).

#### 7.7.2. HF With Preserved Ejection Fraction

Nearly half of the patients with HF have a preserved LVEF ([S7.7.2-1](#)). These patients tend to be older and have more comorbidities than patients with HFrEF. However, although the rate of SCD is lower in patients with HF with preserved ejection fraction (HFpEF) than in patients with HFrEF ([S7.7.2-2](#)), nearly a quarter of all deaths among patients with HFpEF are sudden ([S7.7.2-3–S7.7.2-5](#)). The challenge in preventing SCD in patients with HFpEF is identifying which patients are at a high enough risk to benefit from preventive therapies. Studies exploring noninvasive risk factors for SCD in patients with HFpEF do not identify consistent factors with the exception of ischemic heart disease ([S7.7.2-2,S7.7.2-6](#)). Consequently, there is no accepted noninvasive test to identify high-risk patients with HFpEF. Invasive risk stratification with an electrophysiological study shows promise in this population ([S7.7.2-7,S7.7.2-8](#)). This topic is currently being studied in the PRESERVE-EF (Risk Stratification in Patients With Preserved Ejection Fraction) trial (NCT02124018).

Whether to include a recommendation related to an electrophysiological study in patients with HFpEF and ischemic heart disease was carefully considered by the writing committee. However, evidence was deemed

insufficient to support a formal recommendation. Still, the pros and cons of an electrophysiological study can reasonably be considered in select patients with HFpEF

and ischemic heart disease who are experiencing symptoms suggestive of a VA.

#### 7.7.3. Left Ventricular Assist Device

##### Recommendation for Patients With an LVAD

References that support the recommendation are summarized in [Online Data Supplement 36](#).

COR	LOE	RECOMMENDATION
IIa	C-LD	1. In patients with an LVAD and sustained VA, an ICD can be beneficial ( <a href="#">S7.7.3-1</a> ).

#### Recommendation-Specific Supportive Text

1. Patients with an LVAD have a high risk of VA, particularly those with a history of arrhythmias ([S7.7.3-2–S7.7.3-4](#)). The increased risk of VA may be due to myocardial irritation from insertion of the LVAD inflow cannula, LV compression due to a suctioning effect from the LVAD, inotropic support frequently needed by some patients, and repolarization changes that can occur after LVAD placement. Although VT/VF is tolerated by some patients with an LVAD, others experience a decrease in flow as the RV is unsupported; syncope and hypoperfusion can result. Having an ICD can allow for prompt termination of VA before significant hemodynamic consequences occur. Data on ICDs in patients with an LVAD are from observational series. A systematic review of 6 observational

studies observed that within 7 months, 26% of patients with an LVAD had died ([S7.7.3-1](#)). The death rate was lower among patients who previously had an ICD (16% versus 32%), suggesting a 39% relative-risk reduction in all-cause mortality in an adjusted analysis ([S7.7.3-1](#)). Patients with a history of pre-LVAD VA have nearly a  $\geq 10$ -fold risk of post-LVAD VA ([S7.7.3-2–S7.7.3-4](#)). In many of the initial studies demonstrating ICD benefit, older pulsatile LVAD devices were in use ([S7.7.3-2,S7.7.3-5](#)). Studies of ICD use with the newer, continuous flow LVADs have inconsistently shown benefit ([S7.7.3-1, S7.7.3-4,S7.7.3-6,S7.7.3-7](#)). Of note, approximately 2 of 10 patients with an LVAD develop an LVAD related infection in the first year ([S7.7.3-8,S7.7.3-9](#)).

#### 7.7.4. ICD Use After Heart Transplantation

##### Recommendation for ICD Use After Heart Transplantation

References that support the recommendation are summarized in [Online Data Supplement 37](#).

COR	LOE	RECOMMENDATION
IIb	B-NR	1. In patients with a heart transplant and severe allograft vasculopathy with LV dysfunction, an ICD may be reasonable if meaningful survival of greater than 1 year is expected ( <a href="#">S7.7.4-1–S7.7.4-3</a> ).

#### Recommendation-Specific Supportive Text

1. Development of disease in the transplanted heart places some patients at an increased risk of SCD that has ranged from 10% to 35% in observational studies ([S7.7.4-4,S7.7.4-5](#)). Both rejection and a decreased LVEF are predictors of SCD. The mechanisms underlying SCD in patients with a heart transplant include damage to the conduction system itself and VA due to coronary vasculopathy or during episodes of acute rejection. Several small case series observing

appropriate ICD termination of VA suggest that an ICD can be beneficial in selected patients, particularly those with severe allograft vasculopathy, unexplained syncope, a history of SCA, and severe LV dysfunction ([S7.7.4-1–S7.7.4-3](#)). Additionally, a patient with severe allograft vasculopathy who is being considered for retransplant may be appropriate for an ICD as a bridging device. Secondary prevention indications for an ICD in patients with a heart transplant are identical to those in other patients.

## 7.8. Neuromuscular Disorders

### Recommendations for Neuromuscular Disorders

References that support the recommendations are summarized in [Online Data Supplement 38](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with neuromuscular disorders, primary and secondary prevention ICDs are recommended for the same indications as for patients with NICM if meaningful survival of greater than 1 year is expected ( <a href="#">S7.8-1,S7.8-2</a> ).
IIa	B-NR	2. In patients with Emery-Dreifuss and limb-girdle type IB muscular dystrophies with progressive cardiac involvement, an ICD is reasonable if meaningful survival of greater than 1 year is expected ( <a href="#">S7.8-3–S7.8-8</a> ).
IIa	B-NR	3. In patients with muscular dystrophy, follow-up for development of cardiac involvement is reasonable, even if the patient is asymptomatic at presentation ( <a href="#">S7.8-9–S7.8-12</a> ).
IIb	B-NR	4. In patients with myotonic dystrophy type 1 with an indication for a permanent pacemaker, an ICD may be considered to minimize the risk of SCA from VT if meaningful survival of greater than 1 year is expected ( <a href="#">S7.8-9,S7.8-13,S7.8-14</a> ).

**Table 9**

### Synopsis

The muscular dystrophies are a group of inherited diseases affecting skeletal and cardiac muscle. Some present primarily as a NICM (e.g., Duchenne, Becker, and limb-girdle types 2C, 2F, and 2I), while others present primarily as conduction system degeneration with a variable association with cardiomyopathy (e.g., myotonic dystrophy types 1 and 2, Emery-Dreifuss, limb-girdle type 1B; summarized in [Table 9](#)) ([S7.8-15](#)). Because SCD can occur either due to VA or due to bradyarrhythmias from rapid and unpredictable progression of conduction system disease, the clinician is faced with the challenge of identifying those patients who would benefit from prophylactic pacemaker or ICD implantation. There should be a high level of concern for those patients with muscular dystrophy who present with arrhythmia symptoms ([S7.8-15](#)). The current guideline focuses on VA and indications for implantation of an ICD. The indications for permanent pacemaker are discussed in another ACC/AHA/HRS guideline ([S7.8-16](#)).

### Recommendation-Specific Supportive Text

- In general, the indications for an ICD in patients with muscular dystrophy should follow standard ICD recommendations for patients with NICM (see [Section 7.2.1](#) on Secondary Prevention and [Section 7.2.2](#) on Primary Prevention of SCD with NICM). A high index of suspicion for bundle-branch reentrant tachycardia is warranted in patients with myotonic dystrophy who exhibit wide QRS complex tachycardia or tachycardia-related symptoms ([S7.8-2](#)).

2. In patients with Emery-Dreifuss and limb-girdle type 1B muscular dystrophies associated with *Lamin A/C* mutations, SCD accounts for about one third of all deaths ([S7.8-4](#)). Observational studies show a significant rate of appropriate ICD therapy in patients with cardiac conduction disorders who are gene positive for *Lamin A/C* mutation even if LV function is preserved ([S7.8-3,S7.8-5,S7.8-17](#)). In an observational study in which 38% had isolated skeletal muscular involvement but included patients with conduction defects and other risk factors (including PR interval >240 ms, left bundle-branch block, NSVT, or bradycardia requiring a permanent pacemaker) life-threatening VAs were relatively common; with 52% of patients receiving appropriate ICD therapy including approximately 40% of those patients with an LVEF  $\geq 45\%$  ([S7.8-3](#)). A study of patients who had *Lamin A/C* mutation, in which approximately 21% had a skeletal muscular dystrophy phenotype, SCD and appropriate ICD therapy were associated with NSVT, LVEF  $<45\%$ , male sex, and *Lamin A/C* nonmissense mutations ([S7.8-4](#)). These observational studies support the use of an ICD when a pacing indication is present and likely also when evidence of progressive cardiac involvement such as cardiac conduction defects, NSVT or reduced LVEF is present ([S7.8-8](#)).

There is a paucity of data regarding the rare form of x-linked recessive Emery-Dreifuss muscular dystrophy (related to the *Emerin* gene mutation), but arrhythmias may be less frequent than for the *Lamin A/C* mutations ([S7.8-15](#)).

**TABLE 9 Neuromuscular Disorders Associated With Heart Disease**

Muscular Dystrophy	Inheritance	Gene/Protein Affected	Primary Cardiac Pathology	Frequency of Cardiac Involvement	Causes of Death	Associated With Sudden Death?
Duchenne	X-linked recessive	Dystrophin	NICM	>90%	Respiratory, HF	Yes, uncertain etiology
Becker	X-linked recessive	Dystrophin	NICM	60%–75%	HF, respiratory	Yes, uncertain etiology
Limb-girdle type 1B	Autosomal dominant	Lamin A/C	Conduction system disease and NICM	>90%	Sudden, HF	Yes
Limb-girdle type 2C-2F	Autosomal recessive	Sarcoglycan	NICM	<25%	Respiratory, HF	Uncertain
Limb-girdle type 2I	Autosomal recessive	Fukutin-related protein	NICM	20%–80%	Respiratory, HF	Uncertain
Myotonic type 1	Autosomal dominant	CTG repeat expansion	Conduction system disease and NICM	60%–80%	Respiratory, sudden, HF	30% of deaths, uncertain bradycardia versus tachycardia
Myotonic type 2	Autosomal dominant	CCTG repeat expansion	Conduction system disease	10%–25%	Normal causes	Reported
Emery-Dreifuss	X-linked and autosomal dominant or recessive	Emerin, Lamin A/C	Conduction system disease and NICM	>90%	Sudden, HF	Yes
Facioscapulohumeral	Autosomal dominant	D4Z4 repeat contraction	Possibly conduction disease	5%–15%	Normal causes, respiratory rarely	Not reported

HF indicates heart failure; and NICM, nonischemic cardiomyopathy.

Adapted with permission from Groh, et al. (S7.8-15).

3. Cardiac involvement can occur in a number of neuromuscular dystrophies (Table 9). To determine cardiac involvement, a 12-lead ECG and echocardiogram are important for the initial clinical assessment, independent of symptom status. In general, the more extensive the cardiac involvement, including evidence of distal conduction disease, ventricular dysfunction, and atrial arrhythmias, the more likely a VA will occur. The initial evaluation for myotonic dystrophy patients includes ambulatory monitoring. In asymptomatic patients, some experts advocate for annual follow-up during the concealed phase of the disease with an annual 12-lead ECG to screen for development of conduction abnormalities. However, the optimal frequency of electrocardiographic screening is unknown (S7.8-18). Once cardiac involvement is present, either on the basis of conduction delay, atrial arrhythmias, or ventricular dysfunction, a low threshold for investigating symptoms or electrocardiographic findings by the clinician to determine the need for pacemaker implantation,

invasive electrophysiological studies, or ICD implantation is optimal.

4. Up to one third of deaths in myotonic dystrophy patients are sudden (S7.8-9). Although commonly attributed to conduction block and asystole, SCD due to VT/VF has been recognized in patients with functioning permanent pacemakers, and spontaneous VA have been documented in some (S7.8-13,S7.8-19). The risk of SCD in patients with pacemakers suggests that an ICD may be preferred to a pacemaker. However, these patients are also at high risk of respiratory failure as a competing cause of death. Therefore, in patients with severe skeletal muscle involvement, a pacemaker or ICD may not improve outcomes (S7.8-15). A shared decision-making approach to selecting ICD or pacing therapy is warranted. Compared with myotonic type 1 patients, myotonic dystrophy type 2 patients are not well studied but may also benefit from the same approach.

## 7.9. Cardiac Channelopathies

### Recommendations for Cardiac Channelopathies

References that support the recommendations are summarized in Online Data Supplement 39.

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In first-degree relatives of patients who have a causative mutation for long QT syndrome, catecholaminergic polymorphic ventricular tachycardia, short QT syndrome, or Brugada syndrome, genetic counseling and mutation-specific genetic testing are recommended (S7.9-1–S7.9-6).
I	B-NR	2. In patients with a cardiac channelopathy and SCA, an ICD is recommended if meaningful survival of greater than 1 year is expected (S7.9-7–S7.9-13).

## Synopsis

Implantation of an ICD in asymptomatic low-risk patients with a cardiac channelopathy for a positive family history of SCD as the sole indication is unsupported by published data ([S7.9-13–S7.9-18](#)).

## Recommendation-Specific Supportive Text

- Clinical screening of first-degree relatives of patients with inherited arrhythmia syndromes is crucial to identifying affected family members. Due to the increased risk of adverse cardiac events in genotype positive patients with long QT syndrome, catecholaminergic polymorphic ventricular tachycardia, and Brugada syndrome, targeted screening for the identified family-specific mutation can identify individuals who are at risk for these adverse outcomes ([S7.9-2–S7.9-5](#)). Screening ECGs may be insufficient for diagnosis, because the resting ECG in patients with catecholaminergic polymorphic ventricular tachycardia is normal, and as many as 25% of genotype-positive patients with long QT syndrome have QTc intervals  $\leq 440$  ms ([S7.9-2](#)). Due to the increased risk of adverse cardiac events in young patients with long QT syndrome and catecholaminergic polymorphic ventricular tachycardia ([S7.9-2,S7.9-19–S7.9-22](#)), screening infants and young children is particularly important to guide therapy and institute preventive measures, including the avoidance of possible provocative medications ([www.crediblemeds.org](http://www.crediblemeds.org)) ([S7.9-23](#)). However, because up to 15% of mutations previously associated with catecholaminergic polymorphic ventricular tachycardia do not appear to cause disease ([S7.9-24](#)), caution is advised to avoid unnecessary treatment or sports restriction in

phenotype-negative catecholaminergic polymorphic ventricular tachycardia mutation positive individuals. Notably, some patients may prefer not to undergo genetic testing, so genetic counseling should focus on this issue.

- Patients with cardiac channelopathies (i.e., long QT syndrome, catecholaminergic polymorphic ventricular tachycardia, Brugada syndrome, early repolarization syndrome, and short QT syndrome) and prior SCA have a significantly increased risk of subsequent SCA or SCD ([S7.9-7–S7.9-13,S7.9-25–S7.9-28](#)). Implantation of an ICD reduces the risk of death in high-risk patients ([S7.9-9,S7.9-29–S7.9-31](#)). Appropriate ICD therapy for VF/fast VT is reported in 8% to 33% of channelopathy patients, while inappropriate shocks and device complications are reported in 8% to 35% ([S7.9-10,S7.9-29,S7.9-30,S7.9-32–S7.9-36](#)). To minimize inappropriate shocks, concurrent beta blockers in long QT syndrome and catecholaminergic polymorphic ventricular tachycardia patients, optimal device programming, and appropriate lead selection are necessary. Ventricular pacing without ICD implantation was associated with a significant risk of recurrent SCA or SCD in long QT syndrome patients ([S7.9-37–S7.9-39](#)). In selected patients with LQT1 in whom the SCA occurred in the absence of beta-blocker treatment, beta-blocker therapy is offered as an alternative to ICD implantation in patients who refuse to receive an ICD ([S7.9-40](#)).

### 7.9.1. Specific Cardiac Channelopathy Syndromes

#### 7.9.1.1. Congenital Long QT Syndrome

## Recommendations for Long QT Syndrome

References that support the recommendations are summarized in [Online Data Supplement 40](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with long QT syndrome with a resting QTc greater than 470 ms, a beta blocker is recommended ( <a href="#">S7.9.1.1-1–S7.9.1.1-5</a> ).
I	B-NR	2. In high-risk patients with symptomatic long QT syndrome in whom a beta blocker is ineffective or not tolerated, intensification of therapy with additional medications (guided by consideration of the particular long QT syndrome type), left cardiac sympathetic denervation, and/or an ICD is recommended ( <a href="#">S7.9.1.1-2,S7.9.1.1-6–S7.9.1.1-12</a> ).
I	B-NR	3. In patients with long QT syndrome and recurrent appropriate ICD shocks despite maximum tolerated doses of a beta blocker, intensification of medical therapy with additional medications (guided by consideration of according to the particular long QT syndrome type) or left cardiac sympathetic denervation, is recommended ( <a href="#">S7.9.1.1-6,S7.9.1.1-7,S7.9.1.1-10,S7.9.1.1-13–S7.9.1.1-16</a> ).
I	B-NR	4. In patients with clinically diagnosed long QT syndrome, genetic counseling and genetic testing are recommended ( <a href="#">S7.9.1.1-17–S7.9.1.1-21</a> ).

(continued)

IIa	B-NR	5. In patients with suspected long QT syndrome, ambulatory electrocardiographic monitoring, recording the ECG lying and immediately on standing, and/or exercise treadmill testing can be useful for establishing a diagnosis and monitoring the response to therapy ( <a href="#">S7.9.1.1-22–S7.9.1.1-29</a> ).
IIa	B-NR	6. In asymptomatic patients with long QT syndrome and a resting QTc less than 470 ms, chronic therapy with a beta blocker is reasonable ( <a href="#">S7.9.1.1-3,S7.9.1.1-30,S7.9.1.1-31</a> ).
IIb	B-NR	7. In asymptomatic patients with long QT syndrome and a resting QTc greater than 500 ms while receiving a beta blocker, intensification of therapy with medications (guided by consideration of the particular long QT syndrome type), left cardiac sympathetic denervation or an ICD may be considered ( <a href="#">S7.9.1.1-2,S7.9.1.1-8,S7.9.1.1-11,S7.9.1.1-30</a> ).
III: Harm	B-NR	8. In patients with long QT syndrome, QT-prolonging medications are potentially harmful ( <a href="#">S7.9.1.1-5,S7.9.1.1-12,S7.9.1.1-32–S7.9.1.1-34</a> ).

**Table 10** and **Figures 9, 10, 11, and 12****Recommendation-Specific Supportive Text**

1. Beta blockers reduce adverse cardiac events for long QT syndrome type 1 (**Figure 10**) (>95%), long QT syndrome type 2 (**Figure 11**) (>75%), and females with long QT syndrome type 3 (**Figure 12**) by >60% ([S7.9.1.1-1–S7.9.1.1-5](#)). There are limited data regarding efficacy of beta blockers in males with long QT syndrome type 3 ([S7.9.1.1-3,S7.9.1.1-35,S7.9.1.1-36](#)) but, in selected patients, beta blockers can be protective against SCA ([S7.9.1.1-36,S7.9.1.1-37](#)). Several observational studies have reported effectiveness for risk reduction in long QT syndrome with propranolol, atenolol, and nadolol with appropriate dosing ([S7.9.1.1-26,S7.9.1.1-28, S7.9.1.1-38–S7.9.1.1-40](#)), while metoprolol appears less effective ([S7.9.1.1-41](#)). RCTs to assess comparative efficacy of specific beta blockers are unavailable, although many centers favor the use of nadolol. For long QT syndrome type 1, 1 study reported atenolol reduced risk of VA while nadolol was not associated with risk reduction ([S7.9.1.1-2](#)). For long QT syndrome

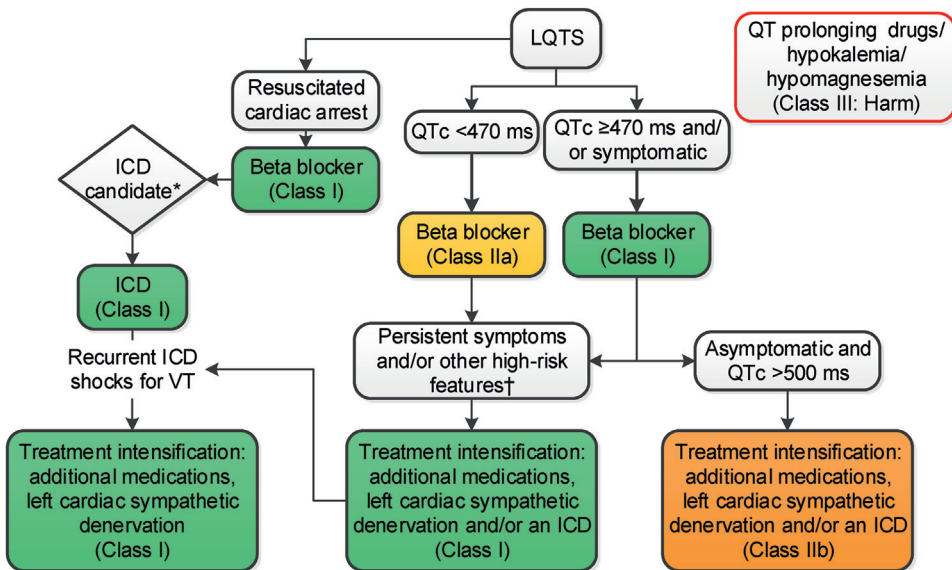
type 2, nadolol was reported to show superior efficacy ([S7.9.1.1-1,S7.9.1.1-2](#)). Patients receiving a beta blocker should undergo ongoing monitoring to assess changes in QTc over time, and adequacy of beta blockade with exertion ([S7.9.1.1-26,S7.9.1.1-28](#)).

2. High-risk patients with long QT syndrome include those with QTc >500 ms, genotypes long QT syndrome type 2 and long QT syndrome type 3, females with genotype long QT syndrome type 2, <40 years of age, onset of symptoms at <10 years of age, and patients with prior cardiac arrest or recurrent syncope ([S7.9.1.1-3, S7.9.1.1-8,S7.9.1.1-11,S7.9.1.1-30,S7.9.1.1-38](#)). Women with long QT syndrome type 2 are at a higher risk of postpartum cardiac arrest/SCD ([S7.9.1.1-42,S7.9.1.1-43](#)) and should receive prepregnancy counseling. Patients with long QT syndrome and recurrent syncope while receiving a beta blocker have an increased risk of SCA or appropriate ICD shocks ([S7.9.1.1-9](#)) and escalation of therapy is warranted to prevent SCD. Earlier studies reported benefit of antibradycardia pacing, with recurrent syncope or cardiac arrest reported in 7% to

**TABLE 10** Commonly Used QT-Prolonging Medications ([S7.9.1.1-59,S7.9.1.1-62](#))**Examples of QT Prolonging Medications\***

Antiarrhythmic Medications	Psychotropic Medications	Antibiotics	Others
Disopyramide	Haloperidol	Erythromycin	Methadone
Procainamide (N-acetylprocainamide)	Phenothiazines	Pentamidine	Probucol
Quinidine	Citalopram	Azithromycin	Droperidol
Dofetilide	Tricyclic antidepressants	Chloroquine	Ondansetron
Dronedarone		Ciprofloxacin	
Ibutilide		Fluconazole	
Sotalol		Levofloxacin	
Amiodarone†		Moxifloxacin	
		Clarithromycin	
		Itraconazole	
		Ketoconazole	

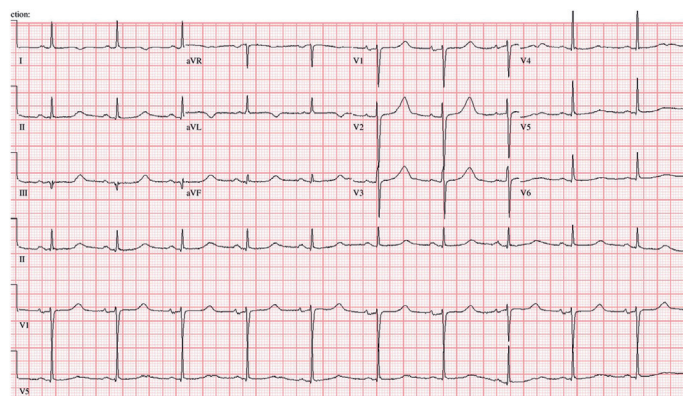
\*A more complete list is maintained at: [www.crediblemeds.org](http://www.crediblemeds.org) ([S7.9.1.1-59](#)). †Amiodarone rarely causes torsades de pointes.

**FIGURE 9** Prevention of SCD in Patients With Long QT Syndrome

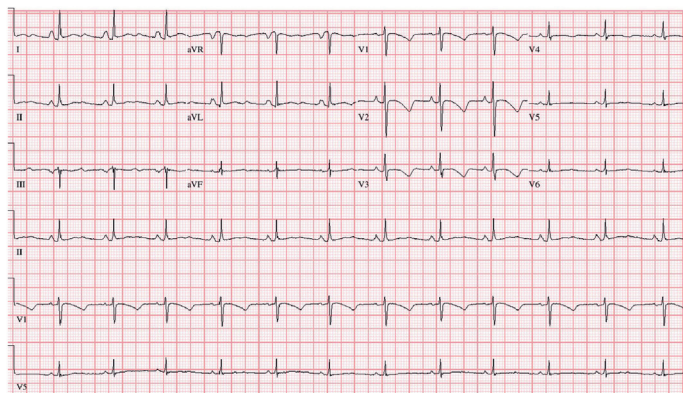
Colors correspond to Class of Recommendation in **Table 1**. See **Section 7.9.1.1** for discussion. \*ICD candidacy as determined by functional status, life expectancy, or patient preference. †High-risk patients with LQTS include those with QTc >500 ms, genotypes LQT2 and LQT3, females with genotype LQT2, <40 years of age, onset of symptoms at <10 years of age, and patients with recurrent syncope. ICD indicates implantable cardioverter-defibrillator; LQTS, long-QT syndrome; VT, ventricular tachycardia.

24% of patients ([S7.9.1.1-44–S7.9.1.1-47](#)). In high-risk patients, observational studies support effectiveness of the ICD in preventing SCD, with consideration of left cardiac sympathetic denervation to reduce the frequency of ICD shocks ([S7.9.1.1-16,S7.9.1.1-48, S7.9.1.1-49](#)). Left cardiac sympathetic denervation

can reduce VA burden, but up to 27% of high-risk patients experience at least 1 recurrence ([S7.9.1.1-16, S7.9.1.1-48,S7.9.1.1-50](#)). Left cardiac sympathetic denervation may be more effective in patients with long QT syndrome type 1 and long QT syndrome type 3 ([S7.9.1.1-16](#)). Complications related to left cardiac

**FIGURE 10** Long-QT Syndrome Type 1

**FIGURE 11** Long-QT Syndrome Type 2



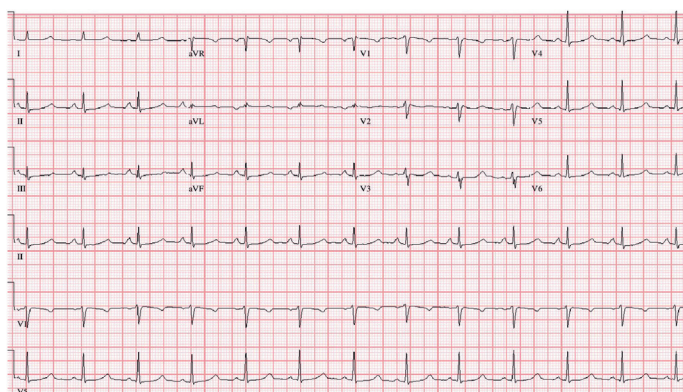
sympathetic denervation occur in 8% to 20% of patients ([S7.9.1.1-48](#),[S7.9.1.1-51](#)). Syncope in patients with long QT syndrome may occur due to vasovagal syncope, noncompliance with medications, or proarrhythmia from concurrent medications ([S7.9.1.1-5](#)). Clinical evaluation that incorporates consideration of genotype, QTc interval, medication compliance, and shared decision-making regarding the need to change or escalate therapy is important. Use of additional medications is guided by long QT syndrome type. In long QT syndrome type 3 ranolazine, mexiletine, and flecainide shorten the QTc and have been used to reduce recurrent arrhythmias ([S7.9.1.1-6](#),[S7.9.1.1-7](#),[S7.9.1.1-10](#)).

3. Mexiletine is an additional medication that can be used in patients with long QT syndrome and recurrent ICD shocks. Left cardiac sympathetic denervation is associated with a reduction the number of appropriate ICD shocks and VA burden ([S7.9.1.1-13](#)–[S7.9.1.1-16](#)).

Reduction of the QTc to <500 ms after left cardiac sympathetic denervation has been correlated with reduced risk of recurrent ICD shocks and frequency of symptoms ([S7.9.1.1-16](#),[S7.9.1.1-52](#)); however, SCD or SCA is reported in 3% to 10% of patients ([S7.9.1.1-15](#),[S7.9.1.1-16](#),[S7.9.1.1-48](#),[S7.9.1.1-50](#)). Although arrhythmia burden is often reduced, up to 27% of high-risk patients experience at least 1 recurrence ([S7.9.1.1-13](#),[S7.9.1.1-14](#),[S7.9.1.1-48](#)). Patient outcomes are improved if the left cardiac sympathetic denervation is performed in centers with surgical expertise in this procedure. Use of additional medications is guided by long QT syndrome type. In long QT syndrome type 3, ranolazine, mexiletine, and flecainide shorten the QTc and have been used to reduce recurrent arrhythmias ([S7.9.1.1-6](#),[S7.9.1.1-7](#),[S7.9.1.1-10](#)).

4. Genetic testing for disease-causing mutations in long QT syndrome offers important diagnostic, prognostic,

**FIGURE 12** Long-QT Syndrome Type 3



and therapeutic information in addition to the clinical evaluation, and a positive test can facilitate establishing risk for family members. The yield of genetic testing in long QT syndrome phenotype-positive patients is 50% to 86%, with the higher range present in patients with marked QT prolongation or positive family history of SCD ([S7.9.1.1-17,S7.9.1.1-21,S7.9.1.1-53](#)). A negative genetic test does not exclude the diagnosis of long QT syndrome, which relies on the clinical evaluation. In asymptomatic patients with otherwise unexplained prolonged QTc  $\geq 480$  ms on serial ECGs, genetic testing may help confirm the diagnosis and supplement prognostic information in addition to clinical symptoms and QTc duration ([S7.9.1.1-5,S7.9.1.1-18–S7.9.1.1-20,S7.9.1.1-30,S7.9.1.1-35,S7.9.1.1-54–S7.9.1.1-56](#)).

5. In a prospective, observational study of patients with suspected long QT syndrome, patients with a history of syncope or cardiac arrest and either an affected first-degree relative or a borderline or prolonged QTc interval underwent exercise treadmill testing and bicycle exercise, with ECGs recorded before, during, and after exercise, as well as in different positions ([S7.9.1.1-27](#)). long QT syndrome was confirmed by genetic testing in all affected individuals. Among patients with borderline-to-normal resting QTc intervals, prolongation of the 4-minute recovery QTc  $\geq 445$  ms had high sensitivity for correctly identifying patients with long QT syndrome ([S7.9.1.1-27](#)). A study in younger patients demonstrated QTc prolongation  $>460$  ms at 7 minutes of recovery predicted long QT syndrome type 1 or long QT syndrome type 2 patients versus controls ([S7.9.1.1-23](#)). In a study using burst bicycle exercise, patients with latent long QT syndrome had a significantly greater increase in QTc with exercise than either controls or those with QTc prolongation at baseline ([S7.9.1.1-24](#)). These findings can be useful in establishing whether long QT syndrome is present. Monitoring adequacy of beta-blocker therapy using exercise testing can be beneficial, particularly in school-aged patients ([S7.9.1.1-26,S7.9.1.1-28](#)). Beta-blocker therapy may be associated with a decrease in supine and peak exercise QTc, with the exception of long QT syndrome type 1 patients with C-loop mutations ([S7.9.1.1-25](#)).
6. Approximately 10% to 36% of genotype-positive patients with long QT syndrome have QTc intervals  $\leq 440$  ms, most commonly patients with long QT syndrome type 1 ([S7.9.1.1-31,S7.9.1.1-35](#)). Patients with long QT syndrome and normal QTc have a lower risk of VA and SCD compared to those with prolonged QTc ([S7.9.1.1-35](#)), but still have an increased risk of SCA or SCD compared with genotype-negative, age- and sex-matched general patients ([S7.9.1.1-31](#)). Beta blockers

reduce the risk of adverse cardiac events substantially ([S7.9.1.1-1–S7.9.1.1-5,S7.9.1.1-30,S7.9.1.1-36,S7.9.1.1-38,S7.9.1.1-41,S7.9.1.1-57](#)). During the periods of highest risk in the first 3 decades of life ([S7.9.1.1-11,S7.9.1.1-18](#)), treatment with a beta blocker may reduce risk of SCA ([S7.9.1.1-26,S7.9.1.1-28,S7.9.1.1-36,S7.9.1.1-38](#)). Changes in QTc occur over time, particularly during puberty and during and after pregnancy, indicating the need for assessment of QTc on ECG annually or with medication changes, and assessing medication efficacy with exercise testing as feasible. Asymptomatic adult (male) long QT syndrome patients with normal QTc intervals may choose to decline beta-blocker therapy ([S7.9.1.1-11,S7.9.1.1-34](#)).

7. The risk of adverse cardiac events from VA is influenced by the patient's resting QTc interval, age, sex, and long QT syndrome genotype/mutation. For asymptomatic males with long QT syndrome, the risk of cardiac events is highest in childhood ([S7.9.1.1-2,S7.9.1.1-8,S7.9.1.1-11,S7.9.1.1-30](#)), during a time when medication compliance is challenging. Young women with LQT2 and QTc  $>500$  ms are at increased risk of SCA ([S7.9.1.1-2,S7.9.1.1-11,S7.9.1.1-18–S7.9.1.1-20,S7.9.1.1-30,S7.9.1.1-35](#)) especially in the 9 months postpartum, and may be candidates for primary prevention ICD placement or use of a wearable cardioverter-defibrillator ([S7.9.1.1-30](#)).
8. The risk of adverse events increases in patients with long QT syndrome with prolongation of the QTc  $>500$  ms ([S7.9.1.1-2,S7.9.1.1-12,S7.9.1.1-26,S7.9.1.1-35,S7.9.1.1-41,S7.9.1.1-58](#)). QT-prolonging medications ([www.crediblemeds.org](http://www.crediblemeds.org)) ([S7.9.1.1-59](#)) should not be used in patients with long QT syndrome unless there is no suitable alternative; careful monitoring of the QTc during therapy is recommended, with consideration for discontinuing therapy with marked QTc prolongation. Concurrent use of stimulant or non-stimulant attention deficit/hyperactivity medications was associated with an increased risk of syncope/cardiac arrest in long QT syndrome, particularly males, in 1 study ([S7.9.1.1-34](#)), but it did not appear to be associated with increased risk in another retrospective study ([S7.9.1.1-60](#)). Episodes of torsades de pointes can be precipitated by exposure to a QT prolonging medication, or hypokalemia induced by diuretics or gastrointestinal illness. Attention to maintaining normal potassium and magnesium balance when medications or situations that promote depletion are encountered is an important component of management. Rare case reports exist of fever prolonging the QT interval in patients with long QT syndrome type 2; fever should be reduced with antipyretics ([S7.9.1.1-61](#)) (**Table 10**).

### 7.9.1.2. Catecholaminergic Polymorphic Ventricular Tachycardia

**Recommendations for Catecholaminergic Polymorphic Ventricular Tachycardia**  
References that support the recommendations are summarized in [Online Data Supplement 41](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with catecholaminergic polymorphic ventricular tachycardia, a beta blocker is recommended ( <a href="#">S7.9.1.2-1,S7.9.1.2-2</a> ).
I	B-NR	2. In patients with catecholaminergic polymorphic ventricular tachycardia and recurrent sustained VT or syncope, while receiving adequate or maximally tolerated beta blocker, treatment intensification with either combination medication therapy (e.g., beta blocker, flecainide), left cardiac sympathetic denervation, and/or an ICD is recommended ( <a href="#">S7.9.1.2-2–S7.9.1.2-6</a> ).
IIa	B-NR	3. In patients with catecholaminergic polymorphic ventricular tachycardia and with clinical VT or exertional syncope, genetic counseling and genetic testing are reasonable ( <a href="#">S7.9.1.2-7</a> ).

*Figure 13*

#### Recommendation-Specific Supportive Text

1. Catecholaminergic polymorphic ventricular tachycardia is characterized by exertion-related polymorphic or bidirectional VT ([Figure 13](#)), associated with syncope and SCA. SCA/SCD is reported in 3% to 13% of patients ([S7.9.1.2-1,S7.9.1.2-2,S7.9.1.2-8](#)). Treatment with beta blockers is associated with a reduction in adverse cardiac events ([S7.9.1.2-1,S7.9.1.2-2](#)). Some experts prefer the use of nadolol over other types of beta blockers; direct comparison data among beta blockers are unavailable. Use of a maximally tolerated dose of a beta blocker is important. Small observational studies suggest possible benefit of nondihydropyridine calcium channel blockers in the treatment of catecholaminergic polymorphic ventricular tachycardia ([S7.9.1.2-9,S7.9.1.2-10](#)).
2. Flecainide in combination with a beta blocker can suppress ventricular ectopy by as much as 76% in patients with catecholaminergic polymorphic ventricular tachycardia during exercise testing or clinical follow-up ([S7.9.1.2-2,S7.9.1.2-6,S7.9.1.2-11](#)). For refractory VA, verapamil or propafenone may also be effective ([S7.9.1.2-9,S7.9.1.2-10,S7.9.1.2-12](#)). ICD implantation in patients with catecholaminergic polymorphic ventricular tachycardia should be reserved for patients with prior SCA, or patients with refractory VAs on combination medical therapy. Inappropriate shocks are reported in 20% to 30% of catecholaminergic polymorphic ventricular tachycardia patients with ICDs ([S7.9.1.2-2,S7.9.1.2-13–S7.9.1.2-16](#)). ICD programming in patients with catecholaminergic polymorphic ventricular tachycardia should be optimized to deliver

therapy for VF and to minimize inappropriate shocks and the risk of potentially fatal electrical storms ([S7.9.1.2-13,S7.9.1.2-15](#)). Left cardiac sympathetic denervation for catecholaminergic polymorphic ventricular tachycardia may reduce the frequency of recurrent ICD shocks by 32% to 75% ([S7.9.1.2-3–S7.9.1.2-5,S7.9.1.2-17,S7.9.1.2-18](#)) although recurrent syncope, SCA, or SCD is reported in 9% to 32% of patients, with other minor complications in 20% to 70% of patients. It is best if the left cardiac sympathetic denervation is performed in centers with expertise in this procedure. Intensification of medical therapy or left cardiac sympathetic denervation is important in treating patients who present with recurrent appropriate ICD shocks ([S7.9.1.2-19](#)).

3. Genetic testing may be useful to confirm the diagnosis of catecholaminergic polymorphic ventricular tachycardia, which is suggested by the development of bidirectional VT with exertion or stress. Recognition of catecholaminergic polymorphic ventricular tachycardia as the cause for exertional symptoms should prompt aggressive therapy to prevent the significant risk of SCD. Therapy for catecholaminergic polymorphic ventricular tachycardia is not guided by genotype status, but screening of first-degree relatives may be facilitated with genetic testing ([S7.9.1.2-20](#)). Ryanodine receptor mutations have been reported in 47% of probands, which were de novo mutations in >70% ([S7.9.1.2-7](#)). Ryanodine genotype status has not correlated with disease severity or response to medications ([S7.9.1.2-7](#)). In very young patients presenting with idiopathic VF, mutations in calmodulin have been identified and are associated with high lethality ([S7.9.1.2-21–S7.9.1.2-24](#)). Studies of proposed

**FIGURE 13** Exercise-Induced Polymorphic VT in Catecholaminergic Polymorphic Ventricular Tachycardia



pathogenic mutations in catecholaminergic polymorphic ventricular tachycardia genes report up to 15% of variants were present in exome databases of the general population, raising questions as to the

monogenic cause of catecholaminergic polymorphic ventricular tachycardia (S7.9.1.2-20,S7.9.1.2-25).

#### 7.9.1.3. Brugada Syndrome

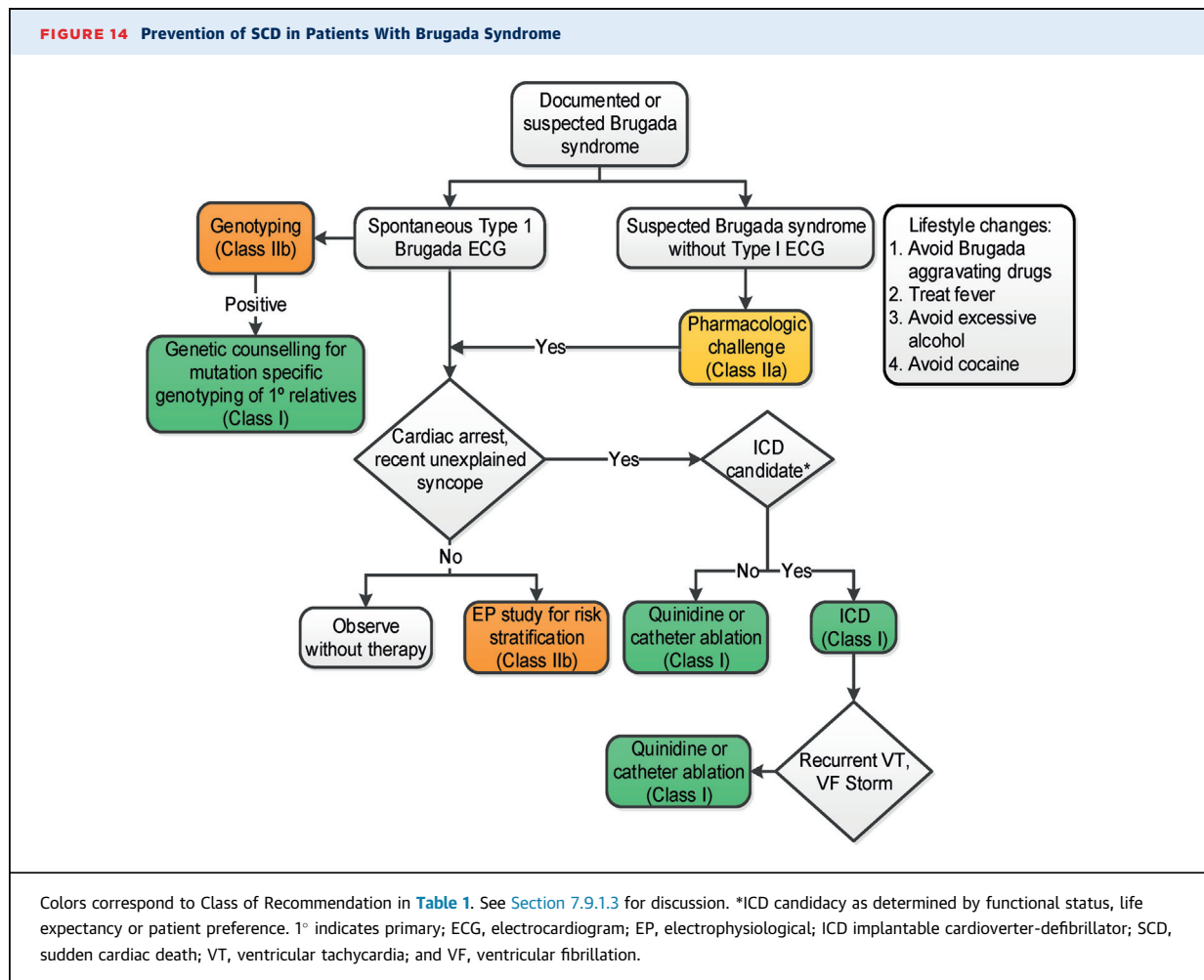
##### Recommendations for Brugada Syndrome

References that support the recommendations are summarized in [Online Data Supplement 42](#) and [Systematic Review Report](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In asymptomatic patients with only inducible type 1 Brugada electrocardiographic pattern, observation without therapy is recommended.
I	B-NR	2. In patients with Brugada syndrome with spontaneous type 1 Brugada electrocardiographic pattern and cardiac arrest, sustained VA or a recent history of syncope presumed due to VA, an ICD is recommended if meaningful survival of greater than 1 year is expected (S7.9.1.3-4,S7.9.1.3-6).
I	B-NR	3. In patients with Brugada syndrome experiencing recurrent ICD shocks for polymorphic VT, intensification of therapy with quinidine or catheter ablation is recommended (S7.9.1.3-7–S7.9.1.3-11).
I	B-NR	4. In patients with spontaneous type 1 Brugada electrocardiographic pattern and symptomatic VA who either are not candidates for or decline an ICD, quinidine or catheter ablation is recommended (S7.9.1.3-7,S7.9.1.3-9–S7.9.1.3-11).
IIa	B-NR	5. In patients with suspected Brugada syndrome in the absence of a spontaneous type 1 Brugada electrocardiographic pattern, a pharmacological challenge using a sodium channel blocker can be useful for diagnosis (S7.9.1.3-12–S7.9.1.3-14).
IIb	B-NR <sup>SR</sup>	6. In patients with asymptomatic Brugada syndrome and a spontaneous type 1 Brugada electrocardiographic pattern, an electrophysiological study with programmed ventricular stimulation using single and double extrastimuli may be considered for further risk stratification (S7.9.1.3-1,S7.9.1.3-6, S7.9.1.3-13,S7.9.1.3-15–S7.9.1.3-17).
IIb	C-EO	7. In patients with suspected or established Brugada syndrome, genetic counseling and genetic testing may be useful to facilitate cascade screening of relatives (S7.9.1.3-18–S7.9.1.3-20).

SR indicated systematic review.

Figures 14 and 15.

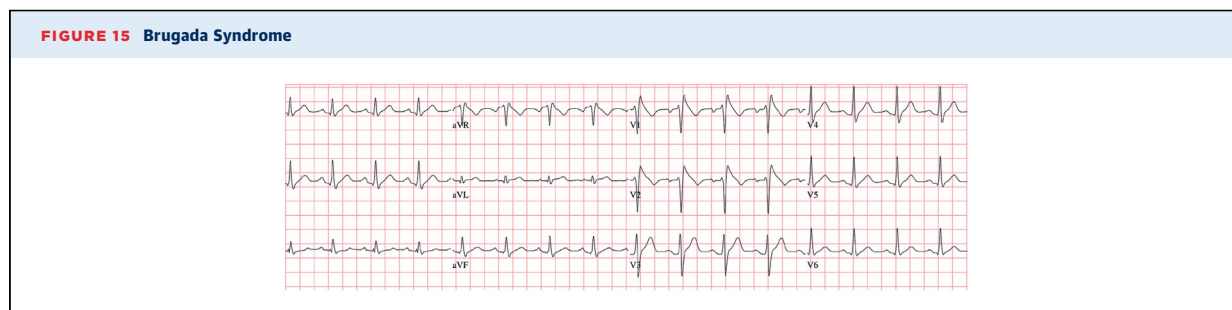


Colors correspond to Class of Recommendation in Table 1. See Section 7.9.1.3 for discussion. \*ICD candidacy as determined by functional status, life expectancy or patient preference. 1<sup>o</sup> indicates primary; ECG, electrocardiogram; EP, electrophysiological; ICD, implantable cardioverter-defibrillator; SCD, sudden cardiac death; VT, ventricular tachycardia; and VF, ventricular fibrillation.

## Synopsis

Refer to the “Systematic Review for the 2017 ACC/AHA/HRS Guideline for Management of Patients With Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death” for the complete systematic evidence review for additional data and analyses (S7.9.1.3-15). The results from the question “For asymptomatic patients with Brugada syndrome, what is the association between an

abnormal EP study and SCD and other arrhythmia endpoints? (Part 1)” and the writing committee’s review of the totality of the literature were used to frame decision-making. Recommendations that are based on a body of evidence that includes the systematic review conducted by the ERC are denoted by the superscript SR (e.g., LOE: B-R<sup>SR</sup>).



Factors identified as potential triggers of VF and SCA in Brugada syndrome include some psychotropic medications, and anesthetic agents, cocaine, excessive alcohol intake, and fever ([www.brugadadrugs.org](http://www.brugadadrugs.org)) (S7.9.1.3-21, S7.9.1.3-22). These agents should be avoided, and fever warrants early and aggressive measures to reduce temperature (S7.9.1.3-23).

#### Recommendation-Specific Supportive Text

1. The risk of major adverse cardiac events in asymptomatic patients without spontaneous type 1 electrocardiographic changes of Brugada syndrome (Figure 15), or with only medication-induced electrocardiographic changes, is low (S7.9.1.3-1–S7.9.1.3-5). A positive family history of Brugada syndrome or SCA is not a significant predictor of adverse events in Brugada syndrome (S7.9.1.3-1,S7.9.1.3-2,S7.9.1.3-4,S7.9.1.3-5). Implantation of an ICD in an asymptomatic patient without a spontaneous type 1 Brugada electrocardiographic has not been shown to confer any benefit.
2. Brugada syndrome is characterized by coved ST elevation in leads V1 or V2 positioned in the second, third, or fourth intercostal space either spontaneously or induced by administration of a sodium channel-blocking drug in the absence of other causes of ST elevation (S7.9.1.3-24) and negative T waves in the right precordial leads, and is associated with syncope or SCA due to VF, predominantly in young males, although it has been reported in all age groups. The type 1 Brugada ECG with coved ST elevation in right precordial leads may be present spontaneously, during fever or vagotonic states, or after medication challenge with sodium channel blockers. QRS complex fractionation is seen in a minority of patients. Patients with spontaneous coved type ST elevation and a history of syncope or prior SCA are at the highest risk for potentially lethal VA. ICD implantation has been shown to reduce mortality in symptomatic patients with Brugada syndrome (S7.9.1.3-25,S7.9.1.3-26).
3. Ablation of abnormal areas of epicardial late activation in the RV can suppress recurrent VA as shown in a small number of patients (S7.9.1.3-8,S7.9.1.3-9,S7.9.1.3-11, S7.9.1.3-27). In these reports, the spontaneous type 1 Brugada pattern on ECG may be eliminated in >75% of patients, and recurrences of VT/VF are markedly reduced (S7.9.1.3-9–S7.9.1.3-11). Experience and follow-up after ablation are limited, and an ICD for patients who have had syncope or SCA is recommended. A series of patients with Brugada syndrome treated with quinidine had no deaths during a mean follow-up of over 9 years, although adverse effects of quinidine were reported in 38% of patients, these authors felt that quinidine could be used as an alternative to the ICD in selected patients (S7.9.1.3-7).
4. Observational studies show that quinidine can suppress VF storm in patients with Brugada syndrome, and a low risk of arrhythmia was observed in a long-term observational study (681). No patient treated with quinidine experienced SCD. Adverse effects of quinidine occur in up to 37% of patients. Catheter ablation targeting the epicardial right ventricular areas of abnormality has also been shown to reduce recurrent VF episodes and normalize the ECG (682, 684, 685).
5. Administration of procainamide, flecainide, or ajmaline may be useful to provoke type 1 ST elevation in patients suspected to have Brugada syndrome as a cause of symptoms but who do not have a type 1 electrocardiographic pattern at baseline. Medication challenge should be terminated with the development of VA, marked QRS widening, or type 1 Brugada electrocardiographic pattern (S7.9.1.3-14,S7.9.1.3-28). The use of high electrocardiographic electrode positioning in the second and third interspaces for electrocardiographic recording improves detection of a type 1 Brugada ECG (S7.9.1.3-29). Asymptomatic patients with a family history of Brugada syndrome may be offered sodium channel blocker challenge for diagnostic evaluation, although a positive test does not require chronic therapy due to a low risk in this setting (S7.9.1.3-12). In asymptomatic patients with type 1 Brugada electrocardiographic findings, medication challenge does not offer additional diagnostic value.
6. Polymorphic VT/VF induced by programmed stimulation has been associated with an increased risk of VA in some patients with spontaneous type 1 Brugada ECG (S7.9.1.3-13). The specificity of programmed stimulation for assessing risk decreases with the inclusion of triple extrastimuli (S7.9.1.3-6,S7.9.1.3-13). The value of programmed stimulation in asymptomatic patients with spontaneous type 1 Brugada ECGs has been the subject of multiple studies (S7.9.1.3-1,S7.9.1.3-2,S7.9.1.3-4,S7.9.1.3-5). A report found that the prognostic value has decreased over time, possibly as patients with less severe phenotypes have been recognized and studied (S7.9.1.3-1). Some experts use the results of programmed ventricular stimulation for informing shared decision-making in consideration of the ICD. In symptomatic patients with Brugada syndrome, programmed ventricular stimulation for risk stratification does not add anything to the evaluation of the patients as an ICD is warranted (S7.9.1.3-2,S7.9.1.3-4, S7.9.1.3-6).
7. The yield of genetic testing in phenotype positive patients is approximately 20% to 30% in Brugada syndrome (S7.9.1.3-4,S7.9.1.3-16,S7.9.1.3-18,S7.9.1.3-19, S7.9.1.3-30,S7.9.1.3-31). SCN5A variants account for most of this subset of genotype positive Brugada syndrome. However, 2% to 10% of otherwise healthy individuals host a rare variant of SCN5A

(S7.9.1.3-20,S7.9.1.3-31). A negative genetic test does not exclude the diagnosis of Brugada syndrome, which is usually based on electrocardiographic and clinical characteristics. Risk stratification is based on symptoms and clinical findings (S7.9.1.3-32); genotype status is not correlated with the risk of adverse events (S7.9.1.3-5,S7.9.1.3-18,S7.9.1.3-19,S7.9.1.3-33). Identification of a pathogenetic mutation may help facilitate recognition of carrier status in family members, allowing for lifestyle modification and potential treatment.

#### Recommendations for Early Repolarization Syndrome

References that support the recommendations are summarized in Online Data Supplement 43.

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In asymptomatic patients with an early repolarization pattern on ECG, observation without treatment is recommended (S7.9.1.4-1,S7.9.1.4-2).
I	B-NR	2. In patients with early repolarization pattern on ECG and cardiac arrest or sustained VA, an ICD is recommended if meaningful survival greater than 1 year is expected (S7.9.1.4-3,S7.9.1.4-4).
III: No Benefit	B-NR	3. In patients with early repolarization pattern on ECG, genetic testing is not recommended (S7.9.1.4-5).

#### Recommendation-Specific Supportive Text

1. The prevalence of an early repolarization pattern on ECG with J point elevation in the inferior or lateral leads of at least 0.1 mV has been reported to be as high as 5.8% in adults (S7.9.1.4-1) and is more common in males. The early repolarization pattern was lost during 10-year follow-up in >60% of young males (S7.9.1.4-2). Patients are determined to have an early repolarization syndrome when, in addition to having early repolarization pattern on an ECG, they either have symptoms such as syncope or present with an arrhythmia. When patients present with an early repolarization pattern on an ECG, it is important to rule out reversible causes such as ischemia. Patients with early repolarization are more susceptible to the development of VF during

acute cardiac ischemia and/or in the presence of QRS abnormalities due to LV hypertrophy or bundle-branch block (S7.9.1.4-6–S7.9.1.4-8).

2. Patients with cardiac arrest or VF in the setting of an electrocardiographic pattern of early repolarization are at increased risk for subsequent recurrent episodes of VF, occurring in at least 40% of patients (S7.9.1.4-3, S7.9.1.4-4,S7.9.1.4-9). Antiarrhythmic medications, with the exception of quindine/hydroquinidine, have limited efficacy in preventing recurrent VA (S7.9.1.4-3, S7.9.1.4-4).

3. To date, genetic testing has not reliably identified mutations predisposing to early repolarization (S7.9.1.4-5).

#### 7.9.1.5. Short QT Syndrome

#### Recommendations for Short QT Syndrome

References that support the recommendations are summarized in Online Data Supplement 44.

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In asymptomatic patients with a short QTc interval, observation without treatment is recommended (S7.9.1.5-1,S7.9.1.5-2).
I	B-NR	2. In patients with short QT syndrome who have a cardiac arrest or sustained VA, an ICD is recommended if meaningful survival greater than 1 year is expected (S7.9.1.5-3–S7.9.1.5-5).

(continued)

IIa	C-LD	3. In patients with short QT syndrome and recurrent sustained VA, treatment with quinidine can be useful (S7.9.1.5-3,S7.9.1.5-5,S7.9.1.5-6).
IIa	C-LD	4. In patients with short QT syndrome and VT/VF storm, isoproterenol infusion can be effective (S7.9.1.5-7).
IIb	C-EO	5. In patients with short QT syndrome, genetic testing may be considered to facilitate screening of first-degree relatives (S7.9.1.5-4).

#### Recommendation-Specific Supportive Text

1. The prevalence of short QTc  $\leq 340$  ms is estimated to be 5 in 10,000 in persons  $< 21$  years of age and is more common in males (S7.9.1.5-1,S7.9.1.5-4,S7.9.1.5-8, S7.9.1.5-9). An incidental finding of a short QTc  $\leq 320$  ms in an asymptomatic patient warrants monitoring and follow-up without prophylactic medication treatment (S7.9.1.5-1,S7.9.1.5-2).
2. Patients with cardiac arrest in the setting of short QT syndrome are known to be at increased risk for recurrent cardiac arrest (S7.9.1.5-3–S7.9.1.5-5). Approximately 18% of the small number of reported patients with short QT and implanted ICDs have experienced appropriate ICD therapies during short-term follow-up (S7.9.1.5-3,S7.9.1.5-5,S7.9.1.5-6). Therapy with quinidine may reduce the number of ICD shocks (S7.9.1.5-3, S7.9.1.5-5,S7.9.1.5-6).
3. Markedly shortened QTc values  $\leq 300$  ms are associated with increased risk of SCD, especially during sleep or rest, in young persons, in whom the median QTc was 285 ms (S7.9.1.5-5,S7.9.1.5-9). A clinical score including QTc duration, clinical history of documented

polymorphic VT or VF, unexplained syncope, family history of autopsy-negative SCD or sudden infant death syndrome, and positive genotype results has been proposed to identify patients at increased risk for SCD (S7.9.1.5-4,S7.9.1.5-10). Treatment with quinidine results in lengthening of the QTc and, in selected patients, may be an alternative to ICD implantation (S7.9.1.5-3,S7.9.1.5-5,S7.9.1.5-6).

4. In the setting of electrical storm with refractory VF and short QT syndrome, infusion of isoproterenol can be effective in restoring/maintaining sinus rhythm (S7.9.1.5-7).
5. Pathogenic mutations in potassium channels have been identified in approximately 10% to 20% of patients with short QT syndrome including in KCNH2 (SQT1), KCNQ1 (SQT2), and KCNJ2 (SQT3) (S7.9.1.5-4). Due to the rarity of the disease, genotype/phenotype correlations are unavailable, limiting the use of knowledge of genotype status.

#### 8. VA IN THE STRUCTURALLY NORMAL HEART

##### Recommendations for VA in the Structurally Normal Heart

References that support the recommendations are summarized in [Online Data Supplement 45](#).

COR	LOE	RECOMMENDATION
I	B-R	1. In patients with symptomatic PVCs in an otherwise normal heart, treatment with a beta blocker or nondihydropyridine calcium channel blocker is useful to reduce recurrent arrhythmias and improve symptoms (S8-1,S8-2).
IIa	B-R	2. In patients with symptomatic VA in an otherwise normal heart, treatment with an antiarrhythmic medication is reasonable to reduce recurrent symptomatic arrhythmias and improve symptoms if beta blockers and nondihydropyridine calcium channel blockers are ineffective or not tolerated (S8-3,S8-4).

#### Synopsis

Most idiopathic VA are due to a focal mechanism of triggered activity or abnormal automaticity, some, notably interfascicular reentrant LV tachycardias, are due to reentry. The clinical manifestations of idiopathic VA are highly variable and range from benign, asymptomatic PVCs to sustained VT or even VF. On initial discovery, an

evaluation for structural heart disease is warranted with physical examination, an ECG, and imaging, usually with echocardiography. In the absence of any abnormality or a family history of SCD, further assessment and treatment are guided by symptoms. If the patient is asymptomatic and does not have evidence of a cardiac channelopathy, reassurance as to the benign nature is sufficient. If the

arrhythmia is suspected of being sufficiently frequent to cause ventricular dysfunction over time, periodic follow-up with reassessment of ventricular function is warranted (see [Section 10.8](#)). For mild symptoms, avoidance of aggravating factors such as excessive consumption of caffeine or sympathomimetic agents, may be sufficient. Therapy with a beta blocker or nondihydropyridine calcium channel blocker reduces symptoms for some patients. Class I antiarrhythmic medications can be effective, but those are generally avoided due to concerns for adverse effects. For patients who require arrhythmia suppression for whom antiarrhythmic medications are ineffective, not tolerated, or undesired, catheter ablation can be a highly effective treatment (see [Section 9](#)). The ablation strategy is to identify the site of origin manifested by the earliest site of electrical activation or, when this is not feasible, by pace-mapping. The most common site of origin for idiopathic VA is from the right ventricular outflow tract (RVOT) or the ostium of the LV, which is comprised of the oval opening of the LV to which the aorta is attached anteriorly and the left atrium is attached posteriorly. The likely origin can be reasonably predicted from the QRS morphology of the VA, which provides a good indication of the type of approach required and the likelihood of success and risks. Ablation failure is often related to the absence of the VA for mapping at the time of the procedure, or origin of the VA in an inaccessible

region of the heart. These foci occasionally produce sustained monomorphic VT ([S8-5–S8-7](#)).

#### Recommendation-Specific Supportive Text

1. In a randomized, double-blinded, placebo-controlled study of 52 patients with symptomatic VA and a mean PVC count of  $21,407 \pm 1740$  beats per 24 hours, atenolol significantly decreased symptom frequency ( $p=0.03$ ) and PVC count ( $p=0.001$ ), whereas placebo had no effect on PVC count ( $p=0.78$ ) or average heart rate ( $p=0.44$ ) ([S8-8](#)). A prospective randomized comparison of antiarrhythmic medications versus catheter ablation, metoprolol or propafenone had modest efficacy to suppress RVOT VA although with a far higher rate of recurrence than catheter ablation ([S8-9](#)).
2. In an RCT of 233 patients with  $\geq 30$  PVCs per hour, d-sotalol was shown to reduce frequent PVCs, but only racemic dl-sotalol is presently available ([S8-10](#)). In a prospective randomized comparison of antiarrhythmic medications versus catheter ablation, therapy with metoprolol or propafenone was shown to have modest efficacy when used to suppress RVOT PVCs although with a far higher rate of recurrence than catheter ablation ([S8-9](#)). Nondihydropyridine calcium channel blockers reduce arrhythmias ([S8-1,S8-2,S8-11,S8-12](#)).

#### 8.1. Outflow Tract and Atrioventricular Annular VA

##### Recommendations for Outflow Tract VA

References that support the recommendations are summarized in [Online Data Supplement 46](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	<ol style="list-style-type: none"><li>1. In patients with symptomatic outflow tract VA in an otherwise normal heart for whom antiarrhythmic medications are ineffective, not tolerated, or not the patient's preference, catheter ablation is useful (<a href="#">S8.1-1–S8.1-3</a>).</li></ol>
I	B-NR	<ol style="list-style-type: none"><li>2. In patients with symptomatic outflow tract VT in an otherwise normal heart, a beta blocker or a calcium channel blocker is useful (<a href="#">S8.1-1–S8.1-3</a>).</li></ol>

#### Recommendation-Specific Supportive Text

1. In 1 RCT, catheter ablation was superior to antiarrhythmic medications at suppressing frequent PVCs arising from the RVOT ([S8.1-4](#)). Observational studies have shown that radiofrequency catheter ablation is effective in the treatment of idiopathic VA arising from the RVOT and LV outflow tract ([S8.1-2,S8.1-5–S8.1-16](#)). The site of ablation may be below or above the pulmonic valve in the RVOT ([S8.1-9,S8.1-13](#)). Although most RVOT VA can be ablated within the RV, 10% may require ablation within the pulmonic sinus cusps ([S8.1-9](#)). Serious complications are infrequent. For LV outflow tract VA, the site of ablation may be within the aortic

cusp sinuses ([S8.1-11,S8.1-14,S8.1-16](#)), below the aortic valve ([S8.1-2,S8.1-6](#)), at the aorto-mitral continuity ([S8.1-1–S8.1-3](#)) or on the epicardial surface of the LV summit ([S8.1-3,S8.1-17,S8.1-18](#)). The mitral and tricuspid annulae are less common sites of idiopathic VA, but these VA can also be effectively treated with catheter ablation ([S8.1-1,S8.1-19,S8.1-20](#)). Approximately 10% of idiopathic VA may arise from the summit of the LV. Some can be ablated from the great cardiac vein or the epicardial surface, but others arise from an inaccessible region in close proximity to the left coronary artery precluding effective ablation ([S8.1-14](#)). Intramural sites of origin are infrequent but may require

ablation on both the endocardial and epicardial surfaces of the LV ostium ([S8.1-3](#)). Complications from ablation of outflow tract VA are infrequent, but bleeding complications related to arterial and venous access, pericardial tamponade, and damage to the coronary arteries can occur.

2. In a prospective randomized comparison of antiarrhythmic medications versus catheter ablation, meto-

prolol or propafenone was shown to have modest effectiveness when used to suppress RVOT PVCs, though with a far higher rate of recurrence than catheter ablation ([S8.1-4](#)). Non-dihydropyridine calcium channel blockers suppress arrhythmia in some patients ([S8.1-4](#)).

## 8.2. Papillary Muscle VA

### Recommendation for Papillary Muscle VA (PVCs and VT)

References that support the recommendation are summarized in [Online Data Supplement 47](#).

COR	LOE	RECOMMENDATION
I	B-NR	1. In patients with symptomatic VA arising from the papillary muscles for whom antiarrhythmic medications are ineffective, not tolerated, or not the patient's preference, catheter ablation is useful ( <a href="#">S8.2-1–S8.2-5</a> ).

### Recommendation-Specific Supportive Text

1. The papillary muscles of the LV or RV can be the site of origin of VA in the presence or absence of structural heart disease ([S8.2-1–S8.2-5](#)). Idiopathic left and right ventricular papillary muscle VA are most commonly PVCs and NSVT, and are usually exercise-related and may be induced by intravenous epinephrine or isoproterenol administration ([S8.2-3](#)). These arrhythmias have a focal, nonreentrant mechanism. Any of the 3 RV papillary muscles may be the site of origin and catheter ablation is usually effective ([S8.2-2](#)). In 1 study, successful ablation was achieved in all 8 patients with a reduction in PVC burden from  $17 \pm 20\%$  to  $0.6 \pm 0.8\%$  ([S8.2-2](#)). In the left ventricle, the site of origin may be

either the posteromedial or the anterolateral papillary muscles ([S8.2-1,S8.2-4,S8.2-5](#)). Multiple VA QRS morphologies were observed in 47% of patients, and ablation on both sides of the papillary muscle is required in some patients ([S8.2-4](#)). Achieving adequate catheter stability can be challenging. Acute ablation success is high, but recurrences are more frequent than for idiopathic outflow tract VA. Serious complications, including valve injury, appear to be infrequent. The risks of catheter ablation include bleeding related to arterial and venous access and a low risk of pericardial tamponade.

## 8.3. Interfascicular Reentrant VT (Belhassen Tachycardia)

### Recommendations for Interfascicular Reentrant VT (Belhassen Tachycardia)

References that support the recommendations are summarized in [Online Data Supplement 48](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients with verapamil-sensitive, idiopathic LVT related to interfascicular reentry for whom antiarrhythmic medications are ineffective, not tolerated, or not the patient's preference, catheter ablation is useful ( <a href="#">S8.3-1–S8.3-3</a> ).
I	B-NR	2. In patients with sustained hemodynamically tolerated verapamil-sensitive, idiopathic LVT related to interfascicular reentry, intravenous verapamil is recommended for VT termination ( <a href="#">S8.3-3–S8.3-6</a> ).
IIa	C-LD	3. In patients with recurrent verapamil-sensitive idiopathic LVT, chronic therapy with oral verapamil can be useful ( <a href="#">S8.3-7–S8.3-10</a> ).

### Recommendation-Specific Supportive Text

1. Idiopathic LVT is due to reentry involving a portion of the LV Purkinje system, usually the left posterior fascicle as the retrograde limb of the circuit and an incompletely defined segment of LV tissue as the anterograde limb, a portion of which is verapamil sensitive ([S8.3-1–S8.3-3](#)). These VTs are typically sustained

with a QRS that has a right bundle-branch block configuration with a superior axis. Less frequently an inferior axis VT or a relatively narrow QRS VT occurs as a result of alternate reentry paths, also involving a part of the Purkinje system. Beta blockers or verapamil typically terminate these arrhythmias, but they fail to prevent recurrences in some patients ([S8.3-1–S8.3-3](#)).

The target of catheter ablation for the most common form is usually the distal insertion of the anterograde limb of the Purkinje system along the inferior portion of the LV septum near its junction with the left posterior fascicle. Catheter ablation is acutely successful in >90% of patients with a risk of recurrence of approximately 10%. This VT may resemble fascicular VA that are due to a focal mechanism in the left anterior or left posterior fascicles of the LV His-Purkinje system. These fascicular arrhythmias usually have a focal mechanism with the target of catheter ablation being the site of earliest electrical activation recorded with a presystolic fascicular potential. Catheter ablation is highly effective for intrafascicular and fascicular VA. Serious complications are infrequent and include bleeding at the site of arterial or venous access and a small risk of bundle branch block or atrioventricular block.

2. Idiopathic LVT is based on reentrant mechanism involving tissue with slow conduction properties along the LV septum as the anterograde limb and the normal left posterior fascicle of the His-Purkinje system as the retrograde limb. The slow conduction zone is verapamil-sensitive ([S8.3-3–S8.3-6](#)). These arrhythmias typically have a right bundle-branch block morphology with superior axis, though reversal of the circuit may produce a relatively narrow QRS during VT. Verapamil typically terminates these arrhythmias in the anterograde slow conduction zone ([S8.3-3–S8.3-6](#)).
3. Although no RCTs have been published, the chronic use of oral verapamil for verapamil-sensitive idiopathic LVT has been reported to control this tachycardia in many patients, including both adults and children ([S8.3-5,S8.3-8–S8.3-10](#)).

#### 8.4. Idiopathic Polymorphic VT/VF

##### Recommendations for Idiopathic Polymorphic VT/VF

References that support the recommendations are summarized in [Online Data Supplement 49](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	<ol style="list-style-type: none"><li>1. In young patients (&lt;40 years of age) with unexplained SCA, unexplained near drowning, or recurrent exertional syncope, who do not have ischemic or other structural heart disease, further evaluation for genetic arrhythmia syndromes is recommended (<a href="#">S8.4-1–S8.4-8</a>).</li></ol>
I	B-NR	<ol style="list-style-type: none"><li value="2">2. In patients resuscitated from SCA due to idiopathic polymorphic VT or VF, an ICD is recommended if meaningful survival greater than 1 year is expected (<a href="#">S8.4-9–S8.4-13</a>).</li></ol>
I	B-NR	<ol style="list-style-type: none"><li value="3">3. For patients with recurrent episodes of idiopathic VF initiated by PVCs with a consistent QRS morphology, catheter ablation is useful (<a href="#">S8.4-11,S8.4-14</a>).</li></ol>

##### Recommendation-Specific Supportive Text

1. When combined with clinical evaluation, genetic testing can provide a diagnosis in up to 13% to 60% of younger (<40 years of age) survivors of SCA ([S8.4-3](#)), with the most common genotypes identified associated with long QT syndrome, catecholaminergic polymorphic ventricular tachycardia, and Brugada syndrome ([S8.4-8](#)). Drowning/near drowning events are particularly associated with LQT1 and catecholaminergic polymorphic ventricular tachycardia; genetic mutations in long QT syndrome and catecholaminergic polymorphic ventricular tachycardia have been identified in 23% of patients with unexplained near-drowning episodes ([S8.4-15](#)). In 1 study ([S8.4-6](#)), exertion-related cardiac arrest, particularly in children, may be related to long QT syndrome, catecholaminergic polymorphic ventricular tachycardia, or to calmodulin/triadin-mediated long QT syndrome/catecholaminergic polymorphic ventricular tachycardia mutations, which may

require additional specialized genetic testing ([S8.4-1, S8.4-2,S8.4-4,S8.4-16–S8.4-18](#)). Single-driver auto crashes should prompt the consideration of arrhythmic causes. The yield of genetic testing is higher if a family history of SCD at a young age is present. Referral to specialized genetic testing centers is important if local expertise is unavailable.

2. VF in the absence of identifiable structural heart disease or known genetic arrhythmia syndromes such as catecholaminergic polymorphic ventricular tachycardia, long QT syndrome, short QT syndrome, Brugada syndrome, or J wave syndromes is usually the result of short coupled PVCs arising from the Purkinje system in either the right or left ventricles or, less commonly, from the ventricular myocardium ([S8.4-9–S8.4-13](#)). The recurrence risk after resuscitation of idiopathic VF is very high ([S8.4-12](#)). Among 38 consecutive patients from 6 different centers who underwent ablation of primary idiopathic VF initiated by

short coupled PVC, 87% had experienced  $\geq 2$  VF episodes in the preceding year (**S8.4-12**). Because idiopathic VF is associated with a very high risk of recurrent VF, an ICD is indicated to prevent SCD. Catheter ablation of the triggering focus has proved to be highly effective in eliminating the repetitive PVCs which induce VF in these patients (**S8.4-11**). During a median post-procedural follow-up of 63 months, 7 (18%) of 38 patients undergoing catheter ablation of idiopathic VF induced by short coupled PVCs experienced VF recurrence at a median follow-up of 4 months. Five of these 7 patients underwent repeat ablation without VF recurrence. Thus, although catheter ablation is very effective in idiopathic VF, the recurrence risk remains substantial after an apparently successful procedure and the patient should be protected with an ICD. The subcutaneous ICD may not be a good therapy for these patients due to the higher risk of T-wave oversensing seen in this population; however, data are limited (**S8.4-10**).

3. Idiopathic VF may be initiated by PVCs that arise from the outflow tracts or the His-Purkinje system within either the right ventricle or left ventricle (**S8.4-11,S8.4-14, S8.4-19–S8.4-21**). Some patients have clusters of VF episodes (electrical storm) that typically present as PVCs initiating polymorphic VT/VF. The PVCs usually have a consistent QRS morphology and a short coupling

interval and can be targeted for ablation to control the arrhythmia (**S8.4-11**). For PVCs from the Purkinje system, the ablation target is a high-frequency Purkinje potential preceding the PVCs. When episodes are induced by short-coupled PVCs arising from the outflow tracts, the ablation target is the site of earliest ventricular activation. Patients with idiopathic VF often have periods of frequent VT/VF interspersed with periods of relative quiescence (**S8.4-11,S8.4-14**). To maximize the probability of successful ablation, the procedure is best performed during periods of frequent PVCs. Less-frequent episodes of VF may be amenable to ablation if frequent PVCs with a consistent QRS morphology are present. When the PVCs can be identified, ablation is highly successful, but late recurrences are observed in approximately 10% of patients such that implantation of an ICD is prudent even if ablation is acutely successful. The risks of catheter ablation include bleeding at the site of arterial or venous access and a small risk of pericardial tamponade. Therapy with quinidine acutely and chronically can suppress recurrent VF episodes in some patients (**S8.4-22**).

## 9. PVC-INDUCED CARDIOMYOPATHY

### Recommendations for PVC-Induced Cardiomyopathy

References that support the recommendations are summarized in [Online Data Supplement 50](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	<ol style="list-style-type: none"> <li>1. For patients who require arrhythmia suppression for symptoms or declining ventricular function suspected to be due to frequent PVCs (generally <math>&gt;15\%</math> of beats and predominately of 1 morphology) and for whom antiarrhythmic medications are ineffective, not tolerated, or not the patient's preference, catheter ablation is useful (<b>S9-1,S9-2</b>).</li> </ol>
IIa	B-NR	<ol style="list-style-type: none"> <li>2. In patients with PVC-induced cardiomyopathy, pharmacological treatment (e.g., beta blocker, amiodarone) is reasonable to reduce recurrent arrhythmias and improve symptoms and LV function (<b>S9-3,S9-4</b>).</li> </ol>

### Recommendation-Specific Supportive Text

1. Frequent PVCs (usually  $>15\%$  of the total number of beats) may produce a reversible form of LV dysfunction (**S9-5–S9-18**). However, it is sometimes difficult to ascertain whether the PVCs caused LV dysfunction or whether progressive LV dysfunction caused frequent PVCs. LV dysfunction has been associated with greater PVC burden ( $>10\%$  and usually  $>20\%$ ), NSVT, a retrograde P-wave after the PVCs, and interpolated

PVCs (**S9-6,S9-15**). In a prospective study of catheter ablation for PVC-induced cardiomyopathy, ablation was completely successful in 80% of patients (**S9-19**). LV function normalized within 6 months in 82% of the 22 patients who had depressed ventricular dysfunction at baseline. Thus, frequent PVCs may be a reversible cause of LV dysfunction that can be effectively treated with catheter ablation. It is often difficult to determine if apparent LV dysfunction reflects impaired LV function

- or inability to accurately assess LV function due to the frequent ectopic activity. In patients who have a high density of PVCs with normal ventricular function, optimal treatment and surveillance for prevention and detection of decline in ventricular function have not been established.
2. In a double-blind parallel study of 30 patients with or without ischemic heart disease with >30 PVCs per hour comparing sotalol to propranolol, proarrhythmic effects were present in 1 patient on sotalol. There was no significant difference in suppression of PVCs (sotalol 65%, propranolol 44%), with reduction in ventricular couplets being 99% for sotalol and 49% for propranolol. There was a significant increase in QTc in patients on sotalol ([S9-20](#)). In a double-blind, randomized, placebo-controlled study of 674 patients with HF and LVEF <0.40 attributed to ischemic or NICM and ≥10 PVCs per hour, amiodarone significantly reduced VA, slowed heart rate, and was associated with an increase in LVEF by 42% at 2 years with a nonsignificant trend toward reduction in mortality ([S9-4](#)). Whether the VA was contributing to ventricular dysfunction in these patients is unknown.

## 10. VA AND SCD RELATED TO SPECIFIC POPULATIONS

### 10.1. Athletes

In athletes, VAs range from isolated PVCs, couplets, and NSVT, to sustained VT and SCA leading to SCD ([S10.1-1](#)). Infrequent PVCs and short runs of repetitive NSVT, especially in the absence of structural heart disease, are more common in nonathletes, but they are generally benign, requiring only a limited workup and rarely lead to disqualification for sports ([S10.1-2,S10.1-3](#)). In contrast, longer runs of NSVT, especially when exercise-induced, and sustained VT and SCA/SCD are infrequent, but they have a higher incidence in athletes than that reported for the general population in the corresponding age groups. Reported estimates of SCD range from 1 per 53,703 athlete-years in the National Collegiate Athletic Association database ([S10.1-4](#)) to <1 per 200,000 in Minnesota high school students ([S10.1-5](#)). Among those studies judged to have better epidemiological protocols, estimates were in the range of 1 per 40,000 to 1 per 80,000 ([S10.1-6](#)). These figures compare with a general population risk of 1.0 to 1.9/100,000 in adolescents and young adults ([S10.1-7,S10.1-8](#)). Moreover, there appears to be both sport and sex differences in the magnitude of risk, with males being at higher risk than females in most

sports ([S10.1-7,S10.1-9](#)), blacks at higher risk than whites, and male basketball players being the single highest risk group in the United States, 1 per 5200 athlete-years ([S10.1-4](#)).

A study that included both competitive and recreational athletes showed that both groups are at a higher risk for SCD than the general population, with recreational athletes having greater cumulative numbers ([S10.1-7](#)), SCD occurring at an older age, and a different distribution of diseases. Postmortem data on SCD in athletes reveal that 25% to 40% are autopsy-negative, suggesting a role for genetic molecular disorders in these victims ([S10.1-4,S10.1-10,S10.1-11](#)) and for family members ([S10.1-12](#)).

Another limitation of SCD data analysis in athletes centers on noncardiac causes, some of which mimic cardiac events. Noncardiac causes include acute neurological disorders, drug abuse, heat stroke, rhabdomyolysis, sickle cell disorders, suicides, and accidents ([S10.1-13,S10.1-14](#)). Nonetheless, arrhythmias in athletes remain the most common medical cause of death and many occur as the first cardiac event.

The most common structural cause of SCAs and SCDs in athletes in the United States is HCM, followed by anomalous origins of coronary arteries, with myocarditis contributing a smaller but significant proportion ([S10.1-15](#)). Beyond these, the other inherited disorders contribute to the distribution of causes of a SCD in athletes, many of which can be suspected or identified by a careful family history and preparticipation ECGs.

In general, management of arrhythmias in athletes follows that in nonathletes. In regard to interventions, it is now generally recommended that AEDs be available at training and facilities for competitive athletes ([S10.1-16](#)), with less specific statements for AED availability at venues (e.g., tennis courts) or circumstances (e.g., jogging or small group runs) in which recreational athletics are occurring.

Many athletes who have had corrective procedures (repair of congenital or developmental defects such as anomalous origins of coronary arteries) ([S10.1-17,S10.1-18](#)) are on therapy for inherited disorders ([S10.1-19](#)) or have ICD implants ([S10.1-1](#)) and are able to participate in athletics depending on the nature and severity of the disease and with appropriate precautions and counseling regarding potential residual risks ([S10.1-19, S10.1-20](#)). For example, athletes with acquired disorders such as myocarditis are advised against exercise for at least 3 to 6 months after disease resolution.

## 10.2. Pregnancy

### Recommendations for Pregnancy

References that support the recommendations are summarized in [Online Data Supplement 51](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In mothers with long QT syndrome, a beta blocker should be continued during pregnancy and throughout the postpartum period including in women who are breastfeeding ( <a href="#">S10.2-1</a> ).
I	C-EO	2. In the pregnant patient with sustained VA, electrical cardioversion is safe and effective and should be used with standard electrode configuration ( <a href="#">S10.2-2,S10.2-3</a> ).
IIa	B-NR	3. In pregnant patients needing an ICD or VT ablation, it is reasonable to undergo these procedures during pregnancy, preferably after the first trimester ( <a href="#">S10.2-4,S10.2-5</a> ).

### Recommendation-Specific Supportive Text

1. Women with long QT syndrome should be counseled about maternal and fetal risks prior to pregnancy to ensure ongoing beta-blocker therapy. The risk of SCA or SCD is significantly higher during the 9 months after delivery, most notably among women with LQT2 ([S10.2-1,S10.2-6,S10.2-7](#)). A large retrospective analysis from the long QT syndrome registry demonstrated an odds ratio of 40.8 for syncope, SCA, or SCD among women with long QT syndrome in the 9 months' postpartum; treatment with beta blockers during pregnancy was independently associated with decreased risk ([S10.2-7](#)). Overall arrhythmic events during pregnancy are not increased among women receiving beta-blocker therapy ([S10.2-1,S10.2-6,S10.2-7](#)). In a case-control study, women with LQT1 who did not receive beta blockers during pregnancy, particularly those with prior syncope, were at significantly increased risk of SCA or syncope ([S10.2-8](#)). Frequency of events returned to prepregnancy levels after 9 months ([S10.2-1](#)). Maternal use of beta blockers during pregnancy is associated with decreased newborn birth weight and hypoglycemia ([S10.2-9](#)), but it is not associated with increased risk of miscarriage ([S10.2-8,S10.2-10](#)). Fetal bradycardia is associated with fetal long QT syndrome and should not independently provoke discontinuation of beta-blocker therapy ([S10.2-11–S10.2-14](#)); these infants are at increased risk of death

and require careful neonatal monitoring and treatment ([S10.2-13](#)). As 50% of offspring may be affected with long QT syndrome, with highest risk of adverse events in infancy and childhood, screening of the newborn at birth and during infancy for long QT syndrome is important ([S10.2-8](#)).

2. Available data on electrical fields associated with properly applied AED patches suggest that the fetus is safe; no observational data are available to the contrary. Anterolateral defibrillator pad placement is preferred with the lateral pad/paddle placed under the breast tissue, which is an important consideration in the pregnant patient.

3. The ICD in pregnant women is safe and effective ([S10.2-4](#)). For the rare circumstance of pregnant women with an immediate indication for an ICD, or less common indications for VT ablation during pregnancy, the radiation risk to the fetus is minimal ([S10.2-5,S10.2-15](#)). The procedure is usually performed after the first trimester unless there are circumstances that demand an earlier procedure. Wearable cardioverter-defibrillators have been used in peripartum cardiomyopathy while awaiting repeat assessment of recovery of ventricular function ([S10.2-16](#)). The subcutaneous implantable cardioverter-defibrillator is a potential alternative to conventional ICDs, although data are unavailable to support a recommendation.

## 10.3. Older Patients With Comorbidities

### Recommendation for Older Patients With Comorbidities

See Systematic Review Report ([S10.3-1](#)).

COR	LOE	RECOMMENDATION
IIa	B-NR <sup>SR</sup>	1. For older patients and those with significant comorbidities, who meet indications for a primary prevention ICD, an ICD is reasonable if meaningful survival of greater than 1 year is expected ( <a href="#">S10.3-1</a> ).

## Synopsis

Refer to the “Systematic Review for the 2017 ACC/AHA/HRS Guideline for Management of Patients With Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death” for the complete systematic evidence review for additional data and analyses ([S10.3-1](#)). The results from the question “What is the impact of ICD implantation for primary prevention in older patients and patients with significant comorbidities? (Part 2)” and the writing committee’s review of the totality of the literature were used to frame our decision-making. Recommendations are based on a body of evidence that includes the systematic review conducted by the ERC and are denoted by the superscript SR (e.g., LOE: B-R<sup>SR</sup>). Comorbidities included various combinations of renal disease, chronic obstructive pulmonary disease, atrial fibrillation, and heart disease, among others.

## Recommendation-Specific Supportive Text

1. Older age is defined as  $\geq 75$  years.

The ERC’s analyses are helpful in clearly demonstrating that neither age nor comorbidities alone should be exclusions for an ICD. However, the data included in the analysis are limited. Firstly, most data are from nonrandomized studies and “both selection and unidentified confounding biases can never be fully adjusted for.” It is likely that the more frail patients are already appropriately not offered ICDs and are thus not included. Secondly, because most of the studies are nonrandomized, these findings signify only an association and not causality.

Also, older adults are prone to higher complication rates, shorter life expectancies (and thus, fewer years during which they could derive benefit from an ICD), and varying preferences ([S10.3-2](#)). For these reasons, it is important to take a particularly nuanced and patient-centered approach to treating these patients.

## 10.4. Chronic Kidney Disease

Patients with chronic kidney disease (CKD) are at an increased risk of SCD compared with the general population, yet the risk versus benefit of primary prevention ICDs has been unclear; data from observational studies have been conflicting, and patients with moderate or severe CKD, especially patients with end-stage renal disease (ESRD) on dialysis were not included in the pivotal RCTs of ICDs ([S10.4-1–S10.4-5](#)). Furthermore, prior data had significant limitations given that patients who received ICDs have been compared inconsistently with a control group with CKD that did not receive primary prevention ICDs and the degree of renal

insufficiency likely influences survival benefit ([S10.4-6](#)). Patients with CKD, especially ESRD on dialysis, appear to be at increased risk of ICD-related complications. A significant number of sudden deaths are unassociated with VA in this population ([S10.4-7](#)). Therefore, the ERC was asked to address the impact of ICDs on mortality in patients with CKD.

The ERC conducted a specific analysis of 5 studies that explored renal dysfunction. A meta-analysis of these studies suggested that an association exists between ICD implantation and improved survival ([S10.4-8](#)). An important limitation is that only 2 studies specifically studied patients with ESRD and most data analyzed were from observational studies ([S10.4-8,S10.4-9](#)). In view of these limitations, the writing committee concluded there was not enough data to inform a recommendation on ICD implantation in patients with ESRD on dialysis. Decisions regarding ICDs in patients with CKD, especially those with ESRD, should be individualized and take into consideration the patient’s functional status, number of comorbidities, and preferences, among other factors.

## 10.5. Valvular Heart Disease

Patients with valvular heart disease should be evaluated and treated according to GDMT for valvular heart disease and, when LVEF is depressed, GDMT that applies to NICM to reduce the risk of SCD ([S10.6-23](#)). VA in patients with valvular heart disease can be caused by any of the mechanisms responsible for VA in other cardiac disease including ischemic heart disease, MI, severe LV hypertrophy, adrenergic-dependent rhythm disturbances, or an inherited molecular abnormality. Patients with valvular heart disease and VA are generally evaluated and treated using current recommendations for each disorder ([S10.6-1](#)). The presence of a VA alone does not constitute an indication for valve repair or replacement. In general, there is more knowledge on the risk for SCD in patients with aortic stenosis than other valvular lesions with a risk of 1% to 1.5% per year ([S10.6-2](#)). Most patients who die suddenly have been symptomatic from their valve disease ([S10.6-3,S10.6-4](#)). Although recurrent NSVT may place a patient with severe aortic stenosis at risk for syncope, the management of such a patient is guided by the severity of the valvular lesion.

Mitral valve prolapse has been implicated as a cause of SCD, although a study of 18,786 patients found no increased risk of SCA for patients with bileaflet mitral valve prolapse versus single leaflet mitral valve prolapse or no mitral valve prolapse ([S10.6-5](#)). LV fibrosis in the papillary muscles has been described in some mitral valve

prolapse patients with VA or SCD ([S10.6-6](#)). Further, a possible syndrome for SCD has been described that includes bileaflet mitral valve prolapse, female sex, T wave abnormality, and complex ventricular ectopy ([S10.6-7](#)). Guidance for treatment of patients with NICM, whether valvular or otherwise in origin, is provided in the current guideline (see [Sections 7.2.1 and 7.2.2](#) for primary and secondary prevention).

### **10.6. Sex-Related Differences in the Risk of SCD**

The information on associations between sex and VA and SCD is largely limited to epidemiological, cohort, and observational studies. Various population studies, primarily focused on SCD due to ischemic heart disease, have demonstrated age gradients in SCD risk among men and women ([S10.6-8–S10.6-10](#)). These include a 10-year lag in SCD incidence in women compared with men. However, risk factor burden among women has the same proportional effect as in men, with a 17-fold increase in risk from the lowest to highest deciles ([S10.6-9](#)). Importantly, 69% of the SCDs in women were first cardiac events ([S10.6-8](#)). A study of lifetime risk of SCD stratified at 45, 55, 65, and 75 years of age identified persistently lower and similar proportions of lifetime risk of SCD among women versus men in each of the strata ([S10.6-10](#)). The difference between women and men is somewhat smaller at ages below and above 75 years, largely because of a reduced risk in men. The overall lifetime risk of SCD was 1 in 9 among men and 1 in 30 among women ([S10.6-10](#)).

In studies of outcomes after out-of-hospital cardiac arrest, women were older, had more SCAs in homes, and fewer shockable rhythms (VT/VF) than men ([S10.6-11](#), [S10.6-12](#)). This was associated with a somewhat lower probability of survival overall; however, women with VT/VF and those with pulseless electrical activity had better outcomes than men ([S10.6-12](#)). A retrospective analysis of out-of-hospital cardiac arrest reported that survival improved over a 10-year period, with more favorable outcomes in men as well as younger women ([S10.6-13](#)).

Two studies demonstrated better outcomes in women with VT/VF, despite adverse risk factor profiles in women ([S10.6-14](#),[S10.6-15](#)). Another large study demonstrated that despite similar prehospital return of spontaneous circulation and survival to discharge, younger women had lower 1-month neurologically intact survival than the 50 to 60 age group ([S10.6-16](#)). A 17-year retrospective analysis did not demonstrate any difference between men and women, although total outcomes improved ([S10.6-17](#)).

The proportion of ischemic heart disease-associated SCAs among women surviving out-of-hospital cardiac arrest was significantly lower than in men, but ischemic heart disease remained the most powerful predictor etiologically ([S10.6-18](#)), and women were also significantly less likely to have severe LV dysfunction (LVEF  $\leq$ 35%) or previously recognized ischemic heart disease ([S10.6-19](#)). Women appear to be less likely to benefit from therapeutic hypothermia postcardiac arrest; however, in the younger age group, neurologic recovery in women was better than in older women ([S10.6-20](#)). Women are less likely to have SCA during competitive athletic events. A large study including both recreational and competitive athletes across a large age range noted that SCA in women during athletic events was 1 in 20 of that in men ([S10.6-21](#)).

A large literature review from 1980 to 1992 demonstrated that women accounted for 70% of recorded cases of cardiovascular medication-related arrhythmias ([S10.6-22](#)). This is consistent with QT interval differences among men and women. A retrospective analysis of quinidine discontinuation reported a significant difference in discontinuation between men and women (66% versus 84%) largely due to prolonged QT ([S10.6-23](#)). A study of catheter ablation for VT reported that overall outcome was similar between men and women ([S10.6-24](#)). The only sex difference was the greater probability of women having RVOT VT and a greater probability of men having LV outflow tract VT.

### **10.7. Medication-Induced Arrhythmias**

#### **Recommendations for Medication-Induced Arrhythmias**

References that support the recommendations are summarized in [Online Data Supplement 52 and 53](#).

#### **Digoxin**

COR	LOE	RECOMMENDATION
I	B-NR	1. Administration of digoxin antibodies is recommended for patients who present with sustained VA potentially due to digoxin toxicity ( <a href="#">S10.7-1</a> , <a href="#">S10.7-2</a> ).

(continued)

#### Medication-Induced QT Prolongation and Torsades de Pointes

COR	LOE	RECOMMENDATIONS
I	B-NR	2. In patients with recurrent torsades de pointes associated with acquired QT prolongation and bradycardia that cannot be suppressed with intravenous magnesium administration, increasing the heart rate with atrial or ventricular pacing or isoproterenol are recommended to suppress the arrhythmia ( <a href="#">S10.7-3</a> ).
I	C-LD	3. For patients with QT prolongation due to a medication, hypokalemia, hypomagnesemia, or other acquired factor and recurrent torsades de pointes, administration of intravenous magnesium sulfate is recommended to suppress the arrhythmia ( <a href="#">S10.7-4,S10.7-5</a> ).
I	C-LD	4. For patients with torsades de pointes associated with acquired QT prolongation, potassium repletion to 4.0 mmol per L or more and magnesium repletion to normal values (e.g., $\geq 2.0$ mmol/L) are beneficial ( <a href="#">S10.7-6,S10.7-7</a> ).

#### Sodium Channel Blocker-Related Toxicity

COR	LOE	RECOMMENDATIONS
IIa	C-LD	5. In patients taking sodium channel blockers who present with elevated defibrillation or pacing thresholds, discontinuing the presumed responsible medication or reprogramming the device can be useful to restore effective device therapy ( <a href="#">S10.7-8,S10.7-9</a> ).
III: Harm	B-NR	6. In patients with congenital or acquired long QT syndrome, QT-prolonging medications are potentially harmful ( <a href="#">S10.7-10</a> ).

#### Recommendation-Specific Supportive Text

1. Typical arrhythmias related to digoxin toxicity include enhanced atrial, junctional, or ventricular automaticity (with ectopic beats or tachycardia) often combined with atrioventricular block ([S10.7-11](#)). VT that is fascicular or bidirectional in origin is suggestive of digoxin toxicity ([S10.7-12](#)). Severe digoxin overdose causes hyperkalemia and cardiac standstill. The diagnosis is established by the combination of characteristic rhythm disturbances, ancillary symptoms (visual disturbances, nausea, changes in mentation), and elevated serum concentrations. Potentiating factors may include hypothyroidism, hypokalemia, or renal dysfunction ([S10.7-12](#)). Treatment of digoxin toxicity is based on the severity. In mild cases, discontinuing the medication, monitoring rhythm, and maintaining normal serum potassium may be sufficient ([S10.7-11](#)). Intravenous magnesium is often administered if VAs are present ([S10.7-12](#)). Occasionally, temporary pacing may be needed for atrioventricular block or asystole ([S10.7-13](#)). For more severe intoxication (serum digoxin concentrations exceeding 4 ng/mL and with serious arrhythmias such as VT), the treatment of choice is digoxin-specific Fab antibody ([S10.7-1](#)). In 1 series of 150 severely intoxicated patients, response was rapid (30 minutes to 4 hour), and 54% of patients presenting

with a cardiac arrest survived hospitalization ([S10.7-1](#)). Adverse effects include worsening of the underlying disease (increased ventricular rate during AF, exacerbation of HF) and hypokalemia. Doses lower (and less expensive) than the full neutralizing dose are sufficient as long as cardiac arrest is not imminent ([S10.7-2](#)). Digoxin concentration monitoring is unreliable after antidigoxin antibody administration.

2. Monitoring high-risk patients during initiation of QT-prolonging antiarrhythmic medications and recognition of the syndrome when it occurs are the first steps. Temporary pacing is highly effective in managing torsades de pointes that is recurrent after potassium and magnesium supplementation ([S10.7-3](#)). Isoproterenol can also be used to increase heart rate and abolish postectopic pauses ([S10.7-3](#)).
3. Intravenous magnesium can suppress episodes of torsades de pointes without necessarily shortening QT, even when serum magnesium is normal ([S10.7-4,S10.7-5](#)). Repeated doses may be needed, titrated to suppress ectopy and nonsustained VT episodes while precipitating factors are corrected ([S10.7-4](#)). Magnesium toxicity (areflexia progressing to respiratory depression) can occur at high serum concentrations, but this risk is very small with the doses usually used to treat torsades de pointes, 1 to 2 g intravenously ([S10.7-14](#)).

Allelic variants in clinical long-QT disease genes have been identified in patients with medication-induced torsades de pointes ([S10.7-7](#),[S10.7-15](#)–[S10.7-18](#)). Further, whole exome sequencing implicates an increased burden of rare potassium channel variants in the risk of medication-induced torsades de pointes ([S10.7-17](#),[S10.7-19](#)). These findings do not yet support general genetic screening for prediction of medication-induced torsades de pointes. In long QT syndrome, genetic testing may be performed in the index case who experienced medication-induced torsades de pointes and, if he/she did not survive that event, electrocardiographic screening of first-degree relatives may be performed.

4. Maintaining serum potassium between 4.5 mEq/L and 5 mEq/L shortens QT and may reduce the chance of recurrent torsades de pointes ([S10.7-6](#),[S10.7-7](#)).
5. In large clinical trials, sodium channel blockers increased mortality among patients convalescing from MI ([S10.7-20](#)), but similar trends were also seen with earlier trials of mexiletine ([S10.7-21](#)) and disopyramide ([S10.7-22](#)). Based on CAST, flecainide is contraindicated in patients with ischemia, prior MI, and is avoided in patients with other structural heart diseases ([S10.7-20](#)).

Sodium channel blockers increase defibrillation energy requirement and pacing thresholds ([S10.7-8](#),[S10.7-9](#)); as a consequence, patients may require reprogramming or revision of pacing or ICD systems or changes in their medication regimens (although modern pacing systems that provide automatic pacing threshold testing and adjustment of pacing output have mitigated the risk of loss of capture). Sodium channel blockers can “convert” AF to slow atrial flutter, which can show 1:1 atrioventricular-

ular conduction with wide QRS complexes that can be confused with VT ([S10.7-23](#)).

Sodium channel blockers, like procainamide and flecainide, can occasionally precipitate the typical Brugada syndrome ECG ([S10.7-24](#),[S10.7-25](#)). This has been reported not only with antiarrhythmic medications but also with tricyclic antidepressants ([S10.7-26](#)) and cocaine ([S10.7-27](#)) ([www.brugadadrugs.org](http://www.brugadadrugs.org)) ([S10.7-28](#)). Whether this represents unmasking individuals with clinically unapparent Brugada syndrome (see [Section 7.9.1.3](#)) or one end of a broad spectrum of responses to sodium channel blockers is unknown.

In the setting of sodium-channel blocker toxicity, limited animal data suggest that administration of sodium, as sodium chloride or sodium bicarbonate, may improve conduction slowing or suppress frequent or cardioversion-resistant VT ([S10.7-29](#)). Successful treatment with beta blockers ([S10.7-30](#)) and intravenous fat emulsion and/or extracorporeal membrane oxygenation has also been reported ([S10.7-31](#)).

6. QT-prolonging medications ([www.crediblemeds.org](http://www.crediblemeds.org)) ([S10.7-32](#)) are not used in patients with congenital or acquired long QT syndrome unless there is no suitable alternative or the benefit greatly exceeds the risk. Episodes of torsades de pointes can be precipitated by exposure to a QT-prolonging medication, and underlying prolongation of the QT (from genetic and clinical risk factors) increases this risk ([S10.7-10](#)). Medications implicated in torsades de pointes are found in several medication classes, including antiarrhythmics, antihistamines, antibiotics, antifungals, antidepressants, antipsychotics, opiates, and anticancer agents ([S10.7-10](#)) ([Table 10](#)).

## 10.8. Adult Congenital Heart Disease

### Recommendations for Adult Congenital Heart Disease

References that support the recommendations are summarized in [Online Data Supplement 54](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	<ol style="list-style-type: none"><li>1. Adult patients with repaired complex congenital heart disease presenting with frequent, complex, or sustained VA, or unexplained syncope should undergo evaluation for potential residual anatomic or coronary abnormalities (<a href="#">S10.8-1</a>–<a href="#">S10.8-6</a>).</li></ol>
I	B-NR	<ol style="list-style-type: none"><li>2. In patients with adult congenital heart disease and complex or sustained VA in the presence of important residual hemodynamic lesions, treatment of hemodynamic abnormalities with catheter or surgical intervention as feasible is indicated prior to consideration of ablation or an ICD (<a href="#">S10.8-3</a>,<a href="#">S10.8-7</a>–<a href="#">S10.8-12</a>).</li></ol>
I	B-NR	<ol style="list-style-type: none"><li>3. In patients with adult congenital heart disease and hemodynamically unstable VT, an ICD is recommended after evaluation and appropriate treatment for residual lesions/ventricular dysfunction if meaningful survival of greater than 1 year is expected (<a href="#">S10.8-13</a>–<a href="#">S10.8-17</a>).</li></ol>
I	B-NR	<ol style="list-style-type: none"><li>4. In patients with adult congenital heart disease with SCA due to VT or VF in the absence of reversible causes, an ICD is recommended if meaningful survival of greater than 1 year is expected (<a href="#">S10.8-13</a>–<a href="#">S10.8-17</a>).</li></ol>

(continued)

IIa	B-NR	5. In adults with repaired tetralogy of Fallot physiology with high-risk characteristics and frequent VA, an electrophysiological study can be useful to evaluate the risk of sustained VT/VF ( <a href="#">S10.8-18,S10.8-19</a> ).
IIa	B-NR	6. In adults with repaired tetralogy of Fallot physiology and inducible VT/VF or spontaneous sustained VT, implantation of an ICD is reasonable if meaningful survival greater than 1 year is expected ( <a href="#">S10.8-1,S10.8-19,S10.8-20</a> ).
IIa	B-NR	7. In patients with adult congenital heart disease with recurrent sustained monomorphic VT or recurrent ICD shocks for VT, catheter ablation can be effective ( <a href="#">S10.8-21–S10.8-25</a> ).
IIa	B-NR	8. In adults with repaired severe complexity adult congenital heart disease and frequent or complex VA, a beta blocker can be beneficial to reduce the risk of SCA ( <a href="#">S10.8-26</a> ).
IIa	B-NR	9. In patients with repaired moderate or severe complexity adult congenital heart disease with unexplained syncope and at least moderate ventricular dysfunction or marked hypertrophy, either ICD implantation or an electrophysiological study with ICD implantation for inducible sustained VA is reasonable if meaningful survival of greater than 1 year is expected ( <a href="#">S10.8-5,S10.8-16,S10.8-27–S10.8-29</a> ).
IIb	B-NR	10. In patients with adult congenital heart disease and severe ventricular dysfunction (LVEF <35%) and symptoms of heart failure despite GDMT or additional risk factors, ICD implantation may be considered if meaningful survival of greater than 1 year is expected ( <a href="#">S10.8-14–S10.8-16,S10.8-20</a> ).
III: Harm	B-NR	11. In patients with adult congenital heart disease who have asymptomatic VA, prophylactic antiarrhythmic therapy with class Ic medications (i.e., flecainide, propafenone) or amiodarone is potentially harmful ( <a href="#">S10.8-30–S10.8-32</a> ).

*Table 11 and Figure 16*

**Synopsis**

Tetralogy of Fallot (TOF) is defined as, congenital heart disease with RVOT obstruction and ventricular septal defect, often requiring right ventricle to pulmonary artery conduit placement or pulmonary valve replacement; includes TOF and double-outlet right ventricle. Moderate complexity congenital heart disease is defined as congenital heart disease requiring intracardiac surgical repair, other than isolated atrial and ventricular septal defects; includes TOF, aortic stenosis, coarctation of aorta, and Ebstein anomaly of the tricuspid valve. Severe complexity congenital heart disease is defined as cyanotic congenital heart disease requiring intracardiac repair in infancy, often with staged procedures; includes transposition of the great arteries, truncus arteriosus, and single ventricle anatomy ([Figure 16](#)).

**Recommendation-Specific Supportive Text**

1. The association of VT with RV hemodynamic abnormalities was first established in patients with repaired TOF ([S10.8-33](#)). Multiple studies since that time have demonstrated the correlation of hemodynamic residue and ventricular dysfunction with risk of VT or SCD in patients with congenital heart disease ([S10.8-1, S10.8-3–S10.8-6,S10.8-18,S10.8-34–S10.8-36](#)). Presentation with frequent or complex VA may indicate worsening hemodynamic function, coronary artery compromise, or decreased perfusion in the setting of

ventricular hypertrophy. Evaluation may also include exercise testing to assess functional capacity ([S10.8-35](#)). Careful evaluation of hemodynamic status for optimization of management is important ([S10.8-9](#)). Potentially treatable residual hemodynamic problems may be identified during hemodynamic evaluation, such as outflow tract stenosis or significant regurgitation, which may benefit from either catheter or surgical intervention ([S10.8-3,S10.8-7,S10.8-10, S10.8-12,S10.8-37](#)). Patients with markedly reduced ventricular function, elevated end-diastolic pressures, or pulmonary hypertension should be treated for underlying hemodynamic problems as part of their arrhythmia management.

2. The correlation of residual hemodynamic abnormalities with VA has been most extensively studied in patients with repaired TOF, where RV hypertension, residual pulmonary outflow tract obstruction or regurgitation, and RV dilation are risk factors for VT/SCD ([S10.8-1,S10.8-2,S10.8-4,S10.8-8,S10.8-33,S10.8-34, S10.8-36](#)). In these studies, frequent PVCs correlated with risk of clinical or inducible sustained VT. A combined approach of surgery for structural abnormalities with map-guided arrhythmia surgery has been used with success ([S10.8-3,S10.8-8,S10.8-10, S10.8-12](#)), but elimination of VT circuits may be limited by deep endocardial or LV origin of VT and limitations of operative mapping; an empiric approach to VT surgery is generally not recommended

**TABLE 11** Congenital Heart Disease: Risk Factors for VA/SCD

Congenital Heart Disease	Incidence of VA	Incidence of SCD	Higher Risk Characteristics
<b>Simple complexity</b>			
<b>ASD (S10.8-44,S10.8-47,S10.8-57–S10.8-62)</b>	2%–6%	<1.5%	Ventricular pacing RV dilatation Pulmonary hypertension NKX2.5 gene
<b>VSD (S10.8-27,S10.8-44,S10.8-47,S10.8-57–S10.8-63)</b>	3%–18%	<3%	
<b>Moderate complexity</b>			
<b>Tetralogy of Fallot (S10.8-1,S10.8-2,S10.8-5,S10.8-6,S10.8-28, S10.8-34,S10.8-36,S10.8-44,S10.8-46,S10.8-47, S10.8-54–S10.8-56,S10.8-62–S10.8-65)</b>	14%–31%	1.4%–8.3%	Unexplained syncope Frequent or complex VA Sustained VT QRS duration ≥180 ms Inducible sustained VT Atrial tachycardia Decreased LVEF Dilated right ventricle Severe PR Severe PS
<b>Aortic stenosis (S10.8-27,S10.8-44,S10.8-56)</b>	10%–34%	3%–20%	Unexplained syncope Severe LV hypertrophy Aortic stenosis mean pressure gradient >40 mm Hg Ventricular dysfunction
<b>Coarctation of aorta (S10.8-28,S10.8-29,S10.8-44,S10.8-46, S10.8-56,S10.8-62)</b>	2%	2%	Aneurysm at repair site Aortic stenosis Systemic hypertension Premature coronary artery disease
<b>Ebstein's anomaly (S10.8-45,S10.8-47,S10.8-55)</b>	2%	3%–6%	Cardiomegaly Atrial fibrillation Wide complex tachycardia Mitral regurgitation Dilated RVOT
<b>Severe complexity</b>			
<b>Transposition of the great arteries (S10.8-27,S10.8-44–S10.8-48, S10.8-51,S10.8-55,S10.8-56,S10.8-62)</b>			Atrial switch Mustard repair Prior VSD closure Unexplained syncope Atrial tachycardia Coronary orifice stenosis Systemic ventricular dysfunction Severe tricuspid regurgitation
<b>Atrial switch</b>	2%	3%–9.5%	
<b>Arterial switch</b>	2%	1%	
<b>cc-TGA</b>	10%	17%–25%	
<b>Truncus arteriosus (S10.8-66,S10.8-67)</b>	10%	4%	Multiple surgical repairs Coronary anomalies Ventricular dysfunction and/or hypertrophy
<b>Fontan repair for univentricular physiology* (S10.8-27,S10.8-37, S10.8-44,S10.8-45,S10.8-47,S10.8-55,S10.8-68)</b>	5%–17%	2.8%–5.4%	Atrial tachycardia Longer duration of follow-up Ascites Protein-losing enteropathy

\*Univentricular physiology includes: Tricuspid atresia, Double inlet left ventricle, Mitral atresia, Hypoplastic left heart, Unbalanced AV septal defect.

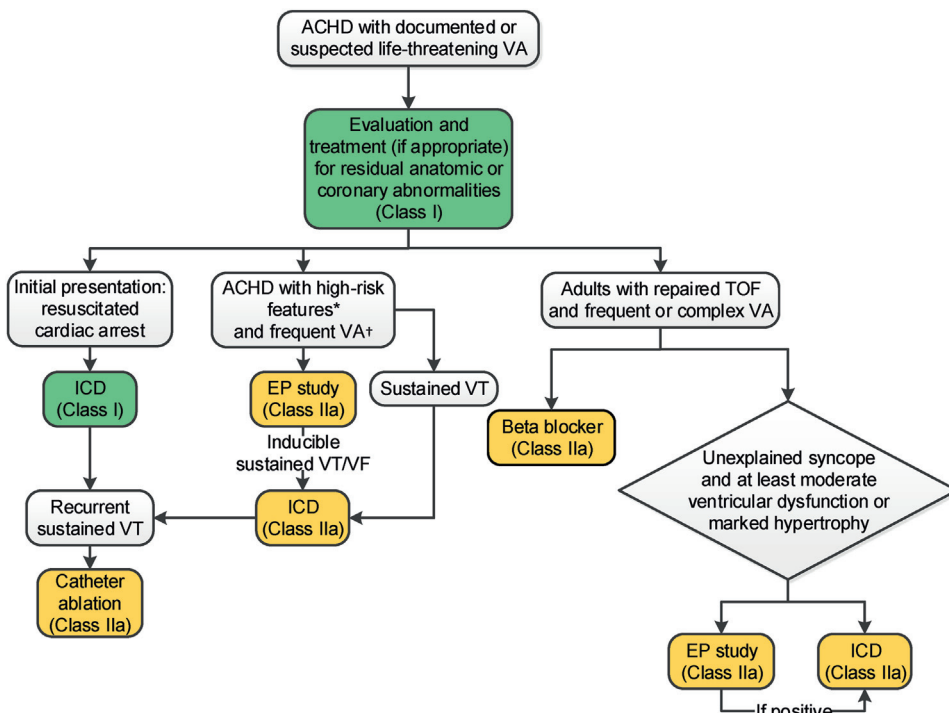
ASD indicates atrial septal defect; cc-TGA, congenitally corrected transposition of the great arteries; LV, left ventricular; LVEF, left ventricular ejection fraction; PR, pulmonary regurgitation; PS, pulmonary stenosis; RV, right ventricular; RVOT, right ventricular outflow tract; SCD, sudden cardiac death; VA, ventricular arrhythmia; VSD, ventricular septal defect; and VT, ventricular tachycardia.

as it has limited effectiveness and carries risk of ventricular proarrhythmia (S10.8-38). Pulmonary valve replacement in patients with TOF may result in improved hemodynamics and functional status, but it may not eliminate the risk of VT (S10.8-3,S10.8-12); postoperative reassessment for the need for an ICD is performed after the early recovery period.

3. Correction of residual hemodynamic/structural abnormalities contributing to VT may improve ventricular function and reduce symptoms, but it may inadequately prevent the risk of subsequent VT or SCA. The use of ICDs in adult congenital heart disease

patients for secondary prevention accounts for approximately 50% of implantations presently, at a mean age of 36 to 41 years (S10.8-13–S10.8-17). Patients with adult congenital heart disease experience appropriate shock rates of 3% to 6% per year, with equivalent or slightly increased frequency of appropriate shocks for secondary prevention indications (S10.8-14,S10.8-15,S10.8-17). Patients with adult congenital heart disease experience a higher rate of complications and inappropriate shocks compared with other adult populations (S10.8-13–S10.8-17,S10.8-39).

**FIGURE 16** Prevention of SCD in Patients With Adult Congenital Heart Disease



Colors correspond to Class of Recommendation in Table 1. See Section 10.8 for discussion. \*High-risk features: prior palliative systemic to pulmonary shunts, unexplained syncope, frequent PVC, atrial tachycardia, QRS duration  $\geq 180$  ms, decreased LVEF or diastolic dysfunction, dilated right ventricle, severe pulmonary regurgitation or stenosis, or elevated levels of BNP. †Frequent VA refers to frequent PVCs and/or nonsustained VT. AChD indicates adult congenital heart disease; BNP, B-type natriuretic peptide; EP, electrophysiological; ICD, implantable cardioverter-defibrillator; LVEF, left ventricular ejection fraction; PVC, premature ventricular complexes; SCD, sudden cardiac death; TOF, tetralogy of Fallot; VA, ventricular arrhythmia; and VT, ventricular tachycardia.

4. Challenges of ICD implantation in patients with adult congenital heart disease may include anatomic complexity, intracardiac shunts, and limited vascular access to the ventricle. Patients with adult congenital heart disease receiving an ICD have an increased rate of complications of 26% to 45%, as well as inappropriate shocks in 15% to 25% of patients (S10.8-13–S10.8-16,S10.8-40). Limited studies on the use of subcutaneous implantable cardioverter-defibrillator implantation, particularly in patients with single ventricle anatomy (S10.8-41), report improved success by using right in addition to left parasternal lead positioning for screening (S10.8-42). Patients with a single ventricle or a systemic right ventricle may not tolerate defibrillation threshold testing, resulting in multiorgan system failure. Patients with complex anatomy, such as older patients with univentricular physiology, or patients with significantly reduced ventricular function, marked hypertrophy, or multiple prior surgeries, may benefit from earlier consideration of heart transplantation before renal or liver dysfunction progresses.
5. Patients with repaired TOF who are at an increased risk of sustained VT include those with prior palliative systemic to pulmonary shunts, unexplained syncope, frequent PVCs, atrial tachycardia, QRS duration  $\geq 180$  ms, decreased LVEF or diastolic dysfunction, dilated right ventricle, severe pulmonary regurgitation or stenosis, or elevated levels of BNP. Patients with TOF physiology and suboptimal hemodynamic status are more likely to have inducible sustained VT (S10.8-18,S10.8-19,S10.8-33,S10.8-35), and inducible sustained VT correlated with an increased risk of SCA in a multicenter cohort study (S10.8-19). Evaluation of hemodynamics for residual abnormalities is important, with catheter or surgical treatment of important lesions prior to consideration of ICD implantation.
6. In a multicenter cohort, inducible sustained VT in patients with TOF was an independent risk factor for subsequent clinical VT or SCD (S10.8-19); patients in that early study had cardiomegaly and prior palliative shunts. Patients with repaired TOF account for approximately 50% of ICD implantations in adult

congenital heart disease ([S10.8-13–S10.8-16,S10.8-40](#)). Appropriate ICD shocks occur in up to 7.7% per year of patients with TOF receiving the ICD for primary prevention, compared with 9.8% per year in patients with a secondary prevention ICD ([S10.8-20](#)). In another study including patients with TOF as well as other lesions, inducible sustained VT did not correlate with subsequent appropriate ICD shocks ([S10.8-14](#)). Because of the high incidence of inappropriate shocks in 20% to 30% and complications in at least 30% of patients with adult congenital heart disease ([S10.8-14–S10.8-17,S10.8-39,S10.8-40,S10.8-43](#)), in addition to financial and psychological burdens, shared decision-making regarding primary prevention ICDs is essential.

7. In patients with recurrent sustained monomorphic VT, catheter ablation of VT can be effective ([S10.8-21–S10.8-25](#)). Hemodynamic repair, at the time that an arrhythmia is being ablated surgically, should be considered. For patients with complex adult congenital heart disease, care should be provided at experienced centers. After successful catheter ablation of VT, implantation of an ICD for those who do not have an ICD is an individualized decision based on overall functional and physiological status and shared decision making. Careful monitoring during follow-up for recurrent arrhythmias is essential.
8. The highest risk of SCD associated with repaired congenital heart disease reported from large contemporaneous cohorts is in patients with transposition of the great arteries with atrial baffle repair, Ebstein anomaly of the tricuspid valve, aortic stenosis, and univentricular physiology ([S10.8-44–S10.8-47](#)). Patients with Senning or Mustard atrial baffle repairs are at an increased risk for SCA, particularly during exertion ([S10.8-48](#)). The atrial baffle is noncompliant restricting ability to augment volume and may be associated with pulmonary vein stenosis and increased end-diastolic pressures. RV ischemia and infarction occur, with perfusion defects identified by myocardial perfusion studies in >40% of patients in this population ([S10.8-49,S10.8-50](#)). Risk factors for cardiac arrest in patients with transposition and atrial baffle repairs include prior ventricular septal defect closure, symptoms of HF, atrial arrhythmia, RVEF <30% to 35%, and QRS duration ≥140 ms ([S10.8-48, S10.8-51](#)). In the single multicenter study assessing outcomes after implantation of an ICD in patients with prior atrial baffle repair of transposition of the great arteries, the lack of beta blockers was associated with a high risk of appropriate ICD therapy ([S10.8-26](#)). Atrial arrhythmias frequently precede VT in transposition patients, and treatments for atrial tachycardia including catheter ablation, antitachycardia

pacing algorithms, and beta blockers are important to reduce ICD shocks ([S10.8-26,S10.8-52,S10.8-53](#)).

9. The risk of SCD is increased among patients with adult congenital heart disease compared with the general population, with the median age at death ranging from 30 to 49 years of age ([S10.8-27,S10.8-44,S10.8-47, S10.8-54,S10.8-55](#)). The risk of SCD is highest among patients with moderate or severe complexity congenital heart disease, and accounts for approximately 25% of cardiac causes of death ([S10.8-5, S10.8-27,S10.8-28,S10.8-44–S10.8-46,S10.8-55,S10.8-56](#)). Patients with septal defects and a positive family history of septal defects, cardiomyopathy, or bundle-branch block/conduction defects may have the gene mutation *NKX2.5*, which portends an increased risk of early SCD; genetic testing and early consideration of ICD implantation if positive is warranted ([S10.8-57–S10.8-59](#)). Patients with repaired complex forms of congenital heart disease have undergone multiple intracardiac surgeries in the first few decades of life with resultant hypertrophy and risk for sub-endocardial ischemia as well as scar formation contributing to VT/VF. Risk factors for SCD include increasing complexity of heart disease, VA, SVT, progressive increase in QRS duration, systemic ventricular dysfunction, and subpulmonary ventricular dysfunction ([S10.8-1,S10.8-5,S10.8-6,S10.8-14,S10.8-28, S10.8-29,S10.8-36,S10.8-45–S10.8-47,S10.8-55](#)). Extrapolation of data regarding specific measures of ventricular function warranting implantation of primary prevention ICDs from adult patients with NICM is unrealistic. The development of unexplained syncope in patients with moderate or severe complexity adult congenital heart disease may be a harbinger of risk for SCD; electrophysiological study with consideration for an ICD as primary prevention can be beneficial.
10. ICDs implanted in patients with adult congenital heart disease, who are in their 40s and 50s, for primary prevention indications now account for >40% to 67% of implanted devices in patients with adult congenital heart disease ([S10.8-13,S10.8-15,S10.8-16,S10.8-41](#)). In these patients, appropriate shocks are delivered in 14% to 22% of patients in the first 3 to 5 years of follow-up ([S10.8-13,S10.8-15,S10.8-16](#)). In patients with congenital heart disease and severely depressed ventricular function, or single ventricle anatomy, defibrillation threshold testing may pose excessively high risk. In patients without vascular access or prior Fontan repairs, the risk of reoperation with sternotomy for epicardial ICD implantation may outweigh the potential benefits, and consideration for transplant evaluation may be preferable. Subcutaneous implantable cardioverter-defibrillator implantation may be an appropriate option for some patients ([S10.8-42,S10.8-53](#)).

11. Adult patients with complex adult congenital heart disease typically have hypertrophy and ventricular dysfunction of varying degrees, increasing their risk for worsening ventricular function with antiarrhythmic medications. In the only large study of antiarrhythmic medications for congenital heart disease, the use of flecainide was associated with proarrhythmia in 5.8% of patients and SCA in 3.9% of patients (S10.8-30). The use of amiodarone is generally reserved for refractory symptomatic VA or asymptomatic VA that can

aggravate ventricular dysfunction, due to the high risk of adverse effects including thyroid dysfunction, particularly among females and patients with uni-ventricular physiology (S10.8-31,S10.8-32).

## 11. DEFIBRILLATORS OTHER THAN TRANSVENOUS ICDs

### 11.1. Subcutaneous Implantable Cardioverter-Defibrillator

#### Recommendations for Subcutaneous Implantable Cardioverter-Defibrillator

References that support the recommendations are summarized in Online Data Supplement 55.

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In patients who meet criteria for an ICD who have inadequate vascular access or are at high risk for infection, and in whom pacing for bradycardia or VT termination or as part of CRT is neither needed nor anticipated, a subcutaneous implantable cardioverter-defibrillator is recommended (S11.1-1–S11.1-5).
IIa	B-NR	2. In patients who meet indication for an ICD, implantation of a subcutaneous implantable cardioverter-defibrillator is reasonable if pacing for bradycardia or VT termination or as part of CRT is neither needed nor anticipated (S11.1-1–S11.1-4).
III: Harm	B-NR	3. In patients with an indication for bradycardia pacing or CRT, or for whom antitachycardia pacing for VT termination is required, a subcutaneous implantable cardioverter-defibrillator should not be implanted (S11.1-1–S11.1-4,S11.1-6–S11.1-8).

#### Synopsis

In patients being considered for a subcutaneous implantable cardioverter-defibrillator, a preimplant ECG to establish QRS-T wave morphology is needed to reduce the risk of under sensing of VT/VF and the risk of inappropriate shocks (S11.1-9–S11.1-11). The subcutaneous implantable cardioverter-defibrillator is implanted using primarily anatomical landmarks, thereby minimizing the need for fluoroscopy. The subcutaneous implantable cardioverter-defibrillator consists of a pulse generator that is placed at the midaxillary line between the fifth and sixth intercostal spaces and a lead with 2 sensing electrodes and a shocking coil, positioned subcutaneously adjacent to the sternum. As with the transvenous ICD, the pulse generator housing serves as an electrode for defibrillation but, in addition, it can also serve as an optional electrode for sensing. The subcutaneous implantable cardioverter-defibrillator cannot achieve adequate arrhythmia sensing for all patients, and electrocardiographic screening to assess sensing is required prior to implantation (S11.1-10, S11.1-11). Some advocate exercise testing after device implantation to ensure proper sensing with exercise.

Both transvenous and subcutaneous implantable cardioverter-defibrillators have SVT-VT discriminators that can be programmed to facilitate discrimination of SVT from VT; however, these discriminators do not always work. If sustained VT is confirmed, therapy to terminate the

arrhythmia is delivered. All ICDs provide shocks to terminate VT or VF, but shocks in an awake patient are painful and associated with decreased QoL. Transvenous ICDs are capable of bradycardia pacing as well as antitachycardia pacing that can terminate many VTs painlessly. Subcutaneous implantable cardioverter-defibrillators provide limited postshock bradycardia pacing but do not provide either bradycardia or antitachycardia pacing.

The subcutaneous implantable cardioverter-defibrillator recommendations supplant, but do not nullify, the need for waiting periods and other requirements to be satisfied for ICD/CRT implantation specified in other parts of this document.

#### Recommendation-Specific Supportive Text

1. The subcutaneous implantable cardioverter-defibrillator was designed to avoid the need for venous access and some of the complications of inserting transvenous lead(s) (S11.1-1–S11.1-4) that include pneumothorax, hemothorax, and cardiac tamponade (S11.1-12). Difficulties in achieving venous access can prolong the implantation procedure and occasionally result in failed ICD implantation. These difficulties are more likely to be encountered in patients with limited venous access such as patients with ESRD. In a study of 27 patients with ESRD, the subcutaneous implantable cardioverter-defibrillator was not associated with an

increased risk of procedural complications or inappropriate shocks ([S11.1-5](#)). The risk of infection appears to be lower with subcutaneous implantable cardioverter-defibrillators than with transvenous ICDs ([S11.1-1–S11.1-4](#)). Therefore, a subcutaneous implantable cardioverter-defibrillator may be preferred in patients who are at high risk of infection, such as those with a prior device infection, ESRD, diabetes mellitus, or who are chronically immunosuppressed.

2. Nonrandomized studies show that the subcutaneous implantable cardioverter-defibrillator reliably detects and converts VF during defibrillation threshold testing and successfully terminates spontaneous sustained VT that occurs during follow-up ([S11.1-1,S11.1-13](#)). In 1 study of 314 patients, the 180-day complication-free rate was 99%, and the success of VF termination with first shock was >90% ([S11.1-2](#)). All spontaneous episodes of VT/VF recorded in 21 patients (6.7%) were successfully converted, and there were no lead failures, endocarditis or bacteremia, tamponade, cardiac perforation, pneumothorax, or hemothorax associated with the subcutaneous implantable cardioverter-defibrillator ([S11.1-2](#)). In 472 patients enrolled in the EFFORTLESS (Evaluation of Factors Impacting Clinical Outcome and Cost Effectiveness of the S-ICD) registry ([S11.1-3](#)), the complication-free rate was 94%, at 360 days. First shock conversion efficacy was 88% with 100% overall successful clinical conversion after a maximum of 5 shocks. In 882 patients enrolled in

investigational device exemption trials and the EFFORTLESS registry ([S11.1-4](#)), 111 spontaneous VT/VF events were treated in 59 patients; 90.1% were terminated with 1 shock, and 98.2% were terminated within the 5 available shocks. The estimated 3-year inappropriate shock rate was 13.1% most due to oversensing of cardiac signals, and mortality was 4.7%. Device-related complications occurred in 11.1% of patients. An ongoing trial will compare the effect of the subcutaneous implantable cardioverter-defibrillator with that of the transvenous ICD on the outcomes of inappropriate shocks, complications, shock efficacy, and mortality ([S11.1-13](#)).

3. The subcutaneous implantable cardioverter-defibrillator is incapable of bradycardia pacing, biventricular pacing, or antitachycardia pacing. Therefore, patients who need any of these types of pacing from an ICD should not be offered a subcutaneous implantable cardioverter-defibrillator ([S11.1-6](#)). Some clinical scenarios may come up in which a transvenous pacemaker for bradycardia pacing in a patient with a subcutaneous implantable cardioverter-defibrillator which is needed; this can be performed as long as the pacing is not unipolar. Leadless pacing devices for patients who require bradycardia pacing will be evaluated with the subcutaneous implantable cardioverter-defibrillator in the near future.

## 11.2. Wearable Cardioverter-Defibrillator

### Recommendations for Wearable Cardioverter-Defibrillator

References that support the recommendations are summarized in [Online Data Supplement 56](#).

COR	LOE	RECOMMENDATIONS
IIa	B-NR	<p>1. In patients with an ICD and a history of SCA or sustained VA in whom removal of the ICD is required (as with infection), the wearable cardioverter-defibrillator is reasonable for the prevention of SCD (<a href="#">S11.2-1–S11.2-4</a>).</p>
IIb	B-NR	<p>2. In patients at an increased risk of SCD but who are not ineligible for an ICD, such as awaiting cardiac transplant, having an LVEF of 35% or less and are within 40 days from an MI, or have newly diagnosed NICM, revascularization within the past 90 days, myocarditis or secondary cardiomyopathy or a systemic infection, the wearable cardioverter-defibrillator may be reasonable (<a href="#">S11.2-1–S11.2-5</a>).</p>

### Synopsis

The wearable cardioverter-defibrillator is a vestlike device worn under the clothing that continuously monitors the heart rhythm and automatically delivers an electric shock when VF or VT is detected. This device is intended to be worn continuously, 24 hours per day, except when the wearer is bathing or showering. The wearable cardioverter-defibrillator has been approved in

the United States by the U.S. Food and Drug Administration for patients who are “at risk for SCA and are not candidates for or refuse an implantable defibrillator” ([S11.2-6](#)). A science advisory from the AHA summarizes the data and recommendations for the use of the wearable cardioverter-defibrillator ([S11.2-4](#)). Effectiveness of the wearable cardioverter-defibrillator in recognition and defibrillation of VF has been demonstrated in a number of

studies, although no RCTs support the use of the wearable cardioverter-defibrillator. Among 3569 patients who received the device for various reasons, for at least 1 day in the U.S. manufacturer registry, there were 80 VT/VF events in 59 patients, with a frequency of 1.7% per patient-year. First shock efficacy was 99%, with post-shock survival of 90%. Overall, 2% of the patients received an inappropriate shock (S11.2-1).

#### Recommendation-Specific Supportive Text

1. Removal of an ICD for a period of time, most commonly due to infection, exposes the patient to risk of untreated VT/SCD unless monitoring and access to emergency external defibrillation is maintained. In 1 series of 354 patients who received the wearable cardioverter-defibrillator, the indication was infection in 10% (S11.2-3). For patients with a history of SCA or sustained VA, the wearable cardioverter-defibrillator may allow the patient to be discharged from the hospital with protection from VT/SCA until the clinical situation allows reimplantation of an ICD.
2. The patients listed in this recommendation are represented in clinical series and registries that demonstrate the safety and effectiveness of the wearable cardioverter-defibrillator. Patients with recent MI, newly diagnosed NICM, recent revascularization, myocarditis, and secondary cardiomyopathy are at increased risk of VT/SCA. However, the wearable cardioverter-defibrillator is of unproven benefit in these settings, in part because the clinical situation may improve with therapy and time. In patients awaiting transplant, even with anticipated survival <1 year without transplant, and depending on clinical factors such as use of intravenous inotropes and ambient VA, a wearable cardioverter-defibrillator may be an alternative to an ICD.

#### 11.3. Automated External Defibrillator

External defibrillation can save lives when used within minutes of the onset of VF. The AED is an efficient

method of delivering defibrillation to persons experiencing out-of-hospital cardiac arrest, and its use by first responders is safe and effective (S11.3-1–S11.3-3). Federal efforts have been effective in placing AEDs in airports/airplanes and federal buildings, while varying efforts at the state and community levels have been effective in placing AEDs in many, but not all, schools, sporting events, high-density residential sites, and airports as well as in police and fire department vehicles (S11.3-4–S11.3-7). Resuscitation protocols with or without AED placement are required in most states for fitness clubs, although alternate indoor exercise facilities may have higher rates of arrest and provide for increased survival over other indoor public sites (S11.3-8). In a study population of 21 million, survival to hospital discharge was nearly twice as high when an AED was applied for out-of-hospital cardiac arrest (S11.3-9). Expanded and coordinated placement of AEDs in the community, including in high-risk geographic locations such as schools and organized sports arenas, can substantially increase the proportion of patients with cardiac out-of-hospital cardiac arrest who receive AED therapy (S11.3-10). The U.S. Food and Drug Administration has approved over-the-counter sales of AEDs. Approximately 70% of SCAs occur in the home, and the rate of survival to hospital discharge after AED placement by emergency medical services is significantly lower for arrest at home (12%) versus public settings (34%) (S11.3-11). However, in an RCT of AEDs, home AED placement did not improve the survival of patients recovering from an anterior MI (S11.3-12). Appropriate device location to reduce time delay after onset of SCA is critical. In addition to prevention, critical components of survival from SCA include immediate recognition and activation of the emergency response system, early high-quality CPR, and rapid defibrillation for shockable rhythms (S11.3-13).

## 12. SPECIAL CONSIDERATIONS FOR CATHETER ABLATION

#### Recommendations for Catheter Ablation

References that support the recommendations are summarized in [Online Data Supplement 57](#).

COR	LOE	RECOMMENDATIONS
I	C-LD	1. In patients with bundle-branch reentrant VT, catheter ablation is useful for reducing the risk of recurrent VT and ICD shocks (S12-1–S12-3).
IIa	B-NR	2. In patients with structural heart disease who have failed endocardial catheter ablation, epicardial catheter ablation can be useful for reducing the risk of recurrent monomorphic VT (S12-4–S12-6).

## Synopsis

Bundle-branch reentrant VT is due to reentry involving the bundle branches. Catheter ablation is the preferred therapy for this VT, which is encountered in <10% of patients with recurrent sustained monomorphic VT and structural heart disease (see [Section 7.2.3](#)).

### Recommendation-Specific Supportive Text

1. Bundle-branch reentrant VT can occur in any form of heart disease associated with slow infra-Hisian conduction. The most common mechanism involves antegrade conduction over the right bundle branch and retrograde conduction over the left bundle branch, thereby producing left bundle-branch block QRS morphology during VT, which is often rapid and poorly tolerated. Catheter ablation of the right or left bundle branch interrupts the circuit and is usually curative ([S12-1–S12-3](#)). After ablation, severely impaired atrioventricular conduction can be present, requiring permanent pacing, which can have hemodynamic consequences ([S12-4,S12-6](#)). Many patients have other inducible scar related VTs or meet eligibility for an ICD due to severity of associated heart disease.
2. Endocardial catheter ablation failure can be due to location of the arrhythmia substrate in the mid-myocardium or epicardium, and this is more likely in patients with nonischemic rather than ischemic car-

diomyopathy, and in arrhythmogenic right ventricular cardiomyopathy ([S12-7–S12-9](#)). In the HELP-VT trial ([S12-4](#)), epicardial ablation was required in 30% of patients with VT related to NICM compared with 1.2% of patients with ischemic cardiomyopathy. A wide QRS with marked slurring of the initial portion of the QRS and a QS complex in the lateral or inferior leads during VT suggests an epicardial circuit in NICM, but the ECG does not reliably predict epicardial VT locations in patients with prior MI. Preprocedural cardiac MRI and intraprocedural electroanatomic mapping are useful tools to guide the localization of epicardial scar that may be the source of reentrant VT ([S12-8,S12-10](#)). Pericardial adhesions prevent percutaneous access in some patients, notably many with prior cardiac surgery. Percutaneous pericardial access for mapping and ablation is associated with a serious complication rate of approximately 5% and tamponade from RV puncture or laceration that can require emergent surgery or be fatal, coronary artery injury and phrenic nerve injury can occur ([S12-11,S12-12](#)). Reported experience is from tertiary referral centers.

## 13. POSTMORTEM EVALUATION OF SCD

### Recommendations for Postmortem Evaluation of SCD

References that support the recommendations are summarized in [Online Data Supplement 58](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	1. In victims of SCD without obvious causes, a standardized cardiac-specific autopsy is recommended ( <a href="#">S13-1,S13-2</a> ).
I	B-NR	2. In first-degree relatives of SCD victims who were 40 years of age or younger, cardiac evaluation is recommended, with genetic counseling and genetic testing performed as indicated by clinical findings ( <a href="#">S13-3</a> ).
IIa	B-NR	3. In victims of SCD with an autopsy that implicates a potentially heritable cardiomyopathy or absence of structural disease, suggesting a potential cardiac channelopathy, postmortem genetic testing is reasonable ( <a href="#">S13-4–S13-7</a> ).
IIa	C-LD	4. In victims of SCD with a previously identified phenotype for a genetic arrhythmia-associated disorder, but without genotyping prior to death, postmortem genetic testing can be useful for the purpose of family risk profiling ( <a href="#">S13-8</a> ).

### Recommendation-Specific Supportive Text

1. A comprehensive postmortem protocol has been recommended for the routine evaluation of subjects (typically <40 years of age) who die suddenly without a prior diagnosis of a condition and circumstances of death that could be reasonably implicated in the cause of unexpected SCD ([S13-1](#)). One study documented the

added value of postmortem examination at a specialized cardiac pathology center ([S13-2](#)), with particular value for clarifying an apparent overdiagnosis of cardiomyopathy by nonspecialized centers. Pathological findings limited to the specialized conduction system were demonstrated in 22% of cases ([S13-9](#)). A misdiagnosis of cardiomyopathy was reported in 37% of

referred cases that were ultimately determined to be structurally normal. The etiologic data for specialized cardiac evaluation are not generalizable to the overall population because of skewing of age at the time of SCD. In another study of SCD patients at ages ranging from <1 year to >80 years (mean, 38.2 years; median, 38 years), the peak incidence of SCD occurred between the ages of 31 and 60 years, with a 5- to 7-fold excess of males/females in that age range (S13-10). For the overall group, 42% of SCD were due to ischemic heart disease, 12% viral myocarditis, and 5% cardiomyopathy, with 15% being unexplained by autopsy. For the subgroup <35 years of age, 13.5% were attributed to ischemic heart disease and 24.9% were unexplained. In the subgroup >55 years of age, only 0.8% were unexplained. In patients who die suddenly despite an ICD, interrogation of the ICD is important to confirm proper device functioning and can provide information on the mechanism of death.

- Comprehensive cardiac screening including 12-lead ECG, possible signal averaged ECG, echocardiogram, and ambulatory rhythm monitoring or exercise testing of first-degree relatives of decedents with sudden unexpected death may identify a probable heritable cardiac cause of death in up to 30% of cases (S13-11–S13-13). Genetic testing should be targeted based on the results of initial evaluation (S13-3). Genetic testing in selected first-degree relatives may result in identification of inherited conditions including long QT syndrome, catecholaminergic polymorphic ventricular tachycardia,

Brugada syndrome, arrhythmogenic right ventricular cardiomyopathy, and HCM in 4% to 30% of families (S13-11,S13-12,S13-14).

- For the purpose of family risk profiling, it is important to use the disease-specific genetic test panel that corresponds to the autopsy findings. Risk profiling of family members of an SCD victim suspected of having an inherited cardiomyopathy at autopsy is important. Although phenotyping of surviving family members is crucial, genotyping of the SCD proband provides a mechanism for efficient follow-up evaluation of those relatives with the disease-causing mutation found in the proband. To be able to harvest quality DNA for such testing, medical examiners, hospital pathologists, and private pathologists need standards for harvesting and storing samples for later genetic testing. Family members of SCD probands who died suddenly (first cardiac event, death from natural causes, last seen alive and well within 12 hours), with autopsy findings showing structural abnormalities of uncertain significance (e.g., ventricular hypertrophy, myocardial fibrosis, or minor ischemic heart disease [n=41]) had a 51% prevalence of genetic variants associated with sudden arrhythmic deaths, compared with 47% among a comparison group in which proband autopsies were completely negative (S13-15).
- Identification of the genotype can facilitate family screening (S13-16).

#### 14. TERMINAL CARE

##### Recommendations for Terminal Care

References that support the recommendations are summarized in Online Data Supplement 59.

COR	LOE	RECOMMENDATIONS
I	C-EO	<ol style="list-style-type: none"><li>At the time of ICD implantation or replacement, and during advance care planning, patients should be informed that their ICD shock therapy can be deactivated at any time if it is consistent with their goals and preferences.</li></ol>
I	C-EO	<ol style="list-style-type: none"><li>In patients with refractory HF symptoms, refractory sustained VA, or nearing the end of life from other illness, clinicians should discuss ICD shock deactivation and consider the patients' goals and preferences.</li></ol>

##### Synopsis

A particularly challenging area of medicine is recognizing when life-prolonging therapies may become burdensome or even harmful. This is particularly true near the end of life for patients with ICDs in whom once life-prolonging shocks may only cause unnecessary morbidity and distress to both patients and loved ones.

##### Recommendation-Specific Supportive Text

- Current evidence suggests that many patients are unaware of the possibility that their ICD can be

deactivated without surgery (S14-1–S14-3). During decision-making, clinicians do not routinely inform patients about ICD deactivation (S14-4). Clinicians even disagree on whether discussions of deactivation should occur when patients are making a decision about an ICD-related procedure (S14-5). As a result, patients often do not include wishes about deactivation in advance care planning documents (S14-6). Consequently, surrogates usually make decisions about ICD deactivation without any prior discussions with the

patient (S14-6). In hypothetical scenarios, patients with ICDs were able to identify scenarios in which they might choose to deactivate their ICD (S14-1,S14-7). This discussion can occur at any time, but it is particularly important to have it at the time of initial ICD implantation, at the time of reimplantation, and during preparation of advance care plans.

- When ICDs are not deactivated at the end of life, patients and families suffer unnecessarily. Families have had unpleasant experiences of watching their loved one die while getting shocked repeatedly by an ICD (S14-8). In 1 survey of hospice staff, half of those surveyed noted that a deceased patient had been shocked by an ICD during the year prior to the survey (S14-9). This is unnecessary and easily preventable by having caring, patient-centered discussions with patients and

their loved ones. In general, patients want their clinicians to initiate these discussions (S14-2,S14-10), so this recommendation is carefully worded to put the responsibility of initiating the discussion on the clinician. Ethically, patients and surrogates are free to choose to deactivate antitachycardia function (S14-11–S14-13). Most patients only elect deactivation of the anti-tachycardia functions while leaving the pacing function on. Even at the end of life, pacing (either for bradycardia or for resynchronization therapy) may be an important aspect of the patient's QoL and may facilitate more alert and meaningful personal interactions. These differences are easily misunderstood, so they need careful explanation.

## 15. SHARED DECISION-MAKING

### Recommendations for Shared Decision-Making

References that support the recommendations are summarized in [Online Data Supplement 60](#).

COR	LOE	RECOMMENDATIONS
I	B-NR	<ol style="list-style-type: none"> <li>In patients with VA or at increased risk for SCD, clinicians should adopt a shared decision-making approach in which treatment decisions are based not only on the best available evidence but also on the patients' health goals, preferences, and values (S15-1–S15-5).</li> </ol>
I	B-NR	<ol style="list-style-type: none"> <li>Patients considering implantation of a new ICD or replacement of an existing ICD for a low battery should be informed of their individual risk of SCD and nonsudden death from HF or noncardiac conditions and the effectiveness, safety, and potential complications of the ICD in light of their health goals, preferences and values (S15-1–S15-5).</li> </ol>

### Synopsis

During most of their lives, people prefer to do everything possible to prevent SCD and prolong life. However, many people may get to a point in their lives where SCD is not the worst outcome. Patients may report a desire to die in their sleep (S15-6). Decisions related to SCD can be quite emotional; according to the patient's wishes, shared decision regarding end-of-life therapy making may involve caregivers such as family members or friends.

### Recommendation-Specific Supportive Text

- Consideration of patient preferences is important for VA diagnosis and management decisions. Patient preferences for invasive therapies and acceptance of SCD risk vary and may evolve throughout the course of their illness. The writing committee endorses a shared decision-making approach as part of the general care for patients at risk for VA and SCD. A commonly accepted definition of the shared decision-making (S15-7) includes 4 components: 1) at least 2 participants, the clinician and patient, be involved; 2) both parties share

information; 3) both parties take steps to build a consensus about the preferred treatment; and 4) an agreement is reached on the treatment to implement. Sharing a decision does not mean giving a patient a list of risks and benefits and telling them to make a decision—a practice some authors have called “abandonment” (S15-8). Notably, a recommendation based on evidence or guidelines alone is not shared decision-making. Rather, a recommendation based both on the evidence as well as an understanding of the patients' health goals, preferences, and values is essential to achieving true shared decision-making. Also, the possibility of deactivation of an existing ICD should be discussed with patients who have terminal illnesses.

- ICDs prolong lives as highlighted in many places within this guideline. However, a patient with HF or advanced noncardiac illness may elect to forgo replacement of an ICD when faced with the prospect of continual decline in health and functional status from either progressive HF or some other competing morbidity.

Unfortunately, research suggests that patients are ill-informed when faced with understanding the risks, benefits, and downstream burdens of their ICDs. Patients with an ICD tend to overestimate the benefit of this therapy and underestimate its risks ([S15-1–S15-3](#)). Likewise, patients who decline an ICD also frequently underestimate their personal risk of VA and SCD ([S15-4,S15-5](#)). Studies of clinician decision-making demonstrate that clinicians often overestimate the benefits while down-playing the potential harms ([S15-3](#)).

In kind, ICD replacement is also an important point in time where patients and clinicians should discuss whether replacing an ICD is still consistent with the patients' goals. What made sense at 70 years of age may not make sense at 80 years of age. Patients may have had progressive disease or developed poor QoL. These factors can all change the risk/benefit ratio of the ICD and the patients' preferences.

## 16. COST AND VALUE CONSIDERATIONS

The key principles of value assessment as part of clinical practice guidelines have been discussed in detail ([S16-1](#)). Economic outcomes of clinical management strategies can be documented empirically using the same research designs as used in establishing clinical outcomes, including RCTs and observational comparisons. In addition, simulation models are often used to assess the value of management strategies, because the standard for cost-effectiveness studies is to compare life-time outcomes, and clinical studies usually have follow-up of a few years at most. Standards for economic modeling in health care have been published by an expert group ([S16-2](#)).

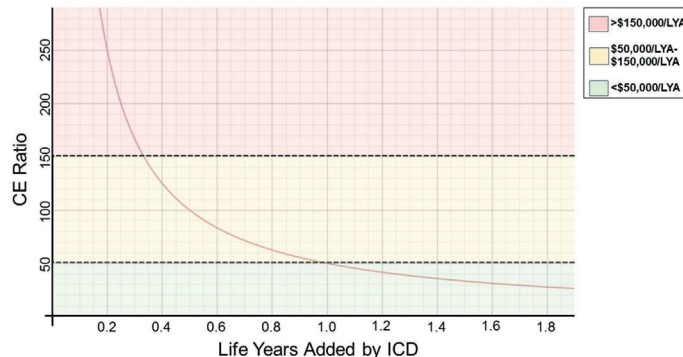
Economic assessments of alternative management strategies for VA and prevention of SCD have primarily evaluated ICDs, including several RCTs ([S16-3–S16-7](#)) and observational studies ([S16-8,S16-9](#)), and simulation models ([S16-10–S16-14](#)). In all studies, patients who received ICDs had higher long-term costs. The high initial cost of the ICD device and the implantation procedure leads to higher long-term costs, because there are few, if any, subsequent cost-savings from implanting an ICD. ICDs without resynchronization capability do not reduce hospital readmissions and may increase late costs due to device monitoring, complications, and replacement. However, the cost of the device and the procedure may change significantly over time.

The trial based assessments of the cost-effectiveness of the ICD are based on 3 to 6 years of follow-up, which is considerably shorter than the lifetime perspective that is standard in cost-effectiveness models. Because most of the incremental cost of the ICD is incurred immediately,

while most of the potential effectiveness (life-years of survival added by the ICD) is accrued over many years, estimates of ICD cost-effectiveness based on limited trial follow-up have a systematic bias toward showing lower value. Trial based economic studies that projected long-term ICD outcomes have consistently found more favorable cost-effectiveness ratios than estimates restricted to the duration of trial follow-up ([S16-4–S16-7](#)). A lifetime simulation model applied to each major trial of primary prevention ICDs also reported consistently more favorable estimates of cost-effectiveness than the estimates based on limited trial follow-up ([S16-11](#)). Because the framework proposed for assessing value in ACC/AHA clinical practice guidelines uses benchmarks based on lifetime estimates ([S16-1](#)), we have generally relied on the model-based estimates of ICD cost-effectiveness in applying value ratings to recommendations in this guideline.

The initial cost of an ICD device is similar regardless of the clinical indication, so variations in ICD cost-effectiveness are driven primarily by potential differences in clinical effectiveness in extending survival in different patient populations. The effect of the years of life added by an ICD on its incremental cost-effectiveness ratio is illustrated in [Figure 17](#): the cost-effectiveness ratio becomes rapidly unfavorable as the extension in survival time falls below 1 year, particularly below 0.5 year. This inverse relation strongly suggests that the value provided by an ICD will be highest when the risk of arrhythmic death due to VT/VF is relatively high and the risk of nonarrhythmic death (either cardiac or noncardiac) is relatively low, such that a meaningful increase in survival can be expected from the ICD. Thus, appropriate patient selection is fundamental to high value care in using the ICD to prevent SCD. It should also be recognized that cost-effectiveness is also influenced by the costs for the ICD and implantation procedure, which are likely to change significantly over time.

The empirical evidence suggests that ICDs are not effective for primary prevention of SCD when implanted early after CABG ([S16-15](#)) or an acute myocardial infarction ([S16-16,S16-17](#)). An analysis of individual patient level data from 3 secondary prevention trials ([S16-18](#)) showed a significant variation ( $p=0.011$ ) in the clinical effectiveness of ICDs between patients with an LVEF  $\leq 35\%$  (hazard ratio: 0.66) and an LVEF  $> 35\%$  (hazard ratio: 1.2). Some studies and simulation models suggest that ICDs might prolong life expectancy to a greater extent when used in higher-risk patients than in lower-risk patients ([S16-19](#)). In contrast, there is little evidence of variation in the effectiveness or cost-effectiveness of the ICD based on factors such as age or

**FIGURE 17 Incremental Cost-Effectiveness of ICD by Years of Life Added\* (Example)**

\*Figure based on formula: Incremental cost-effectiveness ratio = \$50,000/QALYs. CE indicated cost effectiveness, ICD, implantable cardioverter-defibrillator; LYA, life year added; and QALYs, quality-adjusted life-years

sex (S16-20). Most studies of ICD effectiveness and value have been performed on patients with reduced LV function due to prior MI or NICM. There are few data on the effectiveness or value of an ICD for other potential clinical indications, such as cardiac channelopathies or HCM, although studies have suggested that their potential cost effectiveness in such patients will depend on their underlying risk of SCD, with little evidence of value in low-risk patients (S16-14).

## 17. QUALITY OF LIFE

ICD implantation has not had a significant effect on QoL in the overall population of patients enrolled in RCTs (S17-1–S17-3). Several studies have, however, demonstrated that the subset of patients who receive inappropriate ICD shocks have worse QoL than patients who have an ICD but have not had inappropriate shocks (S17-2). Because an ICD is designed to prevent SCD rather than to reduce symptoms, it would not be expected to improve QoL or functional status directly, but may have indirect, negative effects in some patients due to device complications, or indirect, positive effects in some patients due to reassurance of having a protective device in place.

## 18. EVIDENCE GAPS AND FUTURE RESEARCH NEEDS

Despite the numerous advances in risk stratification for SCD and prevention and treatment of SCD and VA, many gaps in knowledge remain. These gaps include:

- Identification of patients who are most likely to benefit from an ICD among all ICD-eligible patients. The role of novel markers (including genetic and imaging markers) and combinations of markers should be studied.
- Characterizing the role of the ICD in patient subgroups not well-represented in the pivotal ICD trials. Such subgroups include patients ≥80 years of age and those with kidney disease, especially patients with ESRD on dialysis, or multiple comorbidities.
- Methods to identify and treat patients at high individual risk for SCD who are not identified by current ICD eligibility criteria, including those who are within 40 days of an MI.
- Defining the role of the ICD in patients with HCM, arrhythmogenic right ventricular cardiomyopathy, cardiac sarcoidosis, and inherited cardiac channelopathies in prospective studies (preferably RCT).
- Determining the best approach to patients due for elective ICD generator replacement due to battery depletion, but who may now be at low risk for SCA, such as if significant LVEF improvement has occurred.
- Obtaining more data on the efficacy and effectiveness of the subcutaneous implantable cardioverter-defibrillator, compared with transvenous ICDs and on the extent of testing required, and its use with other novel technologies, including leadless pacemakers.
- Conducting RCTs on catheter ablation of VT in ischemic heart disease and cardiomyopathies that evaluates procedural end points, mortality, arrhythmia suppression, QoL, and costs.

- Improving identification of individuals without significant ventricular dysfunction who are at risk of SCD.
- Identifying mechanisms and risk factors for SCD in patients with HFpEF.
- Improving emergency response to out-of-hospital cardiac arrest.
- Developing better methods for identifying and ablating the arrhythmia substrate in structural heart disease.
- Developing better risk stratification of diseases and syndromes associated with sudden death, including ischemic heart disease, NICM, adult congenital heart disease, and Brugada syndrome.
- Identifying what causes different types of long QT syndrome, catecholaminergic polymorphic ventricular tachycardia, Brugada syndrome, HCM, and arrhythmogenic right ventricular cardiomyopathy and advancing the genotype-phenotype relationships, genotype-dependent risk, and genotype-based tailoring of therapies for patients with inherited cardiomyopathies and inherited channelopathies.
- Defining the most appropriate and beneficial use of wearable cardioverter-defibrillators.
- Developing methods to identify and treat patients at high personal risk for SCD who are not identified by current ICD eligibility criteria.
- Defining the role of CMR in enhancing risk stratification for SCD.

Increasing research funding in this area, through existing and new mechanisms is critically important. Some have proposed research funding strategies that

would offer business incentives to the insurance industries, while providing support for unresolved research goals. Such approaches should be tested.

## PRESIDENTS AND STAFF

### **American College of Cardiology**

Mary Norine Walsh, MD, FACC, President  
Shalom Jacobovitz, Chief Executive Officer  
William J. Oetgen, MD, MBA, FACC, Executive Vice President, Science, Education, Quality, and Publishing  
Amelia Scholtz, PhD, Publications Manager, Science, Education, Quality, and Publishing

### **American College of Cardiology/American Heart Association**

Katherine A. Sheehan, PhD, Director, Guideline Strategy and Operations  
Abdul R. Abdullah, MD, Science and Medicine Advisor  
Sam Shahid, MBBS, MPH, Associate Science and Medicine Advisor

### **American Heart Association**

John J. Warner, MD, President  
Nancy Brown, Chief Executive Officer  
Rose Marie Robertson, MD, FAHA, Chief Science and Medicine Officer  
Gayle R. Whitman, PhD, RN, FAHA, FAAN, Senior Vice President, Office of Science Operations  
Prashant Nedungadi, PhD, Associate Science and Medicine Advisor, Office of Science Operations  
Jody Hundley, Production and Operations Manager, Scientific Publications, Office of Science Operations

## REFERENCES

## PREAMBLE

**P-1.** Committee on Standards for Developing Trustworthy Clinical Practice Guidelines, Institute of Medicine (U.S.). Clinical Practice Guidelines We Can Trust. Washington, DC: National Academies Press, 2011.

**P-2.** Committee on Standards for Systematic Reviews of Comparative Effectiveness Research, Institute of Medicine (U.S.). Finding What Works in Health Care: Standards for Systematic Reviews. Washington, DC: National Academies Press, 2011.

**P-3.** Anderson JL, Heidenreich PA, Barnett PG, et al. ACC/AHA statement on cost/value methodology in clinical practice guidelines and performance measures: a report of the American College of Cardiology/American Heart Association Task Force on Performance Measures and Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;63:2304–22.

**P-4.** ACCF/AHA Task Force on Practice Guidelines. Methodology Manual and Policies From the ACCF/AHA Task Force on Practice Guidelines. American College of Cardiology and American Heart Association, 2010. Available at: [http://assets.cardiosource.com/Methodology\\_Manual\\_for\\_ACC\\_AHA\\_Writing\\_Committees.pdf](http://assets.cardiosource.com/Methodology_Manual_for_ACC_AHA_Writing_Committees.pdf) and [http://professional.heart.org/idc/groups/ahamah-public/@wcm/@sop/documents/downloadable/ucm\\_319826.pdf](http://professional.heart.org/idc/groups/ahamah-public/@wcm/@sop/documents/downloadable/ucm_319826.pdf). Accessed October 1, 2017.

**P-5.** Arnett DK, Goodman RA, Halperin JL, et al. AHA/ACC/HHS strategies to enhance application of clinical practice guidelines in patients with cardiovascular disease and comorbid conditions: from the American Heart Association, American College of Cardiology, and U.S. Department of Health and Human Services. *J Am Coll Cardiol.* 2014;64:1851–6.

**P-6.** Halperin JL, Levine GN, Al-Khatib SM, et al. Further evolution of the ACC/AHA clinical practice guideline recommendation classification system: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Clinical Practice Guidelines. *J Am Coll Cardiol.* 2016;67:1572–4.

**P-7.** Jacobs AK, Anderson JL, Halperin JL. The evolution and future of ACC/AHA clinical practice guidelines: a 30-year journey: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;64:1373–84.

**P-8.** Jacobs AK, Kushner FG, Ettinger SM, et al. ACCF/AHA clinical practice guideline methodology summit report: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2013;61:213–65.

## 1. INTRODUCTION

**S1.4-1.** Kusumoto FM, Bailey KR, Chaouki AS, et al. Systematic review for the 2017 AHA/ACC/HRS guideline for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2018;72:1653–76.

**S1.4-2.** Anderson JL, Heidenreich PA, Barnett PG, et al. ACC/AHA statement on cost/value methodology in clinical practice guidelines and performance measures: a report of the American College of Cardiology/American Heart Association Task Force on Performance Measures and Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;63:2304–22.

**S1.4-3.** World Health Organization. CHOosing Interventions that are Cost Effective (WHO-CHOICE): cost-effectiveness thresholds. Available at: <http://www.who.int/choice/en/>. Accessed March 26, 2013.

**S1.4-4.** Zipes DP, Camm AJ, Borggrefe M, et al. ACC/AHA/ESC 2006 guidelines for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology/American Heart Association Task Force and the European Society of Cardiology Committee for Practice Guidelines (Writing Committee to Develop Guidelines for Management of Patients With Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death). Developed in collaboration with the European Heart Rhythm Association and the Heart Rhythm Society. *J Am Coll Cardiol.* 2006;48:e247–346.

**S1.4-5.** Epstein AE, DiMarco JP, Ellenbogen KA, et al. ACC/AHA/HRS 2008 guidelines for device-based therapy of cardiac rhythm abnormalities: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines (Writing Committee to Revise the ACC/AHA/NASPE 2002 Guideline Update for Implantation of Cardiac Pacemakers and Antiarrhythmia Devices). Developed in collaboration with the American Association for Thoracic Surgery and Society of Thoracic Surgeons. *J Am Coll Cardiol.* 2008;51:e1–62.

**S1.4-6.** Gersh BJ, Maron BJ, Bonow RO, et al. 2011 ACCF/AHA guideline for the diagnosis and treatment of hypertrophic cardiomyopathy: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Developed in collaboration with the American Association for Thoracic Surgery, American Society of Echocardiography, American Society of Nuclear Cardiology, Heart Failure Society of America, Heart Rhythm Society, Society for Cardiovascular Angiography and Interventions, and Society of Thoracic Surgeons. *J Am Coll Cardiol.* 2011;58:e212–60.

**S1.4-7.** Buxton AE, Calkins H, Callans DJ, et al. ACC/AHA/HRS 2006 key data elements and definitions for electrophysiological studies and procedures: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Data Standards (ACC/AHA/HRS Writing Committee to Develop Data Standards on Electrophysiology). *J Am Coll Cardiol.* 2006;48:2360–96.

**S1.4-8.** Wilkoff BL, Fauchier L, Stiles MK, et al. 2015 HRS/EHRA/APHRS/SOLAECE expert consensus statement on optimal implantable cardioverter-defibrillator programming and testing. *J Arrhythm.* 2016;32:1–28.

**S1.4-9.** Piccini JP Sr., Allen LA, Kudenchuk PJ, et al. Wearable cardioverter-defibrillator therapy for the prevention of sudden cardiac death: a science advisory from the American Heart Association. *Circulation.* 2016;133:1715–27.

**S1.4-10.** Shen WK, Sheldon RS, Benditt DG, et al. 2017 ACC/AHA/HRS guideline for the evaluation and

management of patients with syncope: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2017;70:e39–110.

**S1.4-11.** Yancy CW, Jessup M, Bozkurt B, et al. 2017 ACC/AHA/HFSA focused update of the 2013 ACCF/AHA guideline for the management of heart failure: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Failure Society of America. *J Am Coll Cardiol.* 2017;70:776–803.

**S1.4-12.** Yancy CW, Jessup M, Bozkurt B, et al. 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: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Failure Society of America. *J Am Coll Cardiol.* 2016;68:1476–88.

**S1.4-13.** Yancy CW, Jessup M, Bozkurt B, et al. 2013 ACCF/AHA guideline for the management of heart failure: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2013;62:e147–239.

**S1.4-14.** Nishimura RA, Otto CM, Bonow RO, et al. 2017 AHA/ACC focused update of the 2014 AHA/ACC guideline for the management of patients with valvular heart disease: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. *J Am Coll Cardiol.* 2017;70:252–89.

**S1.4-15.** Nishimura RA, Otto CM, Bonow RO, et al. 2014 AHA/ACC guideline for the management of patients with valvular heart disease: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;63:e57–185.

**S1.4-16.** Page RL, Jorgler JA, Caldwell MA, et al. 2015 ACC/AHA/HRS guideline for the management of adult patients with supraventricular tachycardia: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2016;67:e27–115.

**S1.4-17.** Priori SG, Blomstrom-Lundqvist C, Mazzanti A, et al. 2015 ESC guidelines for the management of patients with ventricular arrhythmias and the prevention of sudden cardiac death. The Task Force for the Management of Patients with Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death of the European Society of Cardiology. *G Ital Cardiol (Rome).* 2016;17:108–70.

**S1.4-18.** Perkins GD, Jacobs IG, Nadkarni VM, et al. Cardiac arrest and cardiopulmonary resuscitation outcome reports: update of the Utstein Resuscitation Registry Templates for Out-of-Hospital Cardiac Arrest: a statement for healthcare professionals from a task force of the International Liaison Committee on Resuscitation (American Heart Association, European Resuscitation Council, Australian and New Zealand Council on Resuscitation, Heart and Stroke Foundation of Canada, InterAmerican Heart Foundation, Resuscitation Council of Southern Africa,

Resuscitation Council of Asia); and the American Heart Association Emergency Cardiovascular Care Committee and the Council on Cardiopulmonary, Critical Care, Perioperative and Resuscitation. *Circulation.* 2015;132:1286–300.

**S1.4-19.** January CT, Wann LS, Alpert JS, et al. 2014 AHA/ACC/HRS guideline for the management of patients with atrial fibrillation: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2014;64:e1–76.

**S1.4-20.** Amsterdam EA, Wenger NK, Brindis RG, et al. 2014 AHA/ACC guideline for the management of patients with non-ST-elevation acute coronary syndromes: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;64:e139–228.

**S1.4-21.** Andrus B, Lacaille D. 2013 ACC/AHA guideline on the assessment of cardiovascular risk. *J Am Coll Cardiol.* 2014;63:2886.

**S1.4-22.** O’Gara PT, Kushner FG, Ascheim DD, et al. 2013 ACCF/AHA guideline for the management of ST-elevation myocardial infarction: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2013;61:e78–140.

**S1.4-23.** Steg PG, James SK, Atar D, et al. ESC guidelines for the management of acute myocardial infarction in patients presenting with ST-segment elevation. *Eur Heart J.* 2012;33:2569–619.

**S1.4-24.** Tracy CM, Epstein AE, Darbar D, et al. 2012 ACCF/AHA/HRS focused update of the 2008 guidelines for device-based therapy of cardiac rhythm abnormalities: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2012;60:1297–313.

**S1.4-25.** Hillis LD, Smith PK, Anderson JL, et al. 2011 ACCF/AHA guideline for coronary artery bypass graft surgery: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Developed in collaboration with the American Association for Thoracic Surgery, Society of Cardiovascular Anesthesiologists, and Society of Thoracic Surgeons. *J Am Coll Cardiol.* 2011;58:e123–210.

**S1.4-26.** Levine GN, Bates ER, Blankenship JC, et al. 2011 ACCF/AHA/SCAI guideline for percutaneous coronary intervention: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines and the Society for Cardiovascular Angiography and Interventions. *Catheter Cardiovasc Interv.* 2013;82:E266–355.

**S1.4-27.** Smith SC Jr., Benjamin EJ, Bonow RO, et al. AHA/ACCF secondary prevention and risk reduction therapy for patients with coronary and other atherosclerotic vascular disease: 2011 update: a guideline from the American Heart Association and American College of Cardiology Foundation. *J Am Coll Cardiol.* 2011;58:2432–46.

**S1.4-28.** Institute of Medicine. Committee on the Treatment of Cardiac Arrest: Current Status and Future Directions: Strategies to improve cardiac arrest survival: a time to act. Washington, DC: National Academic Press, 2015.

**S1.4-29.** Link MS, Myerburg RJ, Estes NA 3rd. Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities: Task Force 12: emergency action plans, resuscitation, cardiopulmonary resuscitation, and automated external defibrillators: a scientific statement from the American Heart Association and American College of Cardiology. *Circulation.* 2015;132:e334–8.

**S1.4-30.** Pedersen CT, Kay GN, Kalman J, et al. EHRA/HRS/APHRS expert consensus on ventricular arrhythmias. *Heart Rhythm.* 2014;11:e166–96.

**S1.4-31.** Khairy P, Van Hare GF, Balaji S, et al. PACES/HRS expert consensus statement on the recognition and management of arrhythmias in adult congenital heart disease: developed in partnership between the Pediatric and Congenital Electrophysiology Society (PACES) and the Heart Rhythm Society (HRS). *Heart Rhythm.* 2014;11:e102–65.

**S1.4-32.** Kusumoto FM, Calkins H, Boehmer J, et al. HRS/ACC/AHA expert consensus statement on the use of implantable cardioverter-defibrillator therapy in patients who are not included or not well represented in clinical trials. *J Am Coll Cardiol.* 2014;64:1143–77.

**S1.4-33.** Birnie DH, Sauer WH, Bogun F, et al. HRS expert consensus statement on the diagnosis and management of arrhythmias associated with cardiac sarcoidosis. *Heart Rhythm.* 2014;11:1305–23.

**S1.4-34.** Priori SG, Wilde AA, Horie M, et al. HRS/EHRA/APHRS expert consensus statement on the diagnosis and management of patients with inherited primary arrhythmia syndromes. *Heart Rhythm.* 2013;10:1932–63.

## 2. EPIDEMIOLOGY

**S2.2.2-1.** Myerburg RJ, Junnila MJ. Sudden cardiac death caused by coronary heart disease. *Circulation.* 2012;125:1043–52.

**S2.2.2-2.** Buxton AE, Calkins H, Callans DJ, et al. ACC/AHA/HRS 2006 key data elements and definitions for electrophysiological studies and procedures: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Data Standards. *J Am Coll Cardiol.* 2006;48:2360–96.

**S2.2.2-3.** Sapp JL, Wells GA, Parkash R, et al. Ventricular tachycardia ablation versus escalation of antiarrhythmic drugs. *N Engl J Med.* 2016;375:111–21.

**S2.2.2-4.** Priori SG, Wilde AA, Horie M, et al. HRS/EHRA/APHRS expert consensus statement on the diagnosis and management of patients with inherited primary arrhythmia syndromes. *Heart Rhythm.* 2013;10:1932–63.

**S2.2.2-5.** Hiss RG, Lamb LE. Electrocardiographic findings in 122,043 individuals. *Circulation.* 1962;25:947–61.

**S2.2.2-6.** Brodsky M, Wu D, Denes P, et al. Arrhythmias documented by 24 hour continuous electrocardiographic monitoring in 50 male medical students without apparent heart disease. *Am J Cardiol.* 1977;39:390–5.

**S2.2.2-7.** Massing MW, Simpson RJ Jr., Rautaharju PM, et al. Usefulness of ventricular premature complexes to predict coronary heart disease events and mortality (from the Atherosclerosis Risk In Communities cohort). *Am J Cardiol.* 2006;98:1609–12.

**S2.2.2-8.** Ofoma U, He F, Shaffer ML, et al. Premature cardiac contractions and risk of incident ischemic stroke. *J Am Heart Assoc.* 2012;1:e002519.

**S2.2.2-9.** Atakite F, Erquou S, Laukkonen J, et al. Meta-analysis of ventricular premature complexes and their relation to cardiac mortality in general populations. *Am J Cardiol.* 2013;112:1263–70.

**S2.2.2-10.** Lin CY, Chang SL, Lin YJ, et al. Long-term outcome of multiform premature ventricular complexes in structurally normal heart. *Int J Cardiol.* 2015;180:80–5.

**S2.2.2-11.** Lin CY, Chang SL, Chung FP, et al. Long-term outcome of non-sustained ventricular tachycardia in structurally normal hearts. *PloS One.* 2016;11:e0160181.

**S2.2.2-12.** Ruberman W, Weinblatt E, Goldberg JD, et al. Ventricular premature complexes and sudden death after myocardial infarction. *Circulation.* 1981;64:297–305.

**S2.2.2-13.** Ruberman W, Weinblatt E, Goldberg JD, et al. Ventricular premature beats and mortality after myocardial infarction. *N Engl J Med.* 1977;297:750–7.

**S2.2.2-14.** The Cardiac Arrhythmia Suppression Trial II Investigators. Effect of the antiarrhythmic agent moricizine on survival after myocardial infarction. The Cardiac Arrhythmia Suppression Trial II Investigators. *N Engl J Med.* 1992;327:227–33.

**S2.2.2-15.** Echt DS, Liebson PR, Mitchell LB, et al. Mortality and morbidity in patients receiving encainide, flecainide, or placebo. The Cardiac Arrhythmia Suppression Trial. *N Engl J Med.* 1991;324:781–8.

**S2.2.2-16.** Morganroth J, Goin JE. Quinidine-related mortality in the short-to-medium-term treatment of ventricular arrhythmias. A meta-analysis. *Circulation.* 1991;84:1977–83.

**S2.2.2-17.** Waldo AL, Camm AJ, deRuyter H, et al. Effect of d-sotalol on mortality in patients with left ventricular dysfunction after recent and remote myocardial infarction. The SWORD Investigators. *Survival With Oral d-Sotalol.* *Lancet.* 1996;348:7–12.

**S2.2.2-18.** Ling Z, Liu Z, Su L, et al. Radiofrequency ablation versus antiarrhythmic medication for treatment of ventricular premature beats from the right ventricular outflow tract: prospective randomized study. *Circ Arrhythm Electrophysiol.* 2014;7:237–43.

**S2.2.2-19.** Jouven X, Zureik M, Desnos M, et al. Long-term outcome in asymptomatic men with exercise-induced premature ventricular depolarizations. *N Engl J Med.* 2000;343:826–33.

**S2.2.2-20.** Frolkis JP, Pothier CE, Blackstone EH, et al. Frequent ventricular ectopy after exercise as a predictor of death. *N Engl J Med.* 2003;348:781–90.

**S2.2.2-21.** Biffi A, Pelliccia A, Verdile L, et al. Long-term clinical significance of frequent and complex ventricular tachyarrhythmias in trained athletes. *J Am Coll Cardiol.* 2002;40:446–52.

**S2.2.2-22.** Heidbüchel H, Hoogsteen J, Fagard R, et al. High prevalence of right ventricular involvement in endurance athletes with ventricular arrhythmias: role of an electrophysiologic study in risk stratification. *Eur Heart J.* 2003;24:1473–80.

**S2.2.2-23.** Kanei Y, Friedman M, Ogawa N, et al. Frequent premature ventricular complexes originating from the right ventricular outflow tract are associated with left ventricular dysfunction. *Ann Noninvasive Electrocardiol.* 2008;13:81–5.

**S2.2.2-24.** Lee GK, Klarich KW, Grogan M, et al. Premature ventricular contraction-induced cardiomyopathy: a treatable condition. *Circ Arrhythm Electrophysiol.* 2012;5:229–36.

**S2.2.2-25.** Viskin S, Rosso R, Rogowski O, et al. The "short-coupled" variant of right ventricular outflow ventricular tachycardia: a not-so-benign form of benign ventricular tachycardia? *J Cardiovasc Electrophysiol.* 2005;16:912–6.

**S2.2.2-26.** Noda T, Shimizu W, Taguchi A, et al. Malignant entity of idiopathic ventricular fibrillation and polymorphic ventricular tachycardia initiated by premature extrasystoles originating from the right ventricular outflow tract. *J Am Coll Cardiol.* 2005;46:1288–94.

**S2.2.2-27.** Dumas F, Cariou A, Manzo-Silberman S, et al. Immediate percutaneous coronary intervention is associated with better survival after out-of-hospital cardiac arrest: insights from the PROCAT (Parisian Region Out of hospital Cardiac Arrest) registry. *Circ Cardiovasc Interv.* 2010;3:200–7.

**S2.2.2-28.** Gupta S, Pressman GS, Figueredo VM. Incidence of, predictors for, and mortality associated with malignant ventricular arrhythmias in non-ST elevation myocardial infarction patients. *Coron Artery Dis.* 2010;21:460–5.

**S2.2.2-29.** Terkelsen CJ, Sorensen JT, Kaltoft AK, et al. Prevalence and significance of accelerated idioventricular rhythm in patients with ST-elevation myocardial infarction treated with primary percutaneous coronary intervention. *Am J Cardiol.* 2009;104:1641–6.

**S2.2.2-30.** Al-Khatib SM, Granger CB, Huang Y, et al. Sustained ventricular arrhythmias among patients with acute coronary syndromes with no ST-segment elevation: incidence, predictors, and outcomes. *Circulation.* 2002;106:309–12.

**S2.2.2-31.** Jabbari R, Engstrom T, Glinge C, et al. Incidence and risk factors of ventricular fibrillation before primary angioplasty in patients with first ST-elevation myocardial infarction: a nationwide study in Denmark. *J Am Heart Assoc.* 2015;4:e001399.

**S2.2.2-32.** Mehta RH, Starr AZ, Lopes RD, et al. Incidence of and outcomes associated with ventricular tachycardia or fibrillation in patients undergoing primary percutaneous coronary intervention. *JAMA.* 2009;301:1779–89.

**S2.2.2-33.** Volpi A, Cavalli A, Franzosi MG, et al. One-year prognosis of primary ventricular fibrillation complicating acute myocardial infarction. The GISSI (Gruppo Italiano per lo Studio della Streptochinasi nell'Infarto miocardico) investigators. *Am J Cardiol.* 1989;63:1174–8.

**S2.2.2-34.** El-Sherif N, Smith RA, Evans K. Canine ventricular arrhythmias in the late myocardial infarction period. 8. Epicardial mapping of reentrant circuits. *Circ Res.* 1981;49:255–65.

**S2.2.2-35.** Ackerman MJ, Priori SG, Willems S, et al. HRS/EHRA expert consensus statement on the state of genetic testing for the channelopathies and cardiomyopathies: this document was developed as a partnership between the Heart Rhythm Society (HRS) and the European Heart Rhythm Association (EHRA). *Europace.* 2011;13:1077–109.

**S2.2.2-36.** Nannenberg EA, Sijbrands EJ, Diksman LM, et al. Mortality of inherited arrhythmia syndromes:

insight into their natural history. *Circ Cardiovasc Genet.* 2012;5:183–9.

**S2.2.2-37.** Haïssaguerre M, Shah DC, Jais P, et al. Role of Purkinje conducting system in triggering of idiopathic ventricular fibrillation. *Lancet.* 2002;359:677–8.

**S2.2.2-38.** Haïssaguerre M, Shoda M, Jais P, et al. Mapping and ablation of idiopathic ventricular fibrillation. *Circulation.* 2002;106:962–7.

**S2.2.2-39.** Goldberger JJ, Buxton AE, Cain M, et al. Risk stratification for arrhythmic sudden cardiac death: identifying the roadblocks. *Circulation.* 2011;123:2423–30.

**S2.2.2-40.** Fishman GI, Chugh SS, DiMarco JP, et al. Sudden cardiac death prediction and prevention: report from a National Heart, Lung, and Blood Institute and Heart Rhythm Society Workshop. *Circulation.* 2010;122:2335–48.

**S2.2.2-41.** Myerburg RJ. Sudden cardiac death: exploring the limits of our knowledge. *J Cardiovasc Electrophysiol.* 2001;12:369–81.

**S2.2.2-42.** Kong MH, Fonarow GC, Peterson ED, et al. Systematic review of the incidence of sudden cardiac death in the United States. *J Am Coll Cardiol.* 2011;57:794–801.

**S2.2.2-43.** Mozaffarian D, Benjamin EJ, Go AS, et al. Heart disease and stroke statistics—2016 update: a report from the American Heart Association. *Circulation.* 2016;133:e38–360.

**S2.2.2-44.** Benjamin EJ, Blaha MJ, Chiuve SE, et al. Heart disease and stroke statistics—2017 update: a report from the American Heart Association. *Circulation.* 2017;135:e146–603.

**S2.2.2-45.** Merchant RM, Yang L, Becker LB, et al. Incidence of treated cardiac arrest in hospitalized patients in the United States. *Crit Care Med.* 2011;39:2401–6.

**S2.2.2-46.** Institute of Medicine. Committee on the Treatment of Cardiac Arrest: Current Status and Future Directions: Strategies to improve cardiac arrest survival: a time to act. Washington, DC: National Academic Press, 2015.

**S2.2.2-47.** Jollis JG, Granger CB. Improving care of out-of-hospital cardiac arrest: next steps. *Circulation.* 2016;134:2040–2.

**S2.2.2-48.** Daya MR, Schmicker R, May SH, et al. Current burden of cardiac arrest in the United States: report from the Resuscitation Outcomes Consortium. Paper commissioned by the Committee on the Treatment of Cardiac Arrest: Current Status and Future Directions. 2015.

**S2.2.2-49.** Myerburg RJ, Halperin H, Egan DA, et al. Pulseless electric activity: definition, causes, mechanisms, management, and research priorities for the next decade: report from a National Heart, Lung, and Blood Institute workshop. *Circulation.* 2013;128:2532–41.

**S2.2.2-50.** Myerburg RJ, Goldberger JJ. Sudden cardiac arrest risk assessment: population science and the individual risk mandate. *JAMA Cardiol.* 2017;2:689–94.

**S2.2.2-51.** Bogle BM, Ning H, Mehrotra S, et al. Lifetime risk for sudden cardiac death in the community. *J Am Heart Assoc.* 2016;5:e002398.

**S2.2.2-52.** Juntila MJ, Hookana E, Kaikonen KS, et al. Temporal trends in the clinical and pathological

characteristics of victims of sudden cardiac death in the absence of previously identified heart disease. *Circ Arrhythm Electrophysiol.* 2016;9:e003723.

**S2.2.2-53.** Hookana E, Juntila MJ, Puurunen VP, et al. Causes of nonischemic sudden cardiac death in the current era. *Heart Rhythm.* 2011;8:1570–5.

**S2.2.2-54.** Wong MK, Morrison LJ, Qiu F, et al. Trends in short- and long-term survival among out-of-hospital cardiac arrest patients alive at hospital arrival. *Circulation.* 2014;130:1883–90.

### 3. MECHANISMS OF VA

**S3.4-1.** Cherry EM, Fenton FH, Gilmour RF Jr. Mechanisms of ventricular arrhythmias: a dynamical systems-based perspective. *Am J Physiol Heart Circ Physiol.* 2012;302:H2451–63.

**S3.4-2.** Pandit SV, Jalife J. Rotors and the dynamics of cardiac fibrillation. *Circ Res.* 2013;112:849–62.

**S3.4-3.** Tsuji Y, Heijman J, Nattel S, et al. Electrical storm: recent pathophysiological insights and therapeutic consequences. *Basic Res Cardiol.* 2013;108:336.

**S3.4-4.** Roden DM. Predicting drug-induced QT prolongation and torsades de pointes. *J Physiol.* 2016;594:2459–68.

**S3.4-5.** Antzelevitch C, Nesterenko V, Shryock JC, et al. The role of late I<sub>Na</sub> in development of cardiac arrhythmias. *Handb Exp Pharmacol.* 2014;221:137–68.

**S3.4-6.** Lerman BB. Mechanism, diagnosis, and treatment of outflow tract tachycardia. *Nat Rev Cardiol.* 2015;12:597–608.

**S3.4-7.** Priori SG, Chen SR. Inherited dysfunction of sarcoplasmic reticulum Ca<sup>2+</sup> handling and arrhythmogenesis. *Circ Res.* 2011;108:871–83.

**S3.4-8.** Nogami A. Purkinje-related arrhythmias part I: monomorphic ventricular tachycardias. *Pacing Clin Electrophysiol.* 2011;34:624–50.

**S3.4-9.** Haïssaguerre M, Vigmond E, Stuyvers B, et al. Ventricular arrhythmias and the His-Purkinje system. *Nat Rev Cardiol.* 2016;13:155–66.

**S3.4-10.** Tabereaux PB, Dosdall DJ, Ideker RE. Mechanisms of VF maintenance: wandering wavelets, mother rotors, or foci. *Heart Rhythm.* 2009;6:405–15.

**S3.4-11.** Zhang J, Cooper DH, Desouza KA, et al. Electrophysiologic scar substrate in relation to VT: noninvasive high-resolution mapping and risk assessment with ECGI. *Pacing Clin Electrophysiol.* 2016;39:781–91.

**S3.4-12.** Fernandez-Armenta J, Penela D, Acosta J, et al. Substrate modification or ventricular tachycardia induction, mapping, and ablation as the first step? A randomized study. *Heart Rhythm.* 2016;13:1589–95.

### 4. GENERAL EVALUATION OF PATIENTS WITH DOCUMENTED OR SUSPECTED VA

#### 4.1. History and Physical Examination

**S4.1-1.** Middlekauff HR, Stevenson WG, Stevenson LW, et al. Syncope in advanced heart failure: high risk of sudden death regardless of origin of syncope. *J Am Coll Cardiol.* 1993;21:110–6.

**S4.1-2.** Myerburg RJ, Castellanos A. Cardiac arrest and sudden cardiac death - Chapter 39. In: Mann DL,

Zipes DP, Libby P, et al., editors. Braunwald's Heart Disease: A Textbook of Cardiovascular Medicine. 10th ed. Oxford, UK: Elsevier; 2015:821-60.

**S4.1-3.** Shen WK, Sheldon RS, Benditt DG, et al. 2017 ACC/AHA/HRS guideline for the evaluation and management of patients with syncope: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2017;70:e39-110.

**S4.1-4.** Soteriades ES, Evans JC, Larson MG, et al. Incidence and prognosis of syncope. *N Engl J Med.* 2002;347:878-85.

**S4.1-5.** Noda T, Shimizu W, Taguchi A, et al. Malignant entity of idiopathic ventricular fibrillation and polymorphic ventricular tachycardia initiated by premature extrasystoles originating from the right ventricular outflow tract. *J Am Coll Cardiol.* 2005;46:1288-94.

**S4.1-6.** Viskin S, Rosso R, Rogowski O, et al. The "short-coupled" variant of right ventricular outflow ventricular tachycardia: a not-so-benign form of benign ventricular tachycardia? *J Cardiovasc Electrophysiol.* 2005;16:912-6.

**S4.1-7.** Zimetbaum P, Josephson ME. Evaluation of patients with palpitations. *N Engl J Med.* 1998;338:1369-73.

**S4.1-8.** Credible meds. Available at: <http://www.crediblemeds.org>. Accessed December 26, 2016.

**S4.1-9.** Brugada drugs. Available at: <http://www.brugadadrugs.org>. Accessed October 6, 2016.

**S4.1-10.** Konigstein M, Rosso R, Topaz G, et al. Drug-induced Brugada syndrome: clinical characteristics and risk factors. *Heart Rhythm.* 2016;13:1083-7.

**S4.1-11.** Basso C, Perazzolo MM, Rizzo S, et al. Arrhythmic mitral valve prolapse and sudden cardiac death. *Circulation.* 2015;132:556-66.

**S4.1-12.** Nordhues BD, Sontis KC, Scott CG, et al. Bileaflet mitral valve prolapse and risk of ventricular dysrhythmias and death. *J Cardiovasc Electrophysiol.* 2016;27:463-8.

**S4.1-13.** Sriram CS, Syed FF, Ferguson ME, et al. Malignant bileaflet mitral valve prolapse syndrome in patients with otherwise idiopathic out-of-hospital cardiac arrest. *J Am Coll Cardiol.* 2013;62:222-30.

**S4.1-14.** Huikuri HV, Castellanos A, Myerburg RJ. Sudden death due to cardiac arrhythmias. *N Engl J Med.* 2001;345:1473-82.

**S4.1-15.** Krahn AD, Healey JS, Simpson CS, et al. Sentinel symptoms in patients with unexplained cardiac arrest: from the Cardiac Arrest Survivors with Preserved Ejection Fraction Registry (CASPER). *J Cardiovasc Electrophysiol.* 2012;23:60-6.

**S4.1-16.** Ruwald MH, Hansen ML, Lamberts M, et al. The relation between age, sex, comorbidity, and pharmacotherapy and the risk of syncope: a Danish nationwide study. *Europace.* 2012;14:1506-14.

## 4.2. Noninvasive Evaluation

### 4.2.1. 12-lead ECG and Exercise Testing

**S4.2.1-1.** Brugada P, Brugada J, Mont L, et al. A new approach to the differential diagnosis of a regular tachycardia with a wide QRS complex. *Circulation.* 1991;83:1649-59.

**S4.2.1-2.** Wellens HJ, Bar FW, Lie KI. The value of the electrocardiogram in the differential diagnosis of a

tachycardia with a widened QRS complex. *Am J Med.* 1978;64:27-33.

**S4.2.1-3.** Steinman RT, Herrera C, Schuger CD, et al. Wide QRS tachycardia in the conscious adult. Ventricular tachycardia is the most frequent cause. *JAMA.* 1989;261:1013-6.

**S4.2.1-4.** Elhendy A, Chandrasekaran K, Gersh BJ, et al. Functional and prognostic significance of exercise-induced ventricular arrhythmias in patients with suspected coronary artery disease. *Am J Cardiol.* 2002;90:95-100.

**S4.2.1-5.** Grady TA, Chiu AC, Snader CE, et al. Prognostic significance of exercise-induced left bundle-branch block. *JAMA.* 1998;279:153-6.

**S4.2.1-6.** Perez-Rodon J, Martinez-Alday J, Baron-Esquivias G, et al. Prognostic value of the electrocardiogram in patients with syncope: data from the group for syncope study in the emergency room (GESINUR). *Heart Rhythm.* 2014;11:2035-44.

**S4.2.1-7.** Baldasseroni S, Opasich C, Gorini M, et al. Left bundle-branch block is associated with increased 1-year sudden and total mortality rate in 5517 outpatients with congestive heart failure: a report from the Italian network on congestive heart failure. *Am Heart J.* 2002;143:398-405.

**S4.2.1-8.** Buxton AE, Sweeney MO, Wathen MS, et al. QRS duration does not predict occurrence of ventricular tachyarrhythmias in patients with implanted cardioverter-defibrillators. *J Am Coll Cardiol.* 2005;46:310-6.

**S4.2.1-9.** Desai AD, Yaw TS, Yamazaki T, et al. Prognostic significance of quantitative QRS duration. *Am J Med.* 2006;119:600-6.

**S4.2.1-10.** Dhar R, Alsheikh-Ali AA, Estes NA 3rd, et al. Association of prolonged QRS duration with ventricular tachyarrhythmias and sudden cardiac death in the Multicenter Automatic Defibrillator Implantation Trial II (MADIT-II). *Heart Rhythm.* 2008;5:807-13.

**S4.2.1-11.** Freedman RA, Alderman EL, Sheffield LT, et al. Bundle branch block in patients with chronic coronary artery disease: angiographic correlates and prognostic significance. *J Am Coll Cardiol.* 1987;10:73-80.

**S4.2.1-12.** Iuliano S, Fisher SG, Karasik PE, et al. QRS duration and mortality in patients with congestive heart failure. *Am Heart J.* 2002;143:1085-91.

**S4.2.1-13.** Kadish A, Dyer A, Daubert JP, et al. Prophylactic defibrillator implantation in patients with nonischemic dilated cardiomyopathy. *N Engl J Med.* 2004;350:2151-8.

**S4.2.1-14.** Zimetbaum PJ, Buxton AE, Batsford W, et al. Electrocardiographic predictors of arrhythmic death and total mortality in the multicenter unsustained tachycardia trial. *Circulation.* 2004;110:766-9.

**S4.2.1-15.** Costantini O, Hohnloser SH, Kirk MM, et al. The ABCD (Alternans Before Cardioverter Defibrillator) Trial: strategies using T-wave alternans to improve efficiency of sudden cardiac death prevention. *J Am Coll Cardiol.* 2009;53:471-9.

**S4.2.1-16.** Monasterio V, Martinez JP, Laguna P, et al. Prognostic value of average T-wave alternans and QT variability for cardiac events in MADIT-II patients. *J Electrocardiol.* 2013;46:480-6.

**S4.2.1-17.** Chow T, Kereiakes DJ, Onufre J, et al. Does microvolt T-wave alternans testing predict ventricular tachyarrhythmias in patients with ischemic

cardiomyopathy and prophylactic defibrillators? The MASTER (Microvolt T Wave Alternans Testing for Risk Stratification of Post-Myocardial Infarction Patients) trial. *J Am Coll Cardiol.* 2008;52:1607-15.

**S4.2.1-18.** Gupta A, Hoang DD, Karliner L, et al. Ability of microvolt T-wave alternans to modify risk assessment of ventricular tachyarrhythmic events: a meta-analysis. *Am Heart J.* 2012;163:354-64.

**S4.2.1-19.** Bloomfield DM, Steinman RC, Namerow PB, et al. Microvolt T-wave alternans distinguishes between patients likely and patients not likely to benefit from implanted cardiac defibrillator therapy: a solution to the Multicenter Automatic Defibrillator Implantation Trial (MADIT) II conundrum. *Circulation.* 2004;110:1885-9.

### 4.2.2. Ambulatory Electrocardiography

**S4.2.2-1.** Barrett PM, Komatireddy R, Haaser S, et al. Comparison of 24-hour Holter monitoring with 14-day novel adhesive patch electrocardiographic monitoring. *Am J Med.* 2014;127:95-7.

**S4.2.2-2.** de Asmundis C, Conte G, Sieira J, et al. Comparison of the patient-activated event recording system vs. traditional 24 h Holter electrocardiography in individuals with paroxysmal palpitations or dizziness. *Europace.* 2014;16:1231-5.

**S4.2.2-3.** Linzer M, Pritchett EL, Pontinen M, et al. Incremental diagnostic yield of loop electrocardiographic recorders in unexplained syncope. *Am J Cardiol.* 1990;66:214-9.

**S4.2.2-4.** Turakhia MP, Hoang DD, Zimetbaum P, et al. Diagnostic utility of a novel leadless arrhythmia monitoring device. *Am J Cardiol.* 2013;112:520-4.

### 4.2.3. Implanted Cardiac Monitors

**S4.2.3-1.** Bloch Thomsen PE, Jons C, Raatikainen MJ, et al. Long-term recording of cardiac arrhythmias with an implantable cardiac monitor in patients with reduced ejection fraction after acute myocardial infarction: the Cardiac Arrhythmias and Risk Stratification After Acute Myocardial Infarction (CARISMA) study. *Circulation.* 2010;122:1258-64.

**S4.2.3-2.** Krahn AD, Klein GJ, Yee R, et al. Use of an extended monitoring strategy in patients with problematic syncope. *Reveal Investigators.* *Circulation.* 1999;99:406-10.

**S4.2.3-3.** Solbiati M, Costantino G, Casazza G, et al. Implantable loop recorder versus conventional diagnostic workup for unexplained recurrent syncope. *Cochrane Database Syst. Rev.* 2016;4:CD011637.

**S4.2.3-4.** Volosin K, Stadler RW, Wyszynski R, et al. Tachycardia detection performance of implantable loop recorders: results from a large 'real-life' patient cohort and patients with induced ventricular arrhythmias. *Europace.* 2013;15:1215-22.

**S4.2.3-5.** Linzer M, Pritchett EL, Pontinen M, et al. Incremental diagnostic yield of loop electrocardiographic recorders in unexplained syncope. *Am J Cardiol.* 1990;66:214-9.

**S4.2.3-6.** Turakhia MP, Hoang DD, Zimetbaum P, et al. Diagnostic utility of a novel leadless arrhythmia monitoring device. *Am J Cardiol.* 2013;112:520-4.

### 4.2.4. Noninvasive Cardiac Imaging

**S4.2.4-1.** Solomon SD, Zelenkofske S, McMurray JJ, et al. Sudden death in patients with myocardial

infarction and left ventricular dysfunction, heart failure, or both. *N Engl J Med.* 2005;352:2581–8.

**S4.2.4-2.** Gula LJ, Klein GJ, Hellkamp AS, et al. Ejection fraction assessment and survival: an analysis of the Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT). *Am Heart J.* 2008;156:1196–200.

**S4.2.4-3.** Rauch B, Schiele R, Schneider S, et al. OMEGA, a randomized, placebo-controlled trial to test the effect of highly purified omega-3 fatty acids on top of modern guideline-adjusted therapy after myocardial infarction. *Circulation.* 2010;122:2152–9.

**S4.2.4-4.** Shen WK, Sheldon RS, Benditt DG, et al. 2017 ACC/AHA/HRS guideline for the evaluation and management of patients with syncope: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2017;70:e39–110.

**S4.2.4-5.** Coleman GC, Shaw PW, Balfour PC Jr., et al. Prognostic value of myocardial scarring on CMR in patients with cardiac sarcoidosis: a systematic review and meta-analysis. *J Am Coll Cardiol Img.* 2016;10:411–20.

**S4.2.4-6.** Di Marco A, Anguera I, Schmitt M, et al. Late gadolinium enhancement and the risk for ventricular arrhythmias or sudden death in dilated cardiomyopathy: systematic review and meta-analysis. *JACC Heart Fail.* 2017;5:28–38.

**S4.2.4-7.** Greulich S, Deluigi CC, Gloekler S, et al. CMR imaging predicts death and other adverse events in suspected cardiac sarcoidosis. *J Am Coll Cardiol Img.* 2013;6:501–11.

**S4.2.4-8.** Piers SR, Tao Q, van Huls van Taxis CF, et al. Contrast-enhanced MRI-derived scar patterns and associated ventricular tachycardias in nonischemic cardiomyopathy: implications for the ablation strategy. *Circ Arrhythm Electrophysiol.* 2013;6:875–83.

**S4.2.4-9.** White JA, Fine NM, Gula L, et al. Utility of cardiovascular magnetic resonance in identifying substrate for malignant ventricular arrhythmias. *Circ Cardiovasc Imaging.* 2012;5:12–20.

#### 4.2.5. Biomarkers

**S4.2.5-1.** Ahmad T, Fuizat M, Neely B, et al. Biomarkers of myocardial stress and fibrosis as predictors of mode of death in patients with chronic heart failure. *JACC Heart Fail.* 2014;2:260–8.

**S4.2.5-2.** Scott PA, Barry J, Roberts PR, et al. Brain natriuretic peptide for the prediction of sudden cardiac death and ventricular arrhythmias: a meta-analysis. *Eur J Heart Fail.* 2009;11:958–66.

**S4.2.5-3.** Levine YC, Rosenberg MA, Mittleman M, et al. B-type natriuretic peptide is a major predictor of ventricular tachyarrhythmias. *Heart Rhythm.* 2014;11:1109–16.

**S4.2.5-4.** Berger R, Huelsman M, Strecker K, et al. B-type natriuretic peptide predicts sudden death in patients with chronic heart failure. *Circulation.* 2002;105:2392–7.

**S4.2.5-5.** Korngold EC, Januzzi JL Jr., Gantzer ML, et al. Amino-terminal pro-B-type natriuretic peptide and high-sensitivity C-reactive protein as predictors of sudden cardiac death among women. *Circulation.* 2009;119:2868–76.

**S4.2.5-6.** Patton KK, Sotoodehnia N, DeFilippi C, et al. N-terminal pro-B-type natriuretic peptide is associated

with sudden cardiac death risk: the Cardiovascular Health Study. *Heart Rhythm.* 2011;8:228–33.

**S4.2.5-7.** Hussein AA, Gottsdiner JS, Bartz TM, et al. Cardiomyocyte injury assessed by a highly sensitive troponin assay and sudden cardiac death in the community: the Cardiovascular Health Study. *J Am Coll Cardiol.* 2013;62:2112–20.

#### 4.2.6. Genetic Considerations in Arrhythmia Syndromes

**S4.2.6-1.** Andreasen C, Refsgaard L, Nielsen JB, et al. Mutations in genes encoding cardiac ion channels previously associated with sudden infant death syndrome (SIDS) are present with high frequency in new exome data. *Can J Cardiol.* 2013;29:1104–9.

**S4.2.6-2.** Jabbari J, Jabbari R, Nielsen MW, et al. New exome data question the pathogenicity of genetic variants previously associated with catecholaminergic polymorphic ventricular tachycardia. *Circ Cardiovasc Genetics.* 2013;481–9.

**S4.2.6-3.** Paludan-Muller C, Ahlberg G, Ghouse J, et al. Integration of 60,000 exomes and ACMG guidelines question the role of catecholaminergic polymorphic ventricular tachycardia-associated variants. *Clin Genet.* 2017;91:63–72.

**S4.2.6-4.** Refsgaard L, Holst AG, Sadjadieh G, et al. High prevalence of genetic variants previously associated with LQT syndrome in new exome data. *Eur J Hum Genet.* 2012;20:905–8.

**S4.2.6-5.** Risgaard B, Jabbari R, Refsgaard L, et al. High prevalence of genetic variants previously associated with Brugada syndrome in new exome data. *Clin Genet.* 2013;84:489–95.

**S4.2.6-6.** Costa J, Lopes CM, Barsheheshet A, et al. Combined assessment of sex- and mutation-specific information for risk stratification in type 1 long QT syndrome. *Heart Rhythm.* 2012;9:892–8.

**S4.2.6-7.** Hershberger RE, Lindenfeld J, Mestroni L, et al. Genetic evaluation of cardiomyopathy—a Heart Failure Society of America practice guideline. *J Cardiac Fail.* 2009;15:83–97.

**S4.2.6-8.** Kim JA, Lopes CM, Moss AJ, et al. Trigger-specific risk factors and response to therapy in long QT syndrome type 2. *Heart Rhythm.* 2010;7:1797–805.

**S4.2.6-9.** Midgaldovich D, Moss AJ, Lopes CM, et al. Mutation and gender-specific risk in type 2 long QT syndrome: implications for risk stratification for life-threatening cardiac events in patients with long QT syndrome. *Heart Rhythm.* 2011;8:1537–43.

**S4.2.6-10.** Crotti L, Marcou CA, Tester DJ, et al. Spectrum and prevalence of mutations involving BrS1-through BrS12-susceptibility genes in a cohort of unrelated patients referred for Brugada syndrome genetic testing: implications for genetic testing. *J Am Coll Cardiol.* 2012;60:1410–8.

**S4.2.6-11.** Probst V, Wilde AA, Barc J, et al. SCN5A mutations and the role of genetic background in the pathophysiology of Brugada syndrome. *Circ Cardiovasc Genetics.* 2009;2:552–7.

**S4.2.6-12.** Dintzis RZ. Genetic variation and the meiotic process. *Res Publ Assoc Res Nerv Ment Dis.* 1991;69:39–46.

**S4.2.6-13.** Groeneweg JA, Bhonsale A, James CA, et al. Clinical presentation, long-term follow-up, and outcomes of 1001 arrhythmogenic right ventricular

dysplasia cardiomyopathy patients and family members. *Circ Cardiovasc Genet.* 2015;8:437–46.

**S4.2.6-14.** Marcus Fl, Edson S, Towbin JA. Genetics of arrhythmogenic right ventricular cardiomyopathy: a practical guide for physicians. *J Am Coll Cardiol.* 2013;61:1945–8.

**S4.2.6-15.** Quarta G, Muir A, Pantazis A, et al. Familial evaluation in arrhythmogenic right ventricular cardiomyopathy: impact of genetics and revised task force criteria. *Circulation.* 2011;123:2701–9.

**S4.2.6-16.** Rigato I, Baucé B, Rampazzo A, et al. Compound and digenic heterozygosity predicts lifetime arrhythmic outcome and sudden cardiac death in desmosomal gene-related arrhythmogenic right ventricular cardiomyopathy. *Circ Cardiovasc Genet.* 2013;6:533–42.

**S4.2.6-17.** Ackerman MJ, Priori SG, Willems S, et al. HRS/EHRA expert consensus statement on the state of genetic testing for the channelopathies and cardiomyopathies: this document was developed as a partnership between the Heart Rhythm Society (HRS) and the European Heart Rhythm Association (EHRA). *Europace.* 2011;13:1077–109.

**S4.2.6-18.** Gollob MH, Blér L, Brugada R, et al. Recommendations for the use of genetic testing in the clinical evaluation of inherited cardiac arrhythmias associated with sudden cardiac death: Canadian Cardiovascular Society/Canadian Heart Rhythm Society joint position paper. *Can J Cardiol.* 2011;27:232–45.

**S4.2.6-19.** Kumar S, Peters S, Thompson T, et al. Familial cardiological and targeted genetic evaluation: low yield in sudden unexplained death and high yield in unexplained cardiac arrest syndromes. *Heart Rhythm.* 2013;10:1653–60.

**S4.2.6-20.** Priori SG, Wilde AA, Horie M, et al. HRS/EHRA/APHS expert consensus statement on the diagnosis and management of patients with inherited primary arrhythmia syndromes. *Heart Rhythm.* 2013;10:1932–63.

**S4.2.6-21.** Tester DJ, Medeiros-Domingo A, Will ML, et al. Unexplained drownings and the cardiac channelopathies: a molecular autopsy series. *Mayo Clin Proc.* 2011;86:941–7.

**S4.2.6-22.** Tzimas I, Zingraf JC, Bajanowski T, et al. The role of known variants of KCNQ1, KCNH2, KCNE1, SCNSA, and NOS1AP in water-related deaths. *Int J Legal Med.* 2016;130:1575–9.

**S4.2.6-23.** Wang D, Shah KR, Um SY, et al. Cardiac channelopathy testing in 274 ethnically diverse sudden unexplained deaths. *Forensic Sci Int.* 2014;237:90–9.

**S4.2.6-24.** Laedtke AL, O'Neill SM, Rubinstein WS, et al. Family physicians' awareness and knowledge of the Genetic Information Non-Discrimination Act (GINA). *J Genet Couns.* 2012;21:345–52.

**S4.2.6-25.** Vavolizza RD, Kalia I, Erskine AK, et al. Disclosing genetic information to family members about inherited cardiac arrhythmias: an obligation or a choice? *J Genet Couns.* 2015;24:608–15.

**S4.2.6-26.** Christiaans I, van Langen IM, Birnie E, et al. Genetic counseling and cardiac care in predictively tested hypertrophic cardiomyopathy mutation carriers: the patients' perspective. *Am J Med Genet. Part A.* 2009;149A:1444–51.

**S4.2.6-27.** Hamang A, Eide GE, Rokne B, et al. Predictors of heart-focused anxiety in patients

undergoing genetic investigation and counseling of long QT syndrome or hypertrophic cardiomyopathy: a one year follow-up. *J Genet Counsel.* 2012;21:72–84.

#### 4.3. Invasive Testing

##### 4.3.1. Invasive Cardiac Imaging: Cardiac Catheterization or CT Angiography

**S4.3.1-1.** Dumas F, Bougouin W, Geri G, et al. Emergency PCI in post-cardiac arrest patients without ST-segment elevation pattern: insights from the PROCAT II registry. *J Am Coll Cardiol Intv.* 2016;9:1011–8.

**S4.3.1-2.** Cronier P, Vignon P, Bouferrache K, et al. Impact of routine percutaneous coronary intervention after out-of-hospital cardiac arrest due to ventricular fibrillation. *Crit Care.* 2011;15:R122.

**S4.3.1-3.** Spaulding CM, Joly LM, Rosenberg A, et al. Immediate coronary angiography in survivors of out-of-hospital cardiac arrest. *N Engl J Med.* 1997;336:1629–33.

**S4.3.1-4.** Zanuttini D, Armellini I, Nucifora G, et al. Impact of emergency coronary angiography on in-hospital outcome of unconscious survivors after out-of-hospital cardiac arrest. *Am J Cardiol.* 2012;110:1723–8.

**S4.3.1-5.** O’Gara PT, Kushner FG, Ascheim DD, et al. 2013 ACCF/AHA guideline for the management of ST-elevation myocardial infarction: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2013;61:e78–140.

##### 4.3.2. Electrophysiological Study for VA

**S4.3.2-1.** Buxton AE, Lee KL, DiCarlo L, et al. Electrophysiologic testing to identify patients with coronary artery disease who are at risk for sudden death. Multicenter Unsustained Tachycardia Trial Investigators. *N Engl J Med.* 2000;342:1937–45.

**S4.3.2-2.** Buxton AE, Lee KL, Hafley GE, et al. Relation of ejection fraction and inducible ventricular tachycardia to mode of death in patients with coronary artery disease: an analysis of patients enrolled in the multicenter unsustained tachycardia trial. *Circulation.* 2002;106:2466–72.

**S4.3.2-3.** Costantini O, Hohnloser SH, Kirk MM, et al. The ABCD (Alternans Before Cardioverter Defibrillator) Trial: strategies using T-wave alternans to improve efficiency of sudden cardiac death prevention. *J Am Coll Cardiol.* 2009;53:471–9.

**S4.3.2-4.** Bourke JP, Richards DA, Ross DL, et al. Routine programmed electrical stimulation in survivors of acute myocardial infarction for prediction of spontaneous ventricular tachyarrhythmias during follow-up: results, optimal stimulation protocol and cost-effective screening. *J Am Coll Cardiol.* 1991;18:780–8.

**S4.3.2-5.** Bailey JJ, Berson AS, Handelman H, et al. Utility of current risk stratification tests for predicting major arrhythmic events after myocardial infarction. *J Am Coll Cardiol.* 2001;38:1902–11.

**S4.3.2-6.** Schmitt C, Barthel P, Ndreppepa G, et al. Value of programmed ventricular stimulation for prophylactic internal cardioverter-defibrillator implantation in postinfarction patients preselected by noninvasive risk stratifiers. *J Am Coll Cardiol.* 2001;37:1901–7.

**S4.3.2-7.** Hilfiker G, Schoenenberger AW, Erne P, et al. Utility of electrophysiological studies to predict arrhythmic events. *World J Cardiol.* 2015;7:344–50.

**S4.3.2-8.** Bardy GH, Lee KL, Mark DB, et al. Amiodarone or an implantable cardioverter-defibrillator for congestive heart failure. *N Engl J Med.* 2005;352:225–37.

**S4.3.2-9.** Buxton AE, Lee KL, Fisher JD, et al. A randomized study of the prevention of sudden death in patients with coronary artery disease. Multicenter Unsustained Tachycardia Trial Investigators. *N Engl J Med.* 1999;341:1882–90.

**S4.3.2-10.** Moss AJ, Hall WJ, Cannom DS, et al. Improved survival with an implanted defibrillator in patients with coronary disease at high risk for ventricular arrhythmia. Multicenter Automatic Defibrillator Implantation Trial Investigators. *N Engl J Med.* 1996;335:1933–40.

**S4.3.2-11.** Moss AJ, Zareba W, Hall WJ, et al. Prophylactic implantation of a defibrillator in patients with myocardial infarction and reduced ejection fraction. *N Engl J Med.* 2002;346:877–83.

**S4.3.2-12.** Bhandari AK, Shapiro WA, Morady F, et al. Electrophysiologic testing in patients with the long QT syndrome. *Circulation.* 1985;71:63–71.

**S4.3.2-13.** Giustetto C, Di MF, Wolpert C, et al. Short QT syndrome: clinical findings and diagnostic-therapeutic implications. *Eur Heart J.* 2006;27:2440–7.

**S4.3.2-14.** Giustetto C, Schimpff R, Mazzanti A, et al. Long-term follow-up of patients with short QT syndrome. *J Am Coll Cardiol.* 2011;58:587–95.

**S4.3.2-15.** Mahida S, Derval N, Sacher F, et al. Role of electrophysiological studies in predicting risk of ventricular arrhythmia in early repolarization syndrome. *J Am Coll Cardiol.* 2015;65:151–9.

**S4.3.2-16.** Priori SG, Napolitano C, Memmi M, et al. Clinical and molecular characterization of patients with catecholaminergic polymorphic ventricular tachycardia. *Circulation.* 2002;106:69–74.

**S4.3.2-17.** Gatzoulis KA, Vouliotis AI, Tsachiris D, et al. Primary prevention of sudden cardiac death in a non-ischemic dilated cardiomyopathy population: reappraisal of the role of programmed ventricular stimulation. *Circ Arrhythm Electrophysiol.* 2013;6:504–12.

**S4.3.2-18.** Bremilla-Perron B, Suty-Selton C, Beurrier D, et al. Differences in mechanisms and outcomes of syncope in patients with coronary disease or idiopathic left ventricular dysfunction as assessed by electrophysiologic testing. *J Am Coll Cardiol.* 2004;44:594–601.

**S4.3.2-19.** Garson A Jr., Dick M, Fournier A, et al. The long QT syndrome in children. An international study of 287 patients. *Circulation.* 1993;87:1866–72.

**S4.3.2-20.** Hobbs JB, Peterson DR, Moss AJ, et al. Risk of aborted cardiac arrest or sudden cardiac death during adolescence in the long-QT syndrome. *JAMA.* 2006;296:1249–54.

**S4.3.2-21.** Iribarren C, Round AD, Peng JA, et al. Short QT in a cohort of 1.7 million persons: prevalence, correlates, and prognosis. *Ann Noninvasive Electrocardiol.* 2014;19:490–500.

**S4.3.2-22.** Kaufman ES, McNitt S, Moss AJ, et al. Risk of death in the long QT syndrome when a sibling has died. *Heart Rhythm.* 2008;5:831–6.

**S4.3.2-23.** Kimbrough J, Moss AJ, Zareba W, et al. Clinical implications for affected parents and siblings

of probands with long-QT syndrome. *Circulation.* 2001;104:557–62.

**S4.3.2-24.** Mazzanti A, Kanthan A, Monteforte N, et al. Novel insight into the natural history of short QT syndrome. *J Am Coll Cardiol.* 2014;63:1300–8.

**S4.3.2-25.** Wong JA, Gula LJ, Klein GJ, et al. Utility of treadmill testing in identification and genotype prediction in long-QT syndrome. *Circ Arrhythm Electrophysiol.* 2010;3:120–5.

**S4.3.2-26.** Walker BD, Krahn AD, Klein GJ, et al. Burst bicycle exercise facilitates diagnosis of latent long QT syndrome. *Am Heart J.* 2005;150:1059–63.

**S4.3.2-27.** Waks JW, Sitzani CM, Soliman EZ, et al. Global electric heterogeneity risk score for prediction of sudden cardiac death in the general population: the Atherosclerosis Risk in Communities (ARIC) and Cardiovascular Health (CHS) Studies. *Circulation.* 2016;133:2222–34.

**S4.3.2-28.** Priori SG, Napolitano C, Schwartz PJ, et al. Association of long QT syndrome loci and cardiac events among patients treated with beta-blockers. *JAMA.* 2004;292:1341–4.

**S4.3.2-29.** Priori SG, Schwartz PJ, Napolitano C, et al. Risk stratification in the long-QT syndrome. *N Engl J Med.* 2003;348:1866–74.

**S4.3.2-30.** Wedekind H, Burde D, Zumhagen S, et al. QT interval prolongation and risk for cardiac events in genotyped LQTS-index children. *Eur J Pediatr.* 2009;168:1107–15.

**S4.3.2-31.** Goldenberg I, Horr S, Moss AJ, et al. Risk for life-threatening cardiac events in patients with genotype-confirmed long-QT syndrome and normal-range corrected QT intervals. *J Am Coll Cardiol.* 2011;57:51–9.

**S4.3.2-32.** Nannenber EA, Sijbrands EJ, Dijksman LM, et al. Mortality of inherited arrhythmia syndromes: insight into their natural history. *Circ Cardiovasc Genet.* 2012;5:183–9.

## 5. THERAPIES FOR TREATMENT OR PREVENTION OF VA

### 5.1. Medication Therapy

**S5.1.5.2-1.** Vaughan Williams EM. A classification of antiarrhythmic actions reassessed after a decade of new drugs. *J Clin Pharmacol.* 1984;24:129–47.

**S5.1.5.2-2.** Schleifer JW, Sorajja D, Shen WK. Advances in the pharmacologic treatment of ventricular arrhythmias. *Expert Opin Pharmacother.* 2015;16:2637–51.

**S5.1.5.2-3.** Kudenchuk PJ, Brown SP, Daya M, et al. Amiodarone, lidocaine, or placebo in out-of-hospital cardiac arrest. *N Engl J Med.* 2016;371:1171–22.

**S5.1.5.2-4.** Mazzanti A, Maragna R, Faragli A, et al. Gene-specific therapy with mexiletine reduces arrhythmic events in patients with long QT syndrome type 3. *J Am Coll Cardiol.* 2016;67:1053–8.

**S5.1.5.2-5.** Watanabe H, Chopra N, Laver D, et al. Flecainide prevents catecholaminergic polymorphic ventricular tachycardia in mice and humans. *Nat Med.* 2009;15:380–3.

**S5.1.5.2-6.** Gupta T, Khera S, Kolte D, et al. Antiarrhythmic properties of ranolazine: a review of the current evidence. *Int J Cardiol.* 2015;187:66–74.

- S5.1.5.2-7.** Bunch TJ, Mahapatra S, Murdock D, et al. Ranolazine reduces ventricular tachycardia burden and ICD shocks in patients with drug-refractory ICD shocks. *Pacing Clin Electrophysiol.* 2011;34:1600–6.
- S5.1.5.2-8.** Scirica BM, Braunwald E, Belardinelli L, et al. Relationship between nonsustained ventricular tachycardia after non-ST-elevation acute coronary syndrome and sudden cardiac death: observations from the metabolic efficiency with ranolazine for less ischemia in non-ST-elevation acute coronary syndrome-thrombolysis in myocardial infarction 36 (MERLIN-TIMI 36) randomized controlled trial. *Circulation.* 2010;122:455–62.
- S5.1.5.2-9.** Zareba W, Daubert JP, Beck CA, et al. Ranolazine in High-Risk ICD Patients (RAID) Trial. Presented at: Heart Rhythm Society Annual Scientific Sessions. 2017; LBCT02-01.
- S5.1.5.2-10.** Reiter MJ, Reiffel JA. Importance of beta blockade in the therapy of serious ventricular arrhythmias. *Am J Cardiol.* 1998;82:91–191.
- S5.1.5.2-11.** Ellison KE, Hafley GE, Hickey K, et al. Effect of beta-blocking therapy on outcome in the Multicenter UnSustained Tachycardia Trial (MUSTT). *Circulation.* 2002;106:2694–9.
- S5.1.5.2-12.** Reiken S, Wehrens XH, Vest JA, et al. Beta-blockers restore calcium release channel function and improve cardiac muscle performance in human heart failure. *Circulation.* 2003;107:2459–66.
- S5.1.5.2-13.** MERIT-HF Group. Effect of metoprolol CR/XL in chronic heart failure: Metoprolol CR/XL Randomised Intervention Trial in Congestive Heart Failure (MERIT-HF). *Lancet.* 1999;353:2001–7.
- S5.1.5.2-14.** Packer M, Bristow MR, Cohn JN, et al. The effect of carvedilol on morbidity and mortality in patients with chronic heart failure. U.S. Carvedilol Heart Failure Study Group. *N Engl J Med.* 1996;334:1349–55.
- S5.1.5.2-15.** Committees of the Cardiac insufficiency Bisoprolol Study II. The cardiac insufficiency Bisoprolol Study II (CIBIS-II): a randomised trial. *Lancet.* 1999; 353:9–13.
- S5.1.5.2-16.** A randomized trial of propranolol in patients with acute myocardial infarction. Mortality results. *JAMA.* 1982;247:1707–14.
- S5.1.5.2-17.** Kontos MC, Diercks DB, Ho PM, et al. Treatment and outcomes in patients with myocardial infarction treated with acute beta-blocker therapy: results from the American College of Cardiology's NCDR(R). *Am Heart J.* 2011;161:864–70.
- S5.1.5.2-18.** Nademanee K, Taylor R, Bailey WE, et al. Treating electrical storm: sympathetic blockade versus advanced cardiac life support-guided therapy. *Circulation.* 2000;102:742–7.
- S5.1.5.2-19.** Kittayaphong R, Bhuripanyo K, Punlee K, et al. Effect of atenolol on symptomatic ventricular arrhythmia without structural heart disease: a randomized placebo-controlled study. *Am Heart J.* 2002; 144:e10.
- S5.1.5.2-20.** Hirsowitz G, Podrid PJ, Lampert S, et al. The role of beta blocking agents as adjunct therapy to membrane stabilizing drugs in malignant ventricular arrhythmia. *Am Heart J.* 1986;111:852–60.
- S5.1.5.2-21.** Connolly SJ. Meta-analysis of antiarrhythmic drug trials. *Am J Cardiol.* 1999;84:90R–3R.
- S5.1.5.2-22.** Steinberg JS, Martins J, Sadanandan S, et al. Antiarrhythmic drug use in the implantable defibrillator arm of the Antiarrhythmics Versus Implantable Defibrillators (AVID) Study. *Am Heart J.* 2001;142:520–9.
- S5.1.5.2-23.** Farre J, Romero J, Rubio JM, et al. Amiodarone and "primary" prevention of sudden death: critical review of a decade of clinical trials. *Am J Cardiol.* 1999;83:55D–63D.
- S5.1.5.2-24.** Bardy GH, Lee KL, Mark DB, et al. Amiodarone or an implantable cardioverter-defibrillator for congestive heart failure. *N Engl J Med.* 2005;352:225–37.
- S5.1.5.2-25.** Thomas KL, Al-Khatib SM, Lohknygina Y, et al. Amiodarone use after acute myocardial infarction complicated by heart failure and/or left ventricular dysfunction may be associated with excess mortality. *Am Heart J.* 2008;155:87–93.
- S5.1.5.2-26.** Claro JC, Candia R, Rada G, et al. Amiodarone versus other pharmacological interventions for prevention of sudden cardiac death. *Cochrane Database Syst. Rev.* 2015;12:CD008093.
- S5.1.5.2-27.** Kudenchuk PJ, Cobb LA, Copass MK, et al. Amiodarone for resuscitation after out-of-hospital cardiac arrest due to ventricular fibrillation. *N Engl J Med.* 1999;341:871–8.
- S5.1.5.2-28.** Dorian P, Cass D, Schwartz B, et al. Amiodarone as compared with lidocaine for shock-resistant ventricular fibrillation. *N Engl J Med.* 2002; 346:884–90.
- S5.1.5.2-29.** Levine JH, Massumi A, Scheinman MM, et al. Intravenous amiodarone for recurrent sustained hypotensive ventricular tachyarrhythmias. Intravenous Amiodarone Multicenter Trial Group. *J Am Coll Cardiol.* 1996;27:67–75.
- S5.1.5.2-30.** Epstein AE, Olshansky B, Naccarelli GV, et al. Practical management guide for clinicians who treat patients with amiodarone. *Am J Med.* 2016;129: 468–75.
- S5.1.5.2-31.** Kuhlkamp V, Mewis C, Mermi J, et al. Suppression of sustained ventricular tachyarrhythmias: a comparison of d<sub>1</sub>,l-sotalol with no antiarrhythmic drug treatment. *J Am Coll Cardiol.* 1999;33: 46–52.
- S5.1.5.2-32.** Waldo AL, Camm AJ, deRuyter H, et al. Survival with oral d-sotalol in patients with left ventricular dysfunction after myocardial infarction: rationale, design, and methods (the SWORD trial). *Am J Cardiol.* 1995;75:1023–7.
- S5.1.5.2-33.** Page RL. Effects of antiarrhythmic medication on implantable cardioverter-defibrillator function. *Am J Cardiol.* 2000;85:1481–5.
- S5.1.5.2-34.** Buxton AE, Marchlinski FE, Doherty JU, et al. Hazards of intravenous verapamil for sustained ventricular tachycardia. *Am J Cardiol.* 1987; 59:1107–10.
- S5.1.5.2-35.** Belhassen BHorowitz LN. Use of intravenous verapamil for ventricular tachycardia. *Am J Cardiol.* 1984;54:1131–3.
- S5.1.5.2-36.** Griffith MJ, Garratt CJ, Rowland E, et al. Effects of intravenous adenosine on verapamil-sensitive "idiopathic" ventricular tachycardia. *Am J Cardiol.* 1994;73:759–64.
- S5.1.5.2-37.** Gill JS, Blaszyk K, Ward DE, et al. Verapamil for the suppression of idiopathic ventricular tachycardia of left bundle branch block-like morphology. *Am Heart J.* 1993;126:1126–33.
- S5.1.5.2-38.** Badhwar N, Scheinman MM. Idiopathic ventricular tachycardia: diagnosis and management. *Curr Probl Cardiol.* 2007;32:7–43.
- S5.1.5.2-39.** Gill JS, Ward DE, Camm AJ. Comparison of verapamil and diltiazem in the suppression of idiopathic ventricular tachycardia. *Pacing Clin Electrophysiol.* 1992;15:2122–6.
- S5.1.5.2-40.** Investigators in the Magnesium in Coronaries (MAGIC) Trial. Early administration of intravenous magnesium to high-risk patients with acute myocardial infarction in the Magnesium in Coronaries (MAGIC) Trial: a randomised controlled trial. *Lancet.* 2002;360:1189–96.
- S5.1.5.2-41.** Cooper HA, Dries DL, Davis CE, et al. Diuretics and risk of arrhythmic death in patients with left ventricular dysfunction. *Circulation.* 1999;100: 1311–5.
- S5.1.5.2-42.** Kafka H, Langevin L, Armstrong PW. Serum magnesium and potassium in acute myocardial infarction. Influence on ventricular arrhythmias. *Arch Intern Med.* 1987;147:465–9.
- S5.1.5.2-43.** Yan GX, Antzelevitch C. Cellular basis for the normal T wave and the electrocardiographic manifestations of the long-QT syndrome. *Circulation.* 1998;98:1928–36.
- S5.1.5.2-44.** Tzivoni D, Banai S, Schuger C, et al. Treatment of torsade de pointes with magnesium sulfate. *Circulation.* 1988;77:392–7.
- S5.1.5.2-45.** Leier CV, Dei CL, Metra M. Clinical relevance and management of the major electrolyte abnormalities in congestive heart failure: hyponatremia, hypokalemia, and hypomagnesemia. *Am Heart J.* 1994; 128:564–74.
- S5.1.5.2-46.** Cohn JN, Kowey PR, Whelton PK, et al. New guidelines for potassium replacement in clinical practice: a contemporary review by the National Council on Potassium in Clinical Practice. *Arch Intern Med.* 2000;160:2429–36.
- S5.1.5.2-47.** Macdonald JE, Struthers AD. What is the optimal serum potassium level in cardiovascular patients? *J Am Coll Cardiol.* 2004;43:155–61.
- S5.1.5.2-48.** Goyal A, Spertus JA, Gosch K, et al. Serum potassium levels and mortality in acute myocardial infarction. *JAMA.* 2012;307:157–64.
- S5.1.5.2-49.** Leaf A, Kang JX, Xiao YF, et al. Clinical prevention of sudden cardiac death by n-3 polyunsaturated fatty acids and mechanism of prevention of arrhythmias by n-3 fish oils. *Circulation.* 2003;107: 2646–52.
- S5.1.5.2-50.** Mozaffarian D, Rimm EB. Fish intake, contaminants, and human health: evaluating the risks and the benefits. *JAMA.* 2006;296:1885–99.
- S5.1.5.2-51.** Marchioli R, Barzi F, Bomba E, et al. Early protection against sudden death by n-3 polyunsaturated fatty acids after myocardial infarction: time-course analysis of the results of the Gruppo Italiano per lo Studio della Sopravvivenza nell'Infarto Miocardico (GISSI)-Prevenzione. *Circulation.* 2002; 105:1897–903.
- S5.1.5.2-52.** Kromhout D, Giltay EJ, Geleijnse JM. n-3 fatty acids and cardiovascular events after myocardial infarction. *N Engl J Med.* 2010;363:2015–26.
- S5.1.5.2-53.** Rauch B, Schiele R, Schneider S, et al. OMEGA, a randomized, placebo-controlled trial to test the effect of highly purified omega-3 fatty acids on top

of modern guideline-adjusted therapy after myocardial infarction. *Circulation.* 2010;122:2152-9.

**S5.1.5.2-54.** Burr ML, Ashfield-Watt PA, Dunstan FD, et al. Lack of benefit of dietary advice to men with angina: results of a controlled trial. *Eur J Clin Nutr.* 2003;57:193-200.

**S5.1.5.2-55.** Galan P, Kesse-Guyot E, Czernichow S, et al. Effects of B vitamins and omega 3 fatty acids on cardiovascular diseases: a randomised placebo controlled trial. *BMJ.* 2010;341:c6273.

**S5.1.5.2-56.** Bosch J, Gerstein HC, Dagenais GR, et al. n-3 fatty acids and cardiovascular outcomes in patients with dysglycemia. *N Engl J Med.* 2012;367:309-18.

**S5.1.5.2-57.** LaRosa JC, He J, Yuppertuti S. Effect of statins on risk of coronary disease: a meta-analysis of randomized controlled trials. *JAMA.* 1999;282:2340-6.

**S5.1.5.2-58.** Buber J, Goldenberg I, Moss AJ, et al. Reduction in life-threatening ventricular tachyarrhythmias in statin-treated patients with nonischemic cardiomyopathy enrolled in the MADIT-CRT (Multi-center Automatic Defibrillator Implantation Trial with Cardiac Resynchronization Therapy). *J Am Coll Cardiol.* 2012;60:749-55.

**S5.1.5.2-59.** Investigators in TAVI/DA. A comparison of antiarrhythmic-drug therapy with implantable defibrillators in patients resuscitated from near-fatal ventricular arrhythmias. *N Engl J Med.* 1997;337:1576-83.

**S5.1.5.2-60.** Mitchell LB, Powell JL, Gillis AM, et al. Are lipid-lowering drugs also antiarrhythmic drugs? An analysis of the Antiarrhythmics versus Implantable Defibrillators (AVID) trial. *J Am Coll Cardiol.* 2003;42:81-7.

**S5.1.5.2-61.** Dickinson MG, Ip JH, Olshansky B, et al. Statin use was associated with reduced mortality in both ischemic and nonischemic cardiomyopathy and in patients with implantable defibrillators: mortality data and mechanistic insights from the Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT). *Am Heart J.* 2007;153:573-8.

**S5.1.5.2-62.** Goldberger JJ, Subacius H, Schaechter A, et al. Effects of statin therapy on arrhythmic events and survival in patients with nonischemic dilated cardiomyopathy. *J Am Coll Cardiol.* 2006;48:1228-33.

**S5.1.5.2-63.** Kjekshus J, Apetrei E, Barrios V, et al. Rosuvastatin in older patients with systolic heart failure. *N Engl J Med.* 2007;357:2248-61.

**S5.1.5.2-64.** Tavazzi L, Maggioni AP, Marchioli R, et al. Effect of rosuvastatin in patients with chronic heart failure (the GISSI-HF trial): a randomised, double-blind, placebo-controlled trial. *Lancet.* 2008;372:1231-9.

## 5.2. Preventing SCD With HF Medications

**S5.2-1.** Yancy CW, Jessup M, Bozkurt B, et al. 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: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Failure Society of America. *J Am Coll Cardiol.* 2016;68:1476-88.

**S5.2-2.** Committees for Cardiac insufficiency Bisoprolol Study II. The cardiac insufficiency Bisoprolol Study II (CIBIS-II): a randomised trial. *Lancet.* 1999;353:9-13.

**S5.2-3.** Cohn JN, Johnson G, Ziesche S, et al. A comparison of enalapril with hydralazine-isosorbide dinitrate in the treatment of chronic congestive heart failure. *N Engl J Med.* 1991;325:303-10.

**S5.2-4.** Packer M, Bristow MR, Cohn JN, et al. The effect of carvedilol on morbidity and mortality in patients with chronic heart failure. *U.S. Carvedilol Heart Failure Study Group.* *N Engl J Med.* 1996;334:1349-55.

**S5.2-5.** Dargie HJ. Effect of carvedilol on outcome after myocardial infarction in patients with left-ventricular dysfunction: the CAPRICORN randomised trial. *Lancet.* 2001;357:1385-90.

**S5.2-6.** Pitt B, Remme W, Zannad F, et al. Eplerenone, a selective aldosterone blocker, in patients with left ventricular dysfunction after myocardial infarction. *N Engl J Med.* 2003;348:1309-21.

**S5.2-7.** Al Chekakie MO. Traditional heart failure medications and sudden cardiac death prevention: a review. *J Cardiovasc Pharmacol Ther.* 2013;18:412-26.

**S5.2-8.** Pfeffer MA, McMurray JJ, Velazquez EJ, et al. Valsartan, captopril, or both in myocardial infarction complicated by heart failure, left ventricular dysfunction, or both. *N Engl J Med.* 2003;349:1893-906.

**S5.2-9.** Hjalmarson A, Goldstein S, Fagerberg B, et al. Effects of controlled-release metoprolol on total mortality, hospitalizations, and well-being in patients with heart failure: the Metoprolol CR/XL Randomized Intervention Trial in congestive heart failure (MERIT-HF). *MERIT-HF Study Group.* *JAMA.* 2000;283:1295-302.

**S5.2-10.** McMurray J JV, Östergren J, Swedberg K, et al. Effects of candesartan in patients with chronic heart failure and reduced left-ventricular systolic function taking angiotensin-converting-enzyme inhibitors: the CHARM-Added trial. *Lancet.* 2003;362:767-71.

**S5.2-11.** Cohn JN, Tognoni G. A randomized trial of the angiotensin-receptor blocker valsartan in chronic heart failure. *N Engl J Med.* 2001;345:1667-75.

**S5.2-12.** Pitt B, Zannad F, Remme WJ, et al. The effect of spironolactone on morbidity and mortality in patients with severe heart failure. Randomized Aldactone Evaluation Study Investigators. *N Engl J Med.* 1999;341:709-17.

**S5.2-13.** Zannad F, McMurray JJ, Krum H, et al. Eplerenone in patients with systolic heart failure and mild symptoms. *N Engl J Med.* 2011;364:11-21.

**S5.2-14.** Desai AS, McMurray JJ, Packer M, et al. Effect of the angiotensin-receptor-neprilysin inhibitor LCZ696 compared with enalapril on mode of death in heart failure patients. *Eur Heart J.* 2015;36:1990-7.

## 5.4. Catheter Ablation

**S5.4.3-1.** Zeppenfeld K, Schalij MJ, Bartelings MM, et al. Catheter ablation of ventricular tachycardia after repair of congenital heart disease: electroanatomic identification of the critical right ventricular isthmus. *Circulation.* 2007;116:2241-52.

**S5.4.3-2.** Priori SG, Blomstrom-Lundqvist C, Mazzanti A, et al. 2015 ESC guidelines for the management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: the Task Force for the Management of Patients with Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death of the European Society of Cardiology (ESC). *Eur Heart J.* 2015;36:2793-867.

**S5.4.3-3.** Aliot EM, Stevenson WG, Almendral-Garrote JM, et al. EHRA/HRS expert consensus on catheter ablation of ventricular arrhythmias: developed in a partnership with the European Heart Rhythm Association (EHRA), a Registered Branch of the European Society of Cardiology (ESC), and the Heart Rhythm Society (HRS); in collaboration with the American College of Cardiology (ACC) and the American Heart Association (AHA). *Heart Rhythm.* 2009;6:886-933.

## 5.5. Surgery and Revascularization Procedures in Patients With Ischemic Heart Disease

**S5.5-1.** Cook JR, Rizo-Patron C, Curtis AB, et al. Effect of surgical revascularization in patients with coronary artery disease and ventricular tachycardia or fibrillation in the Antiarrhythmics Versus Implantable Defibrillators (AVID) Registry. *Am Heart J.* 2002;143:821-6.

**S5.5-2.** Velazquez EJ, Lee KL, Jones RH, et al. Coronary-artery bypass surgery in patients with ischemic cardiomyopathy. *N Engl J Med.* 2016;374:1511-20.

**S5.5-3.** Every NR, Fahrenbruch CE, Hallstrom AP, et al. Influence of coronary bypass surgery on subsequent outcome of patients resuscitated from out-of-hospital cardiac arrest. *J Am Coll Cardiol.* 1992;19:1435-9.

**S5.5-4.** Dumas F, Bougouin W, Geri G, et al. Emergency PCI in post-cardiac arrest patients without ST-segment elevation pattern: insights from the PROCAT II registry. *J Am Coll Cardiol Intv.* 2016;9:1011-8.

**S5.5-5.** Ngaage DL, Cale AR, Cowen ME, et al. Early and late survival after surgical revascularization for ischemic ventricular fibrillation/tachycardia. *Ann Thorac Surg.* 2008;85:1278-81.

**S5.5-6.** Milojevic M, Head SJ, Parasca CA, et al. Causes of death following PCI versus CABG in complex CAD: 5-year follow-up of SYNTAX. *J Am Coll Cardiol.* 2016;67:42-52.

**S5.5-7.** Carson P, Wertheimer J, Miller A, et al. The STICH trial (Surgical Treatment for Ischemic Heart Failure): mode-of-death results. *JACC Heart Fail.* 2013;1:400-8.

**S5.5-8.** Davis JA, Cecchin F, Jones TK, et al. Major coronary artery anomalies in a pediatric population: incidence and clinical importance. *J Am Coll Cardiol.* 2001;37:593-7.

**S5.5-9.** Bagnall RD, Weintraub RG, Ingles J, et al. A prospective study of sudden cardiac death among children and young adults. *N Engl J Med.* 2016;374:2441-52.

**S5.5-10.** Smallman DP, Webber BJ, Mazuchowski EL, et al. Sudden cardiac death associated with physical exertion in the US military, 2005-2010. *BJ Sports Med.* 2016;50:118-23.

**S5.5-11.** Maron BJ, Doerer JJ, Haas TS, et al. Sudden deaths in young competitive athletes: analysis of 1866 deaths in the United States, 1980-2006. *Circulation.* 2009;119:1085-92.

**S5.5-12.** Krasuski RA, Magyar D, Hart S, et al. Long-term outcome and impact of surgery on adults with coronary arteries originating from the opposite coronary cusp. *Circulation.* 2011;123:154-62.

**S5.5-13.** Layser RB, Savage MP, Halpern EJ. Anomalous coronary arteries: analysis of clinical outcome based upon arterial course and surgical intervention in an adult population. *Acad Radiol.* 2016;23:1015-23.

**S5.5-14.** Taylor AJ, Rogan KM, Virmani R. Sudden cardiac death associated with isolated congenital coronary artery anomalies. *J Am Coll Cardiol.* 1992;20:640-7.

#### 5.5.1. Surgery for Arrhythmia Management

**S5.5.1-1.** Anter E, Hutchinson MD, Deo R, et al. Surgical ablation of refractory ventricular tachycardia in patients with nonischemic cardiomyopathy. *Circ Arrhythm Electrophysiol.* 2011;4:494-500.

**S5.5.1-2.** Bhavani SS, Tchou P, Saliba W, et al. Surgical options for refractory ventricular tachycardia. *J Card Surg.* 2007;22:533-4.

**S5.5.1-3.** Choi EK, Nagashima K, Lin KY, et al. Surgical cryoablation for ventricular tachyarrhythmia arising from the left ventricular outflow tract region. *Heart Rhythm.* 2015;12:1128-36.

**S5.5.1-4.** Kumar S, Barbhaiya CR, Sobieszczyk P, et al. Role of alternative interventional procedures when endo- and epicardial catheter ablation attempts for ventricular arrhythmias fail. *Circ Arrhythm Electrophysiol.* 2015;8:606-15.

**S5.5.1-5.** Mulloy DP, Bhamidipati CM, Stone ML, et al. Cryoablation during left ventricular assist device implantation reduces postoperative ventricular tachyarrhythmias. *J Thorac Cardiovasc Surg.* 2013;145:1207-13.

**S5.5.1-6.** Patel M, Rojas F, Shabari FR, et al. Safety and feasibility of open chest epicardial mapping and ablation of ventricular tachycardia during the period of left ventricular assist device implantation. *J Cardiovasc Electrophysiol.* 2016;27:95-101.

**S5.5.1-7.** Sartipy U, Albage A, Straat E, et al. Surgery for ventricular tachycardia in patients undergoing left ventricular reconstruction by the Dor procedure. *Ann Thorac Surg.* 2006;81:65-71.

#### 5.6. Autonomic Modulation

**S5.6-1.** Kittayaphong R, Bhuripanyo K, Punlee K, et al. Effect of atenolol on symptomatic ventricular arrhythmia without structural heart disease: a randomized placebo-controlled study. *Am Heart J.* 2002;144:e10.

**S5.6-2.** Vaseghi M, Barwad P, Malavassi Corrales FJ, et al. Cardiac sympathetic denervation for refractory ventricular arrhythmias. *J Am Coll Cardiol.* 2017;69:3070-80.

**S5.6-3.** Vaseghi M, Gima J, Kanaan C, et al. Cardiac sympathetic denervation in patients with refractory ventricular arrhythmias or electrical storm: intermediate and long-term follow-up. *Heart Rhythm.* 2014;11:360-6.

**S5.6-4.** Schwartz PJ, Motolese M, Pollavini G. Prevention of sudden cardiac death after a first myocardial infarction by pharmacologic or surgical antiadrenergic interventions. *J Cardiovasc Electrophysiol.* 1992;3:2-16.

**S5.6-5.** Shen MJ, Zipes DP. Role of the autonomic nervous system in modulating cardiac arrhythmias. *Circ Res.* 2014;114:1004-21.

**S5.6-6.** Ajijola OA, Lellouche N, Bourke T, et al. Bilateral cardiac sympathetic denervation for the management of electrical storm. *J Am Coll Cardiol.* 2012;59:91-2.

#### 6. ACUTE MANAGEMENT OF SPECIFIC VA

**S6-1.** Link MS, Berkow LC, Kudenchuk PJ, et al. Part 7: adult advanced cardiovascular life support. *2015*

American Heart Association guidelines update for cardiopulmonary resuscitation and emergency cardiovascular care. *Circulation.* 2015;132 suppl 2:S444-64.

**S6-2.** Stiell IG, Wells GA, Field B, et al. Advanced cardiac life support in out-of-hospital cardiac arrest. *N Engl J Med.* 2004;351:647-56.

**S6-3.** Sisson C, Rogers MA, Dahl J, et al. Predictors of survival from out-of-hospital cardiac arrest: a systematic review and meta-analysis. *Circ Cardiovasc Qual Outcomes.* 2010;3:63-81.

**S6-4.** Dorian P, Cass D, Schwartz B, et al. Amiodarone as compared with lidocaine for shock-resistant ventricular fibrillation. *N Engl J Med.* 2002;346:884-90.

**S6-5.** Kudenchuk PJ, Brown SP, Daya M, et al. Amiodarone, lidocaine, or placebo in out-of-hospital cardiac arrest. *N Engl J Med.* 2016;371:1711-22.

**S6-6.** Kudenchuk PJ, Cobb LA, Copass MK, et al. Amiodarone for resuscitation after out-of-hospital cardiac arrest due to ventricular fibrillation. *N Engl J Med.* 1999;341:871-8.

**S6-7.** Spaulding CM, Joly LM, Rosenberg A, et al. Immediate coronary angiography in survivors of out-of-hospital cardiac arrest. *N Engl J Med.* 1997;336:1629-33.

**S6-8.** Cronier P, Vignon P, Bouferrache K, et al. Impact of routine percutaneous coronary intervention after out-of-hospital cardiac arrest due to ventricular fibrillation. *Crit Care.* 2011;15:R122.

**S6-9.** Zanuttini D, Armellini I, Nucifora G, et al. Impact of emergency coronary angiography on in-hospital outcome of unconscious survivors after out-of-hospital cardiac arrest. *Am J Cardiol.* 2012;110:1723-8.

**S6-10.** Dumas F, Bougouin W, Geri G, et al. Emergency PCI in post-cardiac arrest patients without ST-segment elevation pattern: insights from the PROCAT II registry. *J Am Coll Cardiol Intv.* 2016;9:1011-8.

**S6-11.** Gorgels AP, van den Dool A, Hofs A, et al. Comparison of procainamide and lidocaine in terminating sustained monomorphic ventricular tachycardia. *Am J Cardiol.* 1996;78:43-6.

**S6-12.** Markel DT, Gold LS, Allen J, et al. Procainamide and survival in ventricular fibrillation out-of-hospital cardiac arrest. *Acad Emerg Med.* 2010;17:617-23.

**S6-13.** Ortiz M, Martin A, Arribas F, et al. Randomized comparison of intravenous procainamide vs. intravenous amiodarone for the acute treatment of tolerated wide QRS tachycardia: the PROCAMIO study. *Eur Heart J.* 2017;38:1329-35.

**S6-14.** Herlitz J, Ekstrom L, Wennerblom B, et al. Lidocaine in out-of-hospital ventricular fibrillation. Does it improve survival? *Resuscitation.* 1997;33:199-205.

**S6-15.** Kudenchuk PJ, Newell C, White L, et al. Prophylactic lidocaine for post resuscitation care of patients with out-of-hospital ventricular fibrillation cardiac arrest. *Resuscitation.* 2013;84:1512-8.

**S6-16.** Teo KK, Yusuf S, Furberg CD. Effects of prophylactic antiarrhythmic drug therapy in acute myocardial infarction. An overview of results from randomized controlled trials. *JAMA.* 1993;270:1589-95.

**S6-17.** Nademanee K, Taylor R, Bailey WE, et al. Treating electrical storm: sympathetic blockade versus advanced cardiac life support-guided therapy. *Circulation.* 2000;102:742-7.

**S6-18.** Piccini JP, Hranitzky PM, Kilaru R, et al. Relation of mortality to failure to prescribe beta blockers

acutely in patients with sustained ventricular tachycardia and ventricular fibrillation following acute myocardial infarction (from the VALsartan In Acute myocardial infarction trial [VALIANT] Registry). *Am J Cardiol.* 2008;102:1427-32.

**S6-19.** Callaham M, Madsen CD, Barton CW, et al. A randomized clinical trial of high-dose epinephrine and norepinephrine vs standard-dose epinephrine in prehospital cardiac arrest. *JAMA.* 1992;268:2667-72.

**S6-20.** Donnino MW, Salciccioli JD, Howell MD, et al. Time to administration of epinephrine and outcome after in-hospital cardiac arrest with non-shockable rhythms: retrospective analysis of large in-hospital data registry. *BMJ.* 2014;348:g3028.

**S6-21.** Gueugniaud PY, Mols P, Goldstein P, et al. A comparison of repeated high doses and repeated standard doses of epinephrine for cardiac arrest outside the hospital. *European Epinephrine Study Group.* *N Engl J Med.* 1998;339:1595-601.

**S6-22.** Hagiwara A, Hasegawa M, Abe T, et al. Pre-hospital epinephrine use and survival among patients with out-of-hospital cardiac arrest. *JAMA.* 2012;307:1161-8.

**S6-23.** Jacobs IG, Finn JC, Jelinek GA, et al. Effect of adrenaline on survival in out-of-hospital cardiac arrest: a randomised double-blind placebo-controlled trial. *Resuscitation.* 2011;82:1138-43.

**S6-24.** Koscik C, Pinawin A, McGovern H, et al. Rapid epinephrine administration improves early outcomes in out-of-hospital cardiac arrest. *Resuscitation.* 2013;84:915-20.

**S6-25.** Ho DS, Zecchin RP, Richards DA, et al. Double-blind trial of lignocaine versus sotalol for acute termination of spontaneous sustained ventricular tachycardia. *Lancet.* 1994;344:18-23.

**S6-26.** Somberg JC, Bailin SJ, Haffajee CI, et al. Intravenous lidocaine versus intravenous amiodarone (in a new aqueous formulation) for incessant ventricular tachycardia. *Am J Cardiol.* 2002;90:853-9.

**S6-27.** Hassan TB, Jagger C, Barnett DB. A randomised trial to investigate the efficacy of magnesium sulphate for refractory ventricular fibrillation. *Emerg Med J.* 2002;19:57-62.

**S6-28.** Thel MC, Armstrong AL, McNulty SE, et al. Randomised trial of magnesium in in-hospital cardiac arrest. *Duke Internal Medicine Housestaff.* *Lancet.* 1997;350:1272-6.

**S6-29.** Elizari MV, Martinez JM, Belziti C, et al. Morbidity and mortality following early administration of amiodarone in acute myocardial infarction. *GEMICA study investigators, GEMA Group, Buenos Aires, Argentina. Grupo de Estudios Multicentricos en Argentina.* *Eur Heart J.* 2000;21:198-205.

**S6-30.** Behlissen B, Horowitz LN. Use of intravenous verapamil for ventricular tachycardia. *Am J Cardiol.* 1984;54:1131-3.

**S6-31.** Buxton AE, Marchlinski FE, Doherty JU, et al. Hazards of intravenous verapamil for sustained ventricular tachycardia. *Am J Cardiol.* 1987;59:1107-10.

**S6-32.** Zipes DP, Camm AJ, Borggrefe M, et al. ACC/AHA/ESC 2006 guidelines for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology/American Heart Association Task Force and the European Society of

Cardiology Committee for Practice Guidelines. *J Am Coll Cardiol.* 2006;48:e247-346.

**S6-33.** Cobb LA, Fahrenbruch CE, Walsh TR, et al. Influence of cardiopulmonary resuscitation prior to defibrillation in patients with out-of-hospital ventricular fibrillation. *JAMA.* 1999;281:1182-8.

**S6-34.** Cobb LA, Weaver WD, Fahrenbruch CE, et al. Community-based interventions for sudden cardiac death. Impact, limitations, and changes. *Circulation.* 1992;85 suppl:I98-102.

**S6-35.** Weisfeldt ML, Becker LB. Resuscitation after cardiac arrest: a 3-phase time-sensitive model. *JAMA.* 2002;288:3035-8.

**S6-36.** Wik L, Hansen TB, Fylling F, et al. Delaying defibrillation to give basic cardiopulmonary resuscitation to patients with out-of-hospital ventricular fibrillation: a randomized trial. *JAMA.* 2003;289:1389-95.

**S6-37.** Neumar RW, Otto CW, Link MS, et al. Part 8: adult advanced cardiovascular life support: 2010 American Heart Association guidelines for cardiopulmonary resuscitation and emergency cardiovascular care. *Circulation.* 2010;122 suppl 2:S729-67.

**S6-38.** Connolly SJ, Hallstrom AP, Cappato R, et al. Meta-analysis of the implantable cardioverter defibrillator secondary prevention trials. AVID, CASH and CIDS studies. Antiarrhythmics vs implantable defibrillator study. Cardiac Arrest Study Hamburg. Canadian Implantable Defibrillator Study. *Eur Heart J.* 2000;21:2071-8.

**S6-39.** Kern KB. Optimal treatment of patients surviving out-of-hospital cardiac arrest. *J Am Coll Cardiol Intv.* 2012;5:597-605.

**S6-40.** Perkins GD, Jacobs IG, Nadkarni VM, et al. Cardiac arrest and cardiopulmonary resuscitation outcome reports: update of the Utstein Resuscitation Registry Templates for out-of-hospital cardiac arrest: a statement for healthcare professionals from a task force of the International Liaison Committee on Resuscitation (American Heart Association, European Resuscitation Council, Australian and New Zealand Council on Resuscitation, Heart and Stroke Foundation of Canada, InterAmerican Heart Foundation, Resuscitation Council of Southern Africa, Resuscitation Council of Asia); and the American Heart Association Emergency Cardiovascular Care Committee and the Council on Cardiopulmonary, Critical Care, Perioperative and Resuscitation. *Circulation.* 2015;132:1286-300.

**S6-41.** Link MS, Myerburg RJ, Estes NA 3rd. Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities: Task Force 12: emergency action plans, resuscitation, cardiopulmonary resuscitation, and automated external defibrillators: a scientific statement from the American Heart Association and American College of Cardiology. *Circulation.* 2015;132:e334-8.

**S6-42.** Joglar JA, Page RL. Out-of-hospital cardiac arrest—are drugs ever the answer? *N Engl J Med.* 2016;374:1781-2.

## 7. ONGOING MANAGEMENT OF VA AND SCD RISK RELATED TO SPECIFIC DISEASE STATES

### 7.1. Ischemic Heart Disease

#### 7.1.1. Secondary Prevention of SCD in Patients With Ischemic Heart Disease

**S7.1.1-1.** The AVID Investigators. A comparison of antiarrhythmic-drug therapy with implantable

defibrillators in patients resuscitated from near-fatal ventricular arrhythmias. *N Engl J Med.* 1997;337:1576-83.

**S7.1.1-2.** Connolly SJ, Gent M, Roberts RS, et al. Canadian implantable defibrillator study (CIDS): a randomized trial of the implantable cardioverter defibrillator against amiodarone. *Circulation.* 2000;101:1297-302.

**S7.1.1-3.** Connolly SJ, Hallstrom AP, Cappato R, et al. Meta-analysis of the implantable cardioverter defibrillator secondary prevention trials. AVID, CASH and CIDS studies. Antiarrhythmics vs Implantable Defibrillator study. Cardiac Arrest Study Hamburg. Canadian Implantable Defibrillator Study. *Eur Heart J.* 2000;21:2071-8.

**S7.1.1-4.** Kuck KH, Cappato R, Siebels J, et al. Randomized comparison of antiarrhythmic drug therapy with implantable defibrillators in patients resuscitated from cardiac arrest: the Cardiac Arrest Study Hamburg (CASH). *Circulation.* 2000;102:748-54.

**S7.1.1-5.** Raith MH, Renfroe EG, Epstein AE, et al. "Stable" ventricular tachycardia is not a benign rhythm: insights from the antiarrhythmics versus implantable defibrillators (AVID) registry. *Circulation.* 2001;103:244-52.

**S7.1.1-6.** Owens DK, Sanders GD, Heidenreich PA, et al. Effect of risk stratification on cost-effectiveness of the implantable cardioverter defibrillator. *Am Heart J.* 2002;144:440-8.

**S7.1.1-7.** Bass EB, Elson JJ, Fogoros RN, et al. Long-term prognosis of patients undergoing electrophysiologic studies for syncope of unknown origin. *Am J Cardiol.* 1988;62:1186-91.

**S7.1.1-8.** Wyse DG, Friedman PL, Brodsky MA, et al. Life-threatening ventricular arrhythmias due to transient or correctable causes: high risk for death in follow-up. *J Am Coll Cardiol.* 2001;38:1718-24.

**S7.1.1-9.** Maury P, Baratto F, Zeppenfeld K, et al. Radio-frequency ablation as primary management of well-tolerated sustained monomorphic ventricular tachycardia in patients with structural heart disease and left ventricular ejection fraction over 30%. *Eur Heart J.* 2014;35:1479-85.

**S7.1.1-10.** Paurish M, Cismaru G, Magnin-Poull I, et al. A stepwise approach to the management of postinfarct ventricular tachycardia using catheter ablation as the first-line treatment: a single-center experience. *Circ Arrhythm Electrophysiol.* 2013;6:351-6.

**S7.1.1-11.** Larsen G, Hallstrom A, McAnulty J, et al. Cost-effectiveness of the implantable cardioverter-defibrillator versus antiarrhythmic drugs in survivors of serious ventricular tachyarrhythmias: results of the Antiarrhythmics Versus Implantable Defibrillators (AVID) economic analysis substudy. *Circulation.* 2002;105:2049-57.

**S7.1.1-12.** O'Brien BJ, Connolly SJ, Goeree R, et al. Cost-effectiveness of the implantable cardioverter-defibrillator: results from the Canadian Implantable Defibrillator Study (CIDS). *Circulation.* 2001;103:1416-21.

**S7.1.1-13.** Buxton M, Caine N, Chase D, et al. A review of the evidence on the effects and costs of implantable cardioverter defibrillator therapy in different patient groups, and modelling of cost-effectiveness and cost-utility for these groups in a UK context. *Health Technol Assess.* 2006;10:iii-iv, ix-xi, 1-164.

**S7.1.1-14.** Weiss JP, Saynina O, McDonald KM, et al. Effectiveness and cost-effectiveness of implantable cardioverter defibrillators in the treatment of ventricular arrhythmias among medicare beneficiaries. *Am J Med.* 2002;112:519-27.

**S7.1.1-15.** Anderson JL, Heidenreich PA, Barnett PG, et al. ACC/AHA statement on cost/value methodology in clinical practice guidelines and performance measures: a report of the American College of Cardiology/American Heart Association Task Force on Performance Measures and Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;63:2304-22.

#### 7.1.1.1. Coronary Artery Spasm

**S7.1.1-1-1.** Chevalier P, Dacosta A, Defaye P, et al. Arrhythmic cardiac arrest due to isolated coronary artery spasm: long-term outcome of seven resuscitated patients. *J Am Coll Cardiol.* 1998;31:57-61.

**S7.1.1-1-2.** Myerburg RJ, Kessler KM, Mallon SM, et al. Life-threatening ventricular arrhythmias in patients with silent myocardial ischemia due to coronary-artery spasm. *N Engl J Med.* 1992;326:1451-5.

**S7.1.1-1-3.** Ahn JM, Lee KH, Yoo SY, et al. Prognosis of variant angina manifesting as aborted sudden cardiac death. *J Am Coll Cardiol.* 2016;68:137-45.

**S7.1.1-1-4.** Matsue Y, Suzuki M, Nishizaki M, et al. Clinical implications of an implantable cardioverter-defibrillator in patients with vasospastic angina and lethal ventricular arrhythmia. *J Am Coll Cardiol.* 2012;60:908-13.

**S7.1.1-1-5.** Takagi Y, Yasuda S, Tsunoda R, et al. Clinical characteristics and long-term prognosis of vasospastic angina patients who survived out-of-hospital cardiac arrest: multicenter registry study of the Japanese Coronary Spasm Association. *Circ Arrhythm Electrophysiol.* 2011;4:295-302.

**S7.1.1-1-6.** Meisel SR, Mazur A, Chetboun I, et al. Usefulness of implantable cardioverter-defibrillators in refractory variant angina pectoris complicated by ventricular fibrillation in patients with angiographically normal coronary arteries. *Am J Cardiol.* 2002;89:1114-6.

**S7.1.1-1-7.** JCS Joint Working Group. Guidelines for diagnosis and treatment of patients with vasospastic angina (coronary spastic angina) (JCS 2008). *Circ J.* 2010;74:1745-62.

**S7.1.1-1-8.** Amsterdam EA, Wenger NK, Brindis RG, et al. 2014 AHA/ACC guideline for the management of patients with non-ST-elevation acute coronary syndromes: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;64:e139-228.

**S7.1.1-1-9.** Morikawa Y, Mizuno Y, Yasue H. Letter by Morikawa et al regarding article, "coronary artery spasm: a 2009 update". *Circulation.* 2010;121:e16.

**S7.1.1-1-10.** Sasaki S, Tomita H, Shibusaki S, et al. Usefulness of the wearable cardioverter-defibrillator in patients at high risk for sudden cardiac death. *Circ J.* 2014;78:2987-9.

#### 7.1.1.2. Post CABG VT/VF

**S7.1.1-2-1.** Ascione R, Reeves BC, Santo K, et al. Predictors of new malignant ventricular arrhythmias after coronary surgery: a case-control study. *J Am Coll Cardiol.* 2004;43:1630-8.

**S7.1.1.2-2.** Saxon LA, Wiener I, Natterson PD, et al. Monomorphic versus polymorphic ventricular tachycardia after coronary artery bypass grafting. *Am J Cardiol.* 1995;75:403–5.

**S7.1.1.2-3.** Steinberg JS, Gaur A, Sciacca R, et al. New-onset sustained ventricular tachycardia after cardiac surgery. *Circulation.* 1999;99:903–8.

**S7.1.1.2-4.** Bigger JT Jr. Prophylactic use of implanted cardiac defibrillators in patients at high risk for ventricular arrhythmias after coronary-artery bypass graft surgery. Coronary Artery Bypass Graft (CABG) Patch Trial Investigators. *N Engl J Med.* 1997;337:1569–75.

**S7.1.1.2-5.** Vakil K, Florea V, Koene R, et al. Effect of coronary artery bypass grafting on left ventricular ejection fraction in men eligible for implantable cardioverter-defibrillator. *Am J Cardiol.* 2016;117:957–60.

**S7.1.1.2-6.** Mittal S, Lomnitz DJ, Mirchandani S, et al. Prognostic significance of nonsustained ventricular tachycardia after revascularization. *J Cardiovasc Electrophysiol.* 2002;13:342–6.

**S7.1.1.2-7.** Zishiri ET, Williams S, Cronin EM, et al. Early risk of mortality after coronary artery revascularization in patients with left ventricular dysfunction and potential role of the wearable cardioverter defibrillator. *Circ Arrhythm Electrophysiol.* 2013;6:117–28.

#### 7.1.2. Primary Prevention of SCD in Patients With Ischemic Heart Disease

**S7.1.2-1.** Bardy GH, Lee KL, Mark DB, et al. Amiodarone or an implantable cardioverter-defibrillator for congestive heart failure. *N Engl J Med.* 2005;352:225–37.

**S7.1.2-2.** Moss AJ, Zareba W, Hall WJ, et al. Prophylactic implantation of a defibrillator in patients with myocardial infarction and reduced ejection fraction. *N Engl J Med.* 2002;346:877–83.

**S7.1.2-3.** Moss AJ, Hall WJ, Cannom DS, et al. Improved survival with an implanted defibrillator in patients with coronary disease at high risk for ventricular arrhythmia. Multicenter Automatic Defibrillator Implantation Trial Investigators. *N Engl J Med.* 1996;335:1933–40.

**S7.1.2-4.** Owens DK, Sanders GD, Heidenreich PA, et al. Effect of risk stratification on cost-effectiveness of the implantable cardioverter defibrillator. *Am Heart J.* 2002;144:440–8.

**S7.1.2-5.** Buxton AE, Lee KL, Fisher JD, et al. A randomized study of the prevention of sudden death in patients with coronary artery disease. Multicenter Unstabilized Tachycardia Trial Investigators. *N Engl J Med.* 1999;341:1882–90.

**S7.1.2-6.** Cantero-Pérez EM, Sobrino-Márquez JM, Grande-Trillo A, et al. Implantable cardioverter defibrillator for primary prevention in patients with severe ventricular dysfunction awaiting heart transplantation. *Transplant Proc.* 2013;45:3659–61.

**S7.1.2-7.** Frohlich GM, Holzmeister J, Hubler M, et al. Prophylactic implantable cardioverter defibrillator treatment in patients with end-stage heart failure awaiting heart transplantation. *Heart.* 2013;99:1158–65.

**S7.1.2-8.** Gajdabekh E, Rovani M, Varnous S, et al. Implantable cardioverter-defibrillators in end-stage heart failure patients listed for heart transplantation:

results from a large retrospective registry. *Arch Cardiovasc Dis.* 2016;109:476–85.

**S7.1.2-9.** Vakil K, Duval S, Cogswell R, et al. Impact of implantable cardioverter-defibrillators on waitlist mortality among patients awaiting heart transplantation: an UNOS/OPN analysis. *JACC Clin Electrophysiol.* 2017;3:33–40.

**S7.1.2-10.** Hohnloser SH, Kuck KH, Dorian P, et al. Prophylactic use of an implantable cardioverter-defibrillator after acute myocardial infarction. *N Engl J Med.* 2004;351:2481–8.

**S7.1.2-11.** Steinbeck G, Andresen D, Seidl K, et al. Defibrillator implantation early after myocardial infarction. *N Engl J Med.* 2009;361:1427–36.

**S7.1.2-12.** Mushlin AI, Hall WJ, Zwanziger J, et al. The cost-effectiveness of automatic implantable cardiac defibrillators: results from MADIT. Multicenter Automatic Defibrillator Implantation Trial. *Circulation.* 1998;97:2129–35.

**S7.1.2-13.** Zwanziger J, Hall WJ, Dick AW, et al. The cost-effectiveness of implantable cardioverter-defibrillators: results from the Multicenter Automatic Defibrillator Implantation Trial (MADIT)-II. *J Am Coll Cardiol.* 2006;47:2310–8.

**S7.1.2-14.** Mark DB, Nelson CL, Anstrom KJ, et al. Cost-effectiveness of defibrillator therapy or amiodarone in chronic stable heart failure: results from the Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT). *Circulation.* 2006;114:135–42.

**S7.1.2-15.** Al-Khatib SM, Anstrom KJ, Eisenstein EL, et al. Clinical and economic implications of the Multicenter Automatic Defibrillator Implantation Trial-II. *Ann Intern Med.* 2005;142:593–600.

**S7.1.2-16.** Buxton AE, Calkins H, Callans DJ, et al. ACC/AHA/HRS 2006 key data elements and definitions for electrophysiological studies and procedures: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Data Standards. *J Am Coll Cardiol.* 2006;48:2360–96.

**S7.1.2-17.** Cowie MR, Marshall D, Drummond M, et al. Lifetime cost-effectiveness of prophylactic implantation of a cardioverter defibrillator in patients with reduced left ventricular systolic function: results of Markov modelling in a European population. *Europace.* 2009;11:716–26.

**S7.1.2-18.** Sanders GD, Hlatky MA, Owens DK. Cost-effectiveness of implantable cardioverter-defibrillators. *N Engl J Med.* 2005;353:1471–80.

**S7.1.2-19.** Smith T, Jordae L, Theuns DA, et al. The cost-effectiveness of primary prophylactic implantable defibrillator therapy in patients with ischaemic or non-ischaemic heart disease: a European analysis. *Eur Heart J.* 2013;34:211–9.

**S7.1.2-20.** Anderson JL, Heidenreich PA, Barnett PG, et al. ACC/AHA statement on cost/value methodology in clinical practice guidelines and performance measures: a report of the American College of Cardiology/American Heart Association Task Force on Performance Measures and Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;63:2304–22.

**S7.1.2-21.** Buxton AE, Lee KL, DiCarlo L, et al. Electrophysiologic testing to identify patients with coronary artery disease who are at risk for sudden death. Multicenter Unstabilized

Tachycardia Trial Investigators. *N Engl J Med.* 2000;342:1937–45.

**S7.1.2-22.** Zaman S, Sivagangabalan G, Narayan A, et al. Outcomes of early risk stratification and targeted implantable cardioverter-defibrillator implantation after ST-elevation myocardial infarction treated with primary percutaneous coronary intervention. *Circulation.* 2009;120:194–200.

**S7.1.2-23.** Bristow MR, Saxon LA, Boehmer J, et al. Cardiac-resynchronization therapy with or without an implantable defibrillator in advanced chronic heart failure. *N Engl J Med.* 2004;350:2140–50.

**S7.1.2-24.** Kusumoto FM, Calkins H, Boehmer J, et al. HRS/ACC/AHA expert consensus statement on the use of implantable cardioverter-defibrillator therapy in patients who are not included or not well represented in clinical trials. *J Am Coll Cardiol.* 2014;64:1143–77.

#### 7.1.3. Treatment and Prevention of Recurrent VA in Patients With Ischemic Heart Disease

**S7.1.3-1.** Connolly SJ, Dorian P, Roberts RS, et al. Comparison of beta-blockers, amiodarone plus beta-blockers, or sotalol for prevention of shocks from implantable cardioverter defibrillators: the OPTIC Study: a randomized trial. *JAMA.* 2006;295:165–71.

**S7.1.3-2.** Pacifico A, Hohnloser SH, Williams JH, et al. Prevention of implantable-defibrillator shocks by treatment with sotalol. d,l-Sotalol Implantable Cardioverter-Defibrillator Study Group. *N Engl J Med.* 1999;340:1855–62.

**S7.1.3-3.** Kettner K, Mewis C, Dornberger V, et al. Efficacy of metoprolol and sotalol in the prevention of recurrences of sustained ventricular tachyarrhythmias in patients with an implantable cardioverter defibrillator. *Pacing Clin Electrophysiol.* 2002;25:1571–6.

**S7.1.3-4.** Sapp JL, Wells GA, Parkash R, et al. Ventricular tachycardia ablation versus escalation of antiarrhythmic drugs. *N Engl J Med.* 2016;375:111–21.

**S7.1.3-5.** Mallidi J, Nadkarni GN, Berger RD, et al. Meta-analysis of catheter ablation as an adjunct to medical therapy for treatment of ventricular tachycardia in patients with structural heart disease. *Heart Rhythm.* 2011;8:503–10.

**S7.1.3-6.** Marchlinski FE, Haffajee CI, Beshai JF, et al. Long-term success of irrigated radiofrequency catheter ablation of sustained ventricular tachycardia: post-approval THERMOCOOL VT trial. *J Am Coll Cardiol.* 2016;67:674–83.

**S7.1.3-7.** Stevenson WG, Wilber DJ, Natale A, et al. Irrigated radiofrequency catheter ablation guided by electroanatomic mapping for recurrent ventricular tachycardia after myocardial infarction: the multicenter thermocoil ventricular tachycardia ablation trial. *Circulation.* 2008;118:2773–82.

**S7.1.3-8.** Tanner H, Hindricks G, Volkmer M, et al. Catheter ablation of recurrent scar-related ventricular tachycardia using electroanatomical mapping and irrigated ablation technology: results of the prospective multicenter Euro-VT-study. *J Cardiovasc Electrophysiol.* 2010;21:47–53.

**S7.1.3-9.** Tung R, Vaseghi M, Frankel DS, et al. Freedom from recurrent ventricular tachycardia after catheter ablation is associated with improved survival in patients with structural heart disease: an International VT Ablation Center Collaborative Group study. *Heart Rhythm.* 2015;12:1997–2007.

**S7.1.3-10.** Al-Khatib SM, Daubert JP, Anstrom KJ, et al. Catheter ablation for ventricular tachycardia in

patients with an implantable cardioverter defibrillator (CALYPSO) pilot trial. *J Cardiovasc Electrophysiol.* 2015;26:151-7.

**S7.1.3-11.** Kuck KH, Schaumann A, Eckardt L, et al. Catheter ablation of stable ventricular tachycardia before defibrillator implantation in patients with coronary heart disease (VTACH): a multicentre randomised controlled trial. *Lancet.* 2010;375:31-40.

**S7.1.3-12.** Sesselberg HW, Moss AJ, McNitt S, et al. Ventricular arrhythmia storms in postinfarction patients with implantable defibrillators for primary prevention indications: a MADIT-II substudy. *Heart Rhythm.* 2007;4:1395-402.

**S7.1.3-13.** Echt DS, Liebson PR, Mitchell LB, et al. Mortality and morbidity in patients receiving encainide, flecainide, or placebo. The Cardiac Arrhythmia Suppression Trial. *N Engl J Med.* 1991;324:781-8.

**S7.1.3-14.** Sears SF Jr., Todaro JF, Lewis TS, et al. Examining the psychosocial impact of implantable cardioverter defibrillators: a literature review. *Clin Cardiol.* 1999;22:481-9.

**S7.1.3-15.** Steinberg JS, Gaur A, Sciacca R, et al. New-onset sustained ventricular tachycardia after cardiac surgery. *Circulation.* 1999;99:903-8.

**S7.1.3-16.** Brugada J, Aguinaga L, Mont L, et al. Coronary artery revascularization in patients with sustained ventricular arrhythmias in the chronic phase of a myocardial infarction: effects on the electrophysiologic substrate and outcome. *J Am Coll Cardiol.* 2001; 37:529-33.

**S7.1.3-17.** Claro JC, Candia R, Rada G, et al. Amiodarone versus other pharmacological interventions for prevention of sudden cardiac death. *Cochrane Database Syst Rev.* 2015;12:CD008093.

**S7.1.3-18.** Bunch TJ, Mahapatra S, Murdock D, et al. Ranolazine reduces ventricular tachycardia burden and ICD shocks in patients with drug-refractory ICD shocks. *Pacing Clin Electrophysiol.* 2011;34:1600-6.

**S7.1.3-19.** Goldschlager N, Epstein AE, Naccarelli GV, et al. A practical guide for clinicians who treat patients with amiodarone: 2007. *Heart Rhythm.* 2007; 4:1250-9.

**S7.1.3-20.** Singh SN, Fletcher RD, Fisher SG, et al. Amiodarone in patients with congestive heart failure and asymptomatic ventricular arrhythmia. Survival Trial of Antiarrhythmic Therapy in Congestive Heart Failure. *N Engl J Med.* 1995;333:77-82.

**S7.1.3-21.** The Cardiac Arrhythmia Suppression Trial II Investigators. Effect of the antiarrhythmic agent moricizine on survival after myocardial infarction. The Cardiac Arrhythmia Suppression Trial II Investigators. *N Engl J Med.* 1992;327:227-33.

**S7.1.3-22.** Kuck KH, Cappato R, Siebels J, et al. Randomized comparison of antiarrhythmic drug therapy with implantable defibrillators in patients resuscitated from cardiac arrest: the Cardiac Arrest Study Hamburg (CASH). *Circulation.* 2000;102:748-54.

**S7.1.3-23.** Sears SE Jr., Conti JB. Understanding implantable cardioverter defibrillator shocks and storms: medical and psychosocial considerations for research and clinical care. *Clin Cardiol.* 2003;26: 107-11.

**S7.1.3-24.** Sweeney MO, Sherfesee L, Degroot PJ, et al. Differences in effects of electrical therapy type for ventricular arrhythmias on mortality in implantable

cardioverter-defibrillator patients. *Heart Rhythm.* 2010;7:353-60.

**S7.1.3-25.** Berntsen RF, Gunnes P, Lie M, et al. Surgical revascularization in the treatment of ventricular tachycardia and fibrillation exposed by exercise-induced ischaemia. *Eur Heart J.* 1993;14:1297-303.

**S7.1.3-26.** Wilkoff BL, Fauchier L, Stiles MK, et al. 2015 HRS/EHRA/APHRS/SOLAECE expert consensus statement on optimal implantable cardioverter-defibrillator programming and testing. *J Arrhythm.* 2016;32:1-28.

## 7.2. Nonischemic Cardiomyopathy

**S7.2-1.** Greulich S, Deluigi CC, Gloekler S, et al. CMR imaging predicts death and other adverse events in suspected cardiac sarcoidosis. *J Am Coll Cardiol Img.* 2013;6:501-11.

**S7.2-2.** Kuruvilla S, Adenaw N, Katwal AB, et al. Late gadolinium enhancement on cardiac magnetic resonance predicts adverse cardiovascular outcomes in nonischemic cardiomyopathy: a systematic review and meta-analysis. *Circ Cardiovasc Imaging.* 2014;7:250-8.

**S7.2-3.** Piers SR, Tao Q, van Huls van Taxis CF, et al. Contrast-enhanced MRI-derived scar patterns and associated ventricular tachycardias in nonischemic cardiomyopathy: implications for the ablation strategy. *Circ Arrhythm Electrophysiol.* 2013;6:875-83.

**S7.2-4.** Ackerman MJ, Priori SG, Willems S, et al. HRS/EHRA expert consensus statement on the state of genetic testing for the channelopathies and cardiomyopathies: this document was developed as a partnership between the Heart Rhythm Society (HRS) and the European Heart Rhythm Association (EHRA). *Europace.* 2011;13:1077-109.

**S7.2-5.** Hershberger RE, Morales A, Siegfried JD. Clinical and genetic issues in dilated cardiomyopathy: a review for genetics professionals. *Genet Med.* 2010;12: 655-67.

**S7.2-6.** Coleman GC, Shaw PW, Balfour PC Jr., et al. Prognostic value of myocardial scarring on CMR in patients with cardiac sarcoidosis: a systematic review and meta-analysis. *J Am Coll Cardiol Img.* 2016. 2017;10:411-20.

**S7.2-7.** Anselme F, Moubarak G, Savoure A, et al. Implantable cardioverter-defibrillators in lamin A/C mutation carriers with cardiac conduction disorders. *Heart Rhythm.* 2013;10:1492-8.

**S7.2-8.** Kumar S, Baldinger SH, Gandjbakhch E, et al. Long-term arrhythmic and nonarrhythmic outcomes of lamin A/C mutation carriers. *J Am Coll Cardiol.* 2016; 68:2299-307.

**S7.2-9.** Meune C, Van Berlo JH, Anselme F, et al. Primary prevention of sudden death in patients with lamin A/C gene mutations. *N Engl J Med.* 2006;354: 209-10.

**S7.2-10.** Pasotti M, Klersy C, Pilotto A, et al. Long-term outcome and risk stratification in dilated cardiomyopathies. *J Am Coll Cardiol.* 2008;52: 1250-60.

**S7.2-11.** Van Berlo JH, de Voogt WG, van der Kooi AJ, et al. Meta-analysis of clinical characteristics of 299 carriers of LMNA gene mutations: do lamin A/C mutations portend a high risk of sudden death? *J Mol Med.* 2005;83:79-83.

**S7.2-12.** van Rijsingen IA, Arbustini E, Elliott PM, et al. Risk factors for malignant ventricular arrhythmias in

lamin A/C mutation carriers: a European cohort study. *J Am Coll Cardiol.* 2012;59:493-500.

## 7.2.1. Secondary Prevention of SCD in Patients With NICM

**S7.2.1-1.** The Antiarrhythmics versus Implantable Defibrillators (AVID) Investigators. A comparison of antiarrhythmic-drug therapy with implantable defibrillators in patients resuscitated from near-fatal ventricular arrhythmias. *N Engl J Med.* 1997;337: 1576-83.

**S7.2.1-2.** Connolly SJ, Gent M, Roberts RS, et al. Canadian implantable defibrillator study (CIDS): a randomized trial of the implantable cardioverter defibrillator against amiodarone. *Circulation.* 2000; 101:1297-302.

**S7.2.1-3.** Desai AS, Fang JC, Maisel WH, et al. Implantable defibrillators for the prevention of mortality in patients with nonischemic cardiomyopathy: a meta-analysis of randomized controlled trials. *JAMA.* 2004;292:2874-9.

**S7.2.1-4.** Kuck KH, Cappato R, Siebels J, et al. Randomized comparison of antiarrhythmic drug therapy with implantable defibrillators in patients resuscitated from cardiac arrest: the Cardiac Arrest Study Hamburg (CASH). *Circulation.* 2000;102:748-54.

**S7.2.1-5.** Raitt MH, Renfroe EG, Epstein AE, et al. "Stable" ventricular tachycardia is not a benign rhythm: insights from the Antiarrhythmics versus Implantable Defibrillators (AVID) registry. *Circulation.* 2001;103:244-52.

**S7.2.1-6.** Brilakis ES, Shen WK, Hammill SC, et al. Role of programmed ventricular stimulation and implantable cardioverter defibrillators in patients with idiopathic dilated cardiomyopathy and syncope. *Pacing Clin Electrophysiol.* 2001;24:1623-30.

**S7.2.1-7.** Fonarow GC, Feliciano Z, Boyle NG, et al. Improved survival in patients with nonischemic advanced heart failure and syncope treated with an implantable cardioverter-defibrillator. *Am J Cardiol.* 2000;85:981-5.

**S7.2.1-8.** Knight BP, Goyal R, Pelosi F, et al. Outcome of patients with nonischemic dilated cardiomyopathy and unexplained syncope treated with an implantable defibrillator. *J Am Coll Cardiol.* 1999;33:1964-70.

**S7.2.1-9.** Middlekauff HR, Stevenson WG, Stevenson LW, et al. Syncope in advanced heart failure: high risk of sudden death regardless of origin of syncope. *J Am Coll Cardiol.* 1993;21:110-6.

**S7.2.1-10.** Olshansky B, Poole JE, Johnson G, et al. Syncope predicts the outcome of cardiomyopathy patients: analysis of the SCD-HeFT study. *J Am Coll Cardiol.* 2008;51:1277-82.

**S7.2.1-11.** Ruwald MH, Okumura K, Kimura T, et al. Syncope in high-risk cardiomyopathy patients with implantable defibrillators: frequency, risk factors, mechanisms, and association with mortality: results from the multicenter automatic defibrillator implantation trial-reduce inappropriate therapy (MADIT-RIT) study. *Circulation.* 2014;129:545-52.

**S7.2.1-12.** Piccini JP, Berger JS, O'Connor CM. Amiodarone for the prevention of sudden cardiac death: a meta-analysis of randomized controlled trials. *Eur Heart J.* 2009;30:1245-53.

**S7.2.1-13.** Claro JC, Candia R, Rada G, et al. Amiodarone versus other pharmacological interventions for

prevention of sudden cardiac death. *Cochrane Database Syst Rev.* 2015;12:CD008093.

**S7.2.1-14.** Daubert JP, Winters SL, Subacius H, et al. Ventricular arrhythmia inducibility predicts subsequent ICD activation in nonischemic cardiomyopathy patients: a DEFINITE substudy. *Pacing Clin Electrophysiol.* 2009;32:755–61.

**S7.2.1-15.** Gatzoulis KA, Vouliotis AI, Tsachiris D, et al. Primary prevention of sudden cardiac death in a non-ischemic dilated cardiomyopathy population: reappraisal of the role of programmed ventricular stimulation. *Circ Arrhythm Electrophysiol.* 2013;6:504–12.

**S7.2.1-16.** Hilfiker G, Schoenenberger AW, Erne P, et al. Utility of electrophysiological studies to predict arrhythmic events. *World J Cardiol.* 2015;7:344–50.

**S7.2.1-17.** Thomas KL, Al-Khatib SM, Lohknygina Y, et al. Amiodarone use after acute myocardial infarction complicated by heart failure and/or left ventricular dysfunction may be associated with excess mortality. *Am Heart J.* 2008;155:87–93.

#### 7.2.2. Primary Prevention of SCD in Patients With NICM

**S7.2.2-1.** Bardy GH, Lee KL, Mark DB, et al. Amiodarone or an implantable cardioverter-defibrillator for congestive heart failure. *N Engl J Med.* 2005;352:225–37.

**S7.2.2-2.** Bänsch D, Antz M, Boczar S, et al. Primary prevention of sudden cardiac death in idiopathic dilated cardiomyopathy: the Cardiomyopathy Trial (CAT). *Circulation.* 2002;105:1453–8.

**S7.2.2-3.** Bristow MR, Saxon LA, Boehmer J, et al. Cardiac-resynchronization therapy with or without an implantable defibrillator in advanced chronic heart failure. *N Engl J Med.* 2004;350:2140–50.

**S7.2.2-4.** Desai AS, Fang JC, Maisel WH, et al. Implantable defibrillators for the prevention of mortality in patients with nonischemic cardiomyopathy: a meta-analysis of randomized controlled trials. *JAMA.* 2004;292:2874–9.

**S7.2.2-5.** Kadish A, Dyer A, Daubert JP, et al. Prophylactic defibrillator implantation in patients with non-ischemic dilated cardiomyopathy. *N Engl J Med.* 2004;350:2151–8.

**S7.2.2-6.** Strickberger SA, Hummel JD, Bartlett TG, et al. Amiodarone versus implantable cardioverter-defibrillator: randomized trial in patients with non-ischemic dilated cardiomyopathy and asymptomatic nonsustained ventricular tachycardia—AMIOVIRT. *J Am Coll Cardiol.* 2003;41:1707–12.

**S7.2.2-7.** Anselme F, Moubarak G, Savoure A, et al. Implantable cardioverter-defibrillators in lamin A/C mutation carriers with cardiac conduction disorders. *Heart Rhythm.* 2013;10:1492–8.

**S7.2.2-8.** Pasotti M, Klersy C, Pilotto A, et al. Long-term outcome and risk stratification in dilated cardiomopathies. *J Am Coll Cardiol.* 2008;52:1250–60.

**S7.2.2-9.** Van Berlo JH, de Voogt WG, van der Kooi AJ, et al. Meta-analysis of clinical characteristics of 299 carriers of LMNA gene mutations: do lamin A/C mutations portend a high risk of sudden death? *J Mol Med.* 2005;83:79–83.

**S7.2.2-10.** van Rijssingen IA, Arbustini E, Elliott PM, et al. Risk factors for malignant ventricular

arrhythmias in lamin A/C mutation carriers: a European cohort study. *J Am Coll Cardiol.* 2012;59:493–500.

**S7.2.2-11.** Køber L, Thune JJ, Nielsen JC, et al. Defibrillator implantation in patients with nonischemic systolic heart failure. *N Engl J Med.* 2016;375:1221–30.

**S7.2.2-12.** Al-Khatib SM, Fonarow GC, Joglar JA, et al. Primary prevention implantable cardioverter defibrillators in patients with nonischemic cardiomyopathy: a meta-analysis. *JAMA Cardiol.* 2017;2:685–8.

**S7.2.2-13.** Golwala H, Bajaj NS, Arora G, et al. Implantable cardioverter-defibrillator for nonischemic cardiomyopathy: an updated meta-analysis. *Circulation.* 2017;135:201–3.

#### 7.2.3. Treatment of Recurrent VA in Patients With NICM

**S7.2.3-1.** Connolly SJ, Dorian P, Roberts RS, et al. Comparison of beta-blockers, amiodarone plus beta-blockers, or sotalol for prevention of shocks from implantable cardioverter defibrillators: the OPTIC Study: a randomized trial. *JAMA.* 2006;295:165–71.

**S7.2.3-2.** Dinov B, Fiedler L, Schonbauer R, et al. Outcomes in catheter ablation of ventricular tachycardia in dilated nonischemic cardiomyopathy compared with ischemic cardiomyopathy: results from the Prospective Heart Centre of Leipzig VT (HELP-VT) Study. *Circulation.* 2014;129:728–36.

**S7.2.3-3.** Tung R, Vaseghi M, Frankel DS, et al. Freedom from recurrent ventricular tachycardia after catheter ablation is associated with improved survival in patients with structural heart disease: an International VT Ablation Center Collaborative Group study. *Heart Rhythm.* 2015;12:1997–2007.

**S7.2.3-4.** Piers SR, Tao Q, Tvan Huls van Taxis CF, et al. Contrast-enhanced MRI-derived scar patterns and associated ventricular tachycardias in nonischemic cardiomyopathy: implications for the ablation strategy. *Circ Arrhythm Electrophysiol.* 2013;6:875–83.

**S7.2.3-5.** Desjardins B, Yokokawa M, Good E, et al. Characteristics of intramural scar in patients with nonischemic cardiomyopathy and relation to intramural ventricular arrhythmias. *Circ Arrhythm Electrophysiol.* 2013;6:891–7.

**S7.2.3-6.** Naruse Y, Sekiguchi Y, Nogami A, et al. Systematic treatment approach to ventricular tachycardia in cardiac sarcoidosis. *Circ Arrhythm Electrophysiol.* 2014;7:407–13.

#### 7.3. Arrhythmogenic Right Ventricular Cardiomyopathy

**S7.3-1.** Groeneweg JA, Bhonsale A, James CA, et al. Clinical presentation, long-term follow-up, and outcomes of 1001 arrhythmogenic right ventricular dysplasia cardiomyopathy patients and family members. *Circ Cardiovasc Genet.* 2015;8:437–46.

**S7.3-2.** Marcus Fl, Edson S, Towbin JA. Genetics of arrhythmogenic right ventricular cardiomyopathy: a practical guide for physicians. *J Am Coll Cardiol.* 2013;61:1945–8.

**S7.3-3.** Quarta G, Muir A, Pantazis A, et al. Familial evaluation in arrhythmogenic right ventricular cardiomyopathy: impact of genetics and revised task force criteria. *Circulation.* 2011;123:2701–9.

**S7.3-4.** te Riele AS, James CA, Groeneweg JA, et al. Approach to family screening in arrhythmogenic right

ventricular dysplasia/cardiomyopathy. *Eur Heart J.* 2016;37:755–63.

**S7.3-5.** Liu T, Pursnani A, Sharma UC, et al. Effect of the 2010 task force criteria on reclassification of cardiovascular magnetic resonance criteria for arrhythmogenic right ventricular cardiomyopathy. *J Cardiovasc Magn Reson.* 2014;16:47.

**S7.3-6.** Marcus Fl, McKenna WJ, Sherrill D, et al. Diagnosis of arrhythmogenic right ventricular cardiomyopathy/dysplasia: proposed modification of the task force criteria. *Circulation.* 2010;121:1533–41.

**S7.3-7.** te Riele AS, Bhonsale A, James CA, et al. Incremental value of cardiac magnetic resonance imaging in arrhythmic risk stratification of arrhythmogenic right ventricular dysplasia/cardiomyopathy-associated desmosomal mutation carriers. *J Am Coll Cardiol.* 2013;62:1761–9.

**S7.3-8.** Verma E, Strohm O, Otmani A, et al. Impact of the revision of arrhythmogenic right ventricular cardiomyopathy/dysplasia task force criteria on its prevalence by CMR criteria. *J Am Coll Cardiol Img.* 2011;4:282–7.

**S7.3-9.** Bhonsale A, James CA, Tichnell C, et al. Incidence and predictors of implantable cardioverter-defibrillator therapy in patients with arrhythmogenic right ventricular dysplasia/cardiomyopathy undergoing implantable cardioverter-defibrillator implantation for primary prevention. *J Am Coll Cardiol.* 2011;58:1485–96.

**S7.3-10.** Corrado D, Calkins H, Link MS, et al. Prophylactic implantable defibrillator in patients with arrhythmogenic right ventricular cardiomyopathy/dysplasia and no prior ventricular fibrillation or sustained ventricular tachycardia. *Circulation.* 2010;122:1144–52.

**S7.3-11.** Corrado D, Wichter T, Link MS, et al. Treatment of arrhythmogenic right ventricular cardiomyopathy/dysplasia: an international task force consensus statement. *Circulation.* 2015;132:441–53.

**S7.3-12.** Link MS, Laidlaw D, Polonsky B, et al. Ventricular arrhythmias in the North American multidisciplinary study of ARVC: predictors, characteristics, and treatment. *J Am Coll Cardiol.* 2014;64:119–25.

**S7.3-13.** Piccini JP, Dalal D, Roguin A, et al. Predictors of appropriate implantable defibrillator therapies in patients with arrhythmogenic right ventricular dysplasia. *Heart Rhythm.* 2005;2:1188–94.

**S7.3-14.** Marcus Fl, Zareba W, Calkins H, et al. Arrhythmogenic right ventricular cardiomyopathy/dysplasia clinical presentation and diagnostic evaluation: results from the North American Multidisciplinary Study. *Heart Rhythm.* 2009;6:984–92.

**S7.3-15.** Marcus GM, Glidden DV, Polonsky B, et al. Efficacy of antiarrhythmic drugs in arrhythmogenic right ventricular cardiomyopathy: a report from the North American ARVC Registry. *J Am Coll Cardiol.* 2009;54:609–15.

**S7.3-16.** Corrado D, Bassi C, Thiene G, et al. Spectrum of clinicopathologic manifestations of arrhythmogenic right ventricular cardiomyopathy/dysplasia: a multicenter study. *J Am Coll Cardiol.* 1997;30:1512–20.

**S7.3-17.** James CA, Bhonsale A, Tichnell C, et al. Exercise increases age-related penetrance and

arrhythmic risk in arrhythmogenic right ventricular dysplasia/cardiomyopathy-associated desmosomal mutation carriers. *J Am Coll Cardiol.* 2013;62:1290-7.

**S7.3-18.** Ruwald AC, Marcus F, Estes NA 3rd, et al. Association of competitive and recreational sport participation with cardiac events in patients with arrhythmogenic right ventricular cardiomyopathy: results from the North American multidisciplinary study of arrhythmogenic right ventricular cardiomyopathy. *Eur Heart J.* 2015;36:1735-43.

**S7.3-19.** Saberniak J, Hasselberg NE, Borgquist R, et al. Vigorous physical activity impairs myocardial function in patients with arrhythmogenic right ventricular cardiomyopathy and in mutation positive family members. *Eur J Heart Fail.* 2014;16:1337-44.

**S7.3-20.** Sawant AC, Bhonsale A, te Riele AS, et al. Exercise has a disproportionate role in the pathogenesis of arrhythmogenic right ventricular dysplasia/cardiomyopathy in patients without desmosomal mutations. *J Am Heart Assoc.* 2014;3:e001471.

**S7.3-21.** Sawant AC, te Riele AS, Tichnell C, et al. Safety of American Heart Association-recommended minimum exercise for desmosomal mutation carriers. *Heart Rhythm.* 2016;13:199-207.

**S7.3-22.** Bhonsale A, Groeneweg JA, James CA, et al. Impact of genotype on clinical course in arrhythmogenic right ventricular dysplasia/cardiomyopathy-associated mutation carriers. *Eur Heart J.* 2015;36:847-55.

**S7.3-23.** Bhonsale A, James CA, Tichnell C, et al. Risk stratification in arrhythmogenic right ventricular dysplasia/cardiomyopathy-associated desmosomal mutation carriers. *Circ Arrhythm Electrophysiol.* 2013;6:569-78.

**S7.3-24.** Hershberger RE, Lindenfeld J, Mestroni L, et al. Genetic evaluation of cardiomyopathy—a Heart Failure Society of America practice guideline. *J Card Fail.* 2009;15:83-97.

**S7.3-25.** Kapplinger JD, Landstrom AP, Salisbury BA, et al. Distinguishing arrhythmogenic right ventricular cardiomyopathy/dysplasia-associated mutations from background genetic noise. *J Am Coll Cardiol.* 2011;57:2317-27.

**S7.3-26.** Rigato I, Baucé B, Rampazzo A, et al. Compound and digenic heterozygosity predicts lifetime arrhythmic outcome and sudden cardiac death in desmosomal gene-related arrhythmogenic right ventricular cardiomyopathy. *Circ Cardiovasc Genet.* 2013;6:533-42.

**S7.3-27.** Philips B, te Riele AS, Sawant A, et al. Outcomes and ventricular tachycardia recurrence characteristics after epicardial ablation of ventricular tachycardia in arrhythmogenic right ventricular dysplasia/cardiomyopathy. *Heart Rhythm.* 2015;12:716-25.

**S7.3-28.** Philips B, Madhavan S, James C, et al. Outcomes of catheter ablation of ventricular tachycardia in arrhythmogenic right ventricular dysplasia/cardiomyopathy. *Circ Arrhythm Electrophysiol.* 2012;5:499-505.

**S7.3-29.** Dalal D, Jain R, Tandri H, et al. Long-term efficacy of catheter ablation of ventricular tachycardia in patients with arrhythmogenic right ventricular dysplasia/cardiomyopathy. *J Am Coll Cardiol.* 2007;50:432-40.

**S7.3-30.** Garcia FC, Bazan V, Zado ES, et al. Epicardial substrate and outcome with epicardial ablation of ventricular tachycardia in arrhythmogenic right ventricular cardiomyopathy/dysplasia. *Circulation.* 2009;120:366-75.

**S7.3-31.** Bai R, Di BL, Shivkumar K, et al. Ablation of ventricular arrhythmias in arrhythmogenic right ventricular dysplasia/cardiomyopathy: arrhythmia-free survival after endo-epicardial substrate based mapping and ablation. *Circ Arrhythm Electrophysiol.* 2011;4:478-85.

**S7.3-32.** Berrezo A, Fernandez-Armenta J, Mont L, et al. Combined endocardial and epicardial catheter ablation in arrhythmogenic right ventricular dysplasia incorporating scar dechanneling technique. *Circ Arrhythm Electrophysiol.* 2012;5:111-21.

**S7.3-33.** Santangeli P, Zado ES, Supple GE, et al. Long-term outcome with catheter ablation of ventricular tachycardia in patients with arrhythmogenic right ventricular cardiomyopathy. *Circ Arrhythm Electrophysiol.* 2015;8:1413-21.

**S7.3-34.** Choudhary N, Tompkins C, Polonsky B, et al. Clinical Presentation and Outcomes by Sex in Arrhythmogenic Right Ventricular Cardiomyopathy: findings from the North American ARVC Registry. *J Cardiovasc Electrophysiol.* 2016;27:555-62.

**S7.3-35.** Kamath GS, Zareba W, Delaney J, et al. Value of the signal-averaged electrocardiogram in arrhythmogenic right ventricular cardiomyopathy/dysplasia. *Heart Rhythm.* 2011;8:256-62.

**S7.3-36.** Saguner AM, Medeiros-Domingo A, Schwyzer MA, et al. Usefulness of inducible ventricular tachycardia to predict long-term adverse outcomes in arrhythmogenic right ventricular cardiomyopathy. *Am J Cardiol.* 2013;111:250-7.

**S7.3-37.** Sen-Chowdhry S, Syrris P, Prasad SK, et al. Left-dominant arrhythmogenic cardiomyopathy: an under-recognized clinical entity. *J Am Coll Cardiol.* 2008;52:2175-87.

**S7.3-38.** Te Riele AS, James CA, Philips B, et al. Mutation-positive arrhythmogenic right ventricular dysplasia/cardiomyopathy: the triangle of dysplasia displaced. *J Cardiovasc Electrophysiol.* 2013;24:1311-20.

**S7.3-39.** Rastegar N, Zimmerman SL, Te Riele AS, et al. Spectrum of biventricular involvement on cmr among carriers of arvd/c-associated mutations. *J Am Coll Cardiol Img.* 2015;8:863-4.

**S7.3-40.** Sen-Chowdhry S, Syrris P, Ward D, et al. Clinical and genetic characterization of families with arrhythmogenic right ventricular dysplasia/cardiomyopathy provides novel insights into patterns of disease expression. *Circulation.* 2007;115:1710-20.

**S7.3-41.** Dalal D, Nasir K, Bomma C, et al. Arrhythmogenic right ventricular dysplasia: a United States experience. *Circulation.* 2005;112:3823-32.

**S7.3-42.** Brun F, Groeneweg JA, Gear K, et al. Risk stratification in arrhythmic right ventricular cardiomyopathy without implantable cardioverter-defibrillators. *JACC Clin Electrophysiol.* 2016;2:558-64.

**S7.3-43.** Corrado D, Leoni L, Link MS, et al. Implantable cardioverter-defibrillator therapy for prevention of sudden death in patients with arrhythmogenic right ventricular cardiomyopathy/dysplasia. *Circulation.* 2003;108:3084-91.

**S7.3-44.** van Rijssingen IA, Arbustini E, Elliott PM, et al. Risk factors for malignant ventricular arrhythmias in lamin A/C mutation carriers: a European cohort study. *J Am Coll Cardiol.* 2012;59:493-500.

**S7.3-45.** Cox MG, van der Zwaag PA, van der Werf C, et al. Arrhythmogenic right ventricular dysplasia/cardiomyopathy: pathogenic desmosome mutations in index-patients predict outcome of family screening: Dutch arrhythmogenic right ventricular dysplasia/cardiomyopathy genotype-phenotype follow-up study. *Circulation.* 2011;123:2690-700.

**S7.3-46.** Saeed M, Homoud MK, Wang PJ, et al. Role of invasive electrophysiologic testing in risk stratification for sudden cardiac death. *J Invasive Cardiol.* 2001;13:758-62.

**S7.3-47.** Zhu DW, Sun H, Hill R, et al. The value of electrophysiology study and prophylactic implantation of cardioverter defibrillator in patients with hypertrophic cardiomyopathy. *Pacing Clin Electrophysiol.* 1998;21:299-302.

**S7.3-48.** Wichter T, Borggreve M, Havercamp W, et al. Efficacy of antiarrhythmic drugs in patients with arrhythmogenic right ventricular disease. Results in patients with inducible and noninducible ventricular tachycardia. *Circulation.* 1992;86:29-37.

#### 7.4. Hypertrophic Cardiomyopathy

**S7.4-1.** Elliott PM, Sharma S, Varnava A, et al. Survival after cardiac arrest or sustained ventricular tachycardia in patients with hypertrophic cardiomyopathy. *J Am Coll Cardiol.* 1999;33:1596-601.

**S7.4-2.** Elliott PM, Poloniecki J, Dickie S, et al. Sudden death in hypertrophic cardiomyopathy: identification of high risk patients. *J Am Coll Cardiol.* 2000;36:2212-8.

**S7.4-3.** Elliott PM, Gimeno Blanes JR, Mahon NG, et al. Relation between severity of left-ventricular hypertrophy and prognosis in patients with hypertrophic cardiomyopathy. *Lancet.* 2001;357:420-4.

**S7.4-4.** Elliott PM, Gimeno JR, Tome MT, et al. Left ventricular outflow tract obstruction and sudden death risk in patients with hypertrophic cardiomyopathy. *Eur Heart J.* 2006;27:1933-41.

**S7.4-5.** Maki S, Ikeda H, Muro A, et al. Predictors of sudden cardiac death in hypertrophic cardiomyopathy. *Am J Cardiol.* 1998;82:774-8.

**S7.4-6.** Maron BJ, Spirito P, Shen WK, et al. Implantable cardioverter-defibrillators and prevention of sudden cardiac death in hypertrophic cardiomyopathy. *JAMA.* 2007;298:405-12.

**S7.4-7.** O'Mahony C, Jichi F, Pavlou M, et al. A novel clinical risk prediction model for sudden cardiac death in hypertrophic cardiomyopathy (HCM risk-SCD). *Eur Heart J.* 2014;35:2010-20.

**S7.4-8.** Spirito P, Autore C, Rapezzi C, et al. Syncope and risk of sudden death in hypertrophic cardiomyopathy. *Circulation.* 2009;119:1703-10.

**S7.4-9.** O'Mahony C, Lambiase PD, Quarta G, et al. The long-term survival and the risks and benefits of implantable cardioverter defibrillators in patients with hypertrophic cardiomyopathy. *Heart.* 2012;98:116-25.

**S7.4-10.** Syska P, Przybylski A, Chojnowska L, et al. Implantable cardioverter-defibrillator in patients with hypertrophic cardiomyopathy: efficacy and complications of the therapy in long-term follow-up. *J Cardiovasc Electrophysiol.* 2010;21:883-9.

- S7.4-11.** Adabag AS, Kuskowski MA, Maron BJ. Determinants for clinical diagnosis of hypertrophic cardiomyopathy. *Am J Cardiol.* 2006;98:1507–11.
- S7.4-12.** Afonso LC, Bernal J, Bax JJ, et al. Echocardiography in hypertrophic cardiomyopathy: the role of conventional and emerging technologies. *J Am Coll Cardiol Img.* 2008;1:787–800.
- S7.4-13.** Girolami F, Ho CY, Semsarian C, et al. Clinical features and outcome of hypertrophic cardiomyopathy associated with triple sarcomere protein gene mutations. *J Am Coll Cardiol.* 2010;55:1444–53.
- S7.4-14.** Ingles J, Sarina T, Yeates L, et al. Clinical predictors of genetic testing outcomes in hypertrophic cardiomyopathy. *Genet Med.* 2013;15:972–7.
- S7.4-15.** Jensen MK, Havndrup O, Christiansen M, et al. Penetrance of hypertrophic cardiomyopathy in children and adolescents: a 12-year follow-up study of clinical screening and predictive genetic testing. *Circulation.* 2013;127:48–54.
- S7.4-16.** Klues HG, Schiffrers A, Maron BJ. Phenotypic spectrum and patterns of left ventricular hypertrophy in hypertrophic cardiomyopathy: morphologic observations and significance as assessed by two-dimensional echocardiography in 600 patients. *J Am Coll Cardiol.* 1995;26:1699–708.
- S7.4-17.** Wigle ED, Rakowski H, Kimball BP, et al. Hypertrophic cardiomyopathy. Clinical spectrum and treatment. *Circulation.* 1995;92:1680–92.
- S7.4-18.** Christiaans I, Birnie E, van Langen IM, et al. The yield of risk stratification for sudden cardiac death in hypertrophic cardiomyopathy myosin-binding protein C gene mutation carriers: focus on predictive screening. *Eur Heart J.* 2010;31:842–8.
- S7.4-19.** Olivotto I, Girolami F, Ackerman MJ, et al. Myofilament protein gene mutation screening and outcome of patients with hypertrophic cardiomyopathy. *Mayo Clin Proc.* 2008;83:630–8.
- S7.4-20.** Christiaans I, van Langen IM, Birnie E, et al. Genetic counseling and cardiac care in predictively tested hypertrophic cardiomyopathy mutation carriers: the patients' perspective. *Am J Med Genet.* 2009;149a:1444–51.
- S7.4-21.** Hamang A, Eide GE, Rokne B, et al. Predictors of heart-focused anxiety in patients undergoing genetic investigation and counseling of long QT syndrome or hypertrophic cardiomyopathy: a one year follow-up. *J Genet Counsel.* 2012;21:72–84.
- S7.4-22.** Bos JM, Will ML, Gersh BJ, et al. Characterization of a phenotype-based genetic test prediction score for unrelated patients with hypertrophic cardiomyopathy. *Mayo Clin Proc.* 2014;89:727–37.
- S7.4-23.** Soraja P, Nishimura RA, Ommen SR, et al. Use of echocardiography in patients with hypertrophic cardiomyopathy: clinical implications of massive hypertrophy. *J Am Soc Echocardiogr.* 2006;19:788–95.
- S7.4-24.** Spirito P, Bellone P, Harris KM, et al. Magnitude of left ventricular hypertrophy and risk of sudden death in hypertrophic cardiomyopathy. *N Engl J Med.* 2000;342:1778–85.
- S7.4-25.** Bos JM, Maron BJ, Ackerman MJ, et al. Role of family history of sudden death in risk stratification and prevention of sudden death with implantable defibrillators in hypertrophic cardiomyopathy. *Am J Cardiol.* 2010;106:1481–6.
- S7.4-26.** Maron BJ, Shen WK, Link MS, et al. Efficacy of implantable cardioverter-defibrillators for the prevention of sudden death in patients with hypertrophic cardiomyopathy. *N Engl J Med.* 2000;342:365–73.
- S7.4-27.** Monserrat L, Elliott PM, Gimeno JR, et al. Non-sustained ventricular tachycardia in hypertrophic cardiomyopathy: an independent marker of sudden death risk in young patients. *J Am Coll Cardiol.* 2003;42:873–9.
- S7.4-28.** Olivotto I, Maron BJ, Montereleggi A, et al. Prognostic value of systemic blood pressure response during exercise in a community-based patient population with hypertrophic cardiomyopathy. *J Am Coll Cardiol.* 1999;33:2044–51.
- S7.4-29.** Sadoul N, Prasad K, Elliott PM, et al. Prospective prognostic assessment of blood pressure response during exercise in patients with hypertrophic cardiomyopathy. *Circulation.* 1997;96:2987–91.
- S7.4-30.** McKenna WJ, Oakley CM, Krikler DM, et al. Improved survival with amiodarone in patients with hypertrophic cardiomyopathy and ventricular tachycardia. *Br Heart J.* 1985;53:412–6.
- S7.4-31.** Melacini P, Maron BJ, Bobbo F, et al. Evidence that pharmacological strategies lack efficacy for the prevention of sudden death in hypertrophic cardiomyopathy. *Heart.* 2007;93:708–10.
- S7.4-32.** Kuck KH, Kunze KP, Schluter M, et al. Programmed electrical stimulation in hypertrophic cardiomyopathy. Results in patients with and without cardiac arrest or syncope. *Eur Heart J.* 1988;9:177–85.
- S7.4-33.** Zhu DW, Sun H, Hill R, et al. The value of electrophysiology study and prophylactic implantation of cardioverter defibrillator in patients with hypertrophic cardiomyopathy. *Pacing Clin Electrophysiol.* 1998;21:299–302.
- S7.4-34.** Ackerman MJ, VanDriest SL, Ommen SR, et al. Prevalence and age-dependence of malignant mutations in the beta-myosin heavy chain and troponin T genes in hypertrophic cardiomyopathy: a comprehensive outpatient perspective. *J Am Coll Cardiol.* 2002;39:2042–8.
- S7.4-35.** Lopes LR, Rahman MS, Elliott PM. A systematic review and meta-analysis of genotype-phenotype associations in patients with hypertrophic cardiomyopathy caused by sarcomeric protein mutations. *Heart.* 2013;99:1800–11.
- S7.4-36.** Gersh BJ, Maron BJ, Bonow RO, et al. 2011 ACCF/AHA guideline for the diagnosis and treatment of hypertrophic cardiomyopathy: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Developed in collaboration with the American Association for Thoracic Surgery, American Society of Echocardiography, American Society of Nuclear Cardiology, Heart Failure Society of America, Heart Rhythm Society, Society for Cardiovascular Angiography and Interventions, and Society of Thoracic Surgeons. *J Am Coll Cardiol.* 2011;58:e212–60.
- S7.4-37.** Green JJ, Berger JS, Kramer CM, et al. Prognostic value of late gadolinium enhancement in clinical outcomes for hypertrophic cardiomyopathy. *J Am Coll Cardiol Img.* 2012;5:370–7.
- S7.4-38.** O'Hanlon R, Grasso A, Roughton M, et al. Prognostic significance of myocardial fibrosis in hypertrophic cardiomyopathy. *J Am Coll Cardiol.* 2010;56:867–74.
- S7.4-39.** Rubinshtain R, Glockner JF, Ommen SR, et al. Characteristics and clinical significance of late gadolinium enhancement by contrast-enhanced magnetic resonance imaging in patients with hypertrophic cardiomyopathy. *Circ Heart Fail.* 2010;3:51–8.
- S7.4-40.** Rowin EJ, Maron BJ, Haas TS, et al. Hypertrophic cardiomyopathy with left ventricular apical aneurysm: implications for risk stratification and management. *J Am Coll Cardiol.* 2017;69:761–73.
- S7.4-41.** Maron BJ, Rowin EJ, Casey SA, et al. Risk stratification and outcome of patients with hypertrophic cardiomyopathy  $\geq 60$  years of age. *Circulation.* 2013;127:585–93.
- S7.4-42.** Francia P, Adduci C, Palano F, et al. Eligibility for the subcutaneous implantable cardioverter-defibrillator in patients with hypertrophic cardiomyopathy. *J Cardiovasc Electrophysiol.* 2015;26:893–9.
- S7.4-43.** Christiaans I, Birnie E, Bonsel GJ, et al. Manifest disease, risk factors for sudden cardiac death, and cardiac events in a large nationwide cohort of predictively tested hypertrophic cardiomyopathy mutation carriers: determining the best cardiological screening strategy. *Eur Heart J.* 2011;32:1161–70.
- S7.4-44.** Morita H, Rehm HL, Meneses A, et al. Shared genetic causes of cardiac hypertrophy in children and adults. *N Engl J Med.* 2008;358:1899–908.
- S7.4-45.** Niimura H, Bachinski LL, Sangwanaratne S, et al. Mutations in the gene for cardiac myosin-binding protein C and late-onset familial hypertrophic cardiomyopathy. *N Engl J Med.* 1998;338:1248–57.
- S7.4-46.** Maron BJ, Semsarian C. Emergence of gene mutation carriers and the expanding disease spectrum of hypertrophic cardiomyopathy. *Eur Heart J.* 2010;31:1551–3.
- S7.4-47.** Maron BJ, Yeates L, Semsarian C. Clinical challenges of genotype positive (+)-phenotype negative (-) family members in hypertrophic cardiomyopathy. *Am J Cardiol.* 2011;107:604–8.
- S7.4-48.** Ingles J, Doolan A, Chiu C, et al. Compound and double mutations in patients with hypertrophic cardiomyopathy: implications for genetic testing and counselling. *J Med Genet.* 2005;42:e59.
- S7.4-49.** Rosenzweig A, Watkins H, Hwang DS, et al. Preclinical diagnosis of familial hypertrophic cardiomyopathy by genetic analysis of blood lymphocytes. *N Engl J Med.* 1991;325:1753–60.
- S7.4-50.** O'Mahony C, Jichi F, Monserrat L, et al. Inverted U-shaped relation between the risk of sudden cardiac death and maximal left ventricular wall thickness in hypertrophic cardiomyopathy. *Circ Arrhythm Electrophysiol.* 2016;9:e003818.
- S7.4-51.** Saeed M, Homoud MK, Wang PJ, et al. Role of invasive electrophysiologic testing in risk stratification for sudden cardiac death. *J Invasive Cardiol.* 2001;13:758–62.
- S7.4-52.** Maron BJ, Haas TS, Goodman JS. Hypertrophic cardiomyopathy: one gene ... but many phenotypes. *Am J Cardiol.* 2014;113:1772–3.
- S7.4-53.** Ho CY, Cirino AL, Lakdawala NK, et al. Evolution of hypertrophic cardiomyopathy in sarcomere mutation carriers. *Heart.* 2016;102:1805–12.
- S7.4-54.** Moon JC, McKenna WJ, McCrohon JA, et al. Toward clinical risk assessment in hypertrophic cardiomyopathy with gadolinium cardiovascular magnetic resonance. *J Am Coll Cardiol.* 2003;41:1561–7.

**S7.4-55.** Maron MS, Finley JJ, Bos JM, et al. Prevalence, clinical significance, and natural history of left ventricular apical aneurysms in hypertrophic cardiomyopathy. *Circulation.* 2008;118:1541-9.

**S7.4-56.** Minami Y, Kajimoto K, Terajima Y, et al. Clinical implications of midventricular obstruction in patients with hypertrophic cardiomyopathy. *J Am Coll Cardiol.* 2011;57:2346-55.

## 7.5. Myocarditis

**S7.5-1.** Aoyama N, Izumi T, Hiramori K, et al. National survey of fulminant myocarditis in Japan: therapeutic guidelines and long-term prognosis of using percutaneous cardiopulmonary support for fulminant myocarditis (special report from a scientific committee). *Circ J.* 2002;66:133-44.

**S7.5-2.** Maleszewski JJ, Orellana VM, Hodge DO, et al. Long-term risk of recurrence, morbidity and mortality in giant cell myocarditis. *Am J Cardiol.* 2015;115:1733-8.

**S7.5-3.** Cooper LT Jr., Berry GJ, Shabetai R. Idiopathic giant-cell myocarditis—natural history and treatment. Multicenter Giant Cell Myocarditis Study Group Investigators. *N Engl J Med.* 1997;336:1860-6.

**S7.5-4.** Kandolin R, Lehtonen J, Salmenkivi K, et al. Diagnosis, treatment, and outcome of giant-cell myocarditis in the era of combined immunosuppression. *Circ Heart Fail.* 2013;6:15-22.

**S7.5-5.** Friman G, Wesslen L, Fohlman J, et al. The epidemiology of infectious myocarditis, lymphocytic myocarditis and dilated cardiomyopathy. *Eur Heart J.* 1995;16 suppl O:36-41.

**S7.5-6.** Hufnagel G, Pankweitz S, Richter A, et al. The European Study of Epidemiology and Treatment of Cardiac Inflammatory Diseases (ESET-CID). First epidemiological results. *Herz.* 2000;25:279-85.

**S7.5-7.** Winkel E, Parrillo J. Myocarditis. *Curr Treat Options. Cardiovasc Med.* 2002;4:455-66.

**S7.5-8.** Zawadowski GM, Klarich KW, Moder KG, et al. A contemporary case series of lupus myocarditis. *Lupus.* 2012;21:1378-84.

**S7.5-9.** Caforio AL, Pankweitz S, Arbustini E, et al. Current state of knowledge on aetiology, diagnosis, management, and therapy of myocarditis: a position statement of the European Society of Cardiology Working Group on Myocardial and Pericardial Diseases. *Eur Heart J.* 2013;34:2636-48d.

**S7.5-10.** Elamm C, Fairweather D, Cooper LT. Pathogenesis and diagnosis of myocarditis. *Heart.* 2012;98:835-40.

**S7.5-11.** Sagar S, Liu PP, Cooper LT Jr.. Myocarditis. *Lancet.* 2012;379:738-47.

**S7.5-12.** McCarthy RE 3rd, Boehmer JP, Hruban RH, et al. Long-term outcome of fulminant myocarditis as compared with acute (nonfulminant) myocarditis. *N Engl J Med.* 2000;342:690-5.

**S7.5-13.** Kawai C. From myocarditis to cardiomyopathy: mechanisms of inflammation and cell death: learning from the past for the future. *Circulation.* 1999;99:1091-100.

**S7.5-14.** Kindermann I, Kindermann M, Kandolf R, et al. Predictors of outcome in patients with suspected myocarditis. *Circulation.* 2008;118:639-48.

**S7.5-15.** Tubman TR, Craig B, Mulholland HC. Ventricular tachycardia associated with Coxsackie B4 virus infection. *Acta Paediatr Scand.* 1990;79:572-5.

**S7.5-16.** Tai YT, Lau CP, Fong PC, et al. Incessant automatic ventricular tachycardia complicating acute coxsackie B myocarditis. *Cardiology.* 1992;80:339-44.

**S7.5-17.** Dhar KL, Adlakha A, Phillip PJ. Recurrent seizures and syncope, ventricular arrhythmias with reversible prolonged Q-Tc interval in typhoid myocarditis. *J Indian Med Assoc.* 1987;85:336-7.

**S7.5-18.** McMurray JJ, Adamopoulos S, Anker SD, et al. ESC guidelines for the diagnosis and treatment of acute and chronic heart failure 2012: the Task Force for the Diagnosis and Treatment of Acute and Chronic Heart Failure 2012 of the European Society of Cardiology. Developed in collaboration with the Heart Failure Association (HFA) of the ESC. *Eur J Heart Fail.* 2012;14:803-69.

**S7.5-19.** Cooper LT Jr., Elamm C. Giant cell myocarditis. Diagnosis and treatment. *Herz.* 2012;37:632-6.

**S7.5-20.** Milei J, Grana D, Fernandez AG, et al. Cardiac involvement in acquired immunodeficiency syndrome—a review to push action. The Committee for the Study of Cardiac Involvement in AIDS. *Clin Cardiol.* 1998;21:465-72.

**S7.5-21.** Yunis NA, Stone VE. Cardiac manifestations of HIV/AIDS: a review of disease spectrum and clinical management. *J Acquir Immune Defic Syndr Hum Retrovir.* 1998;18:145-54.

**S7.5-22.** Huang CN, Yu HH, Chiu SN, et al. Acute myocarditis and ventricular fibrillation as initial presentation of pediatric systemic lupus erythematosus. *Rheumatol Int.* 2013;33:1093-6.

**S7.5-23.** Rassi A Jr., Rassi A, Little WC. Chagas' heart disease. *Clin Cardiol.* 2000;23:883-9.

**S7.5-24.** Muratore C, Rabinovich R, Iglesias R, et al. Implantable cardioverter defibrillators in patients with Chagas' disease: are they different from patients with coronary disease? *Pacing Clin Electrophysiol.* 1997;20:194-7.

**S7.5-25.** d'Avila A, Splinter R, Svenson RH, et al. New perspectives on catheter-based ablation of ventricular tachycardia complicating Chagas' disease: experimental evidence of the efficacy of near infrared lasers for catheter ablation of Chagas' VT. *J Interv Card Electrophysiol.* 2002;7:23-38.

**S7.5-26.** Kao AC, Krause SW, Handa R, et al. Wearable defibrillator use in heart failure (WIF): results of a prospective registry. *BMC Cardiovasc Disord.* 2012;12:123.

**S7.5-27.** Feldman AM, Klein H, Tchou P, et al. Use of a wearable defibrillator in terminating tachyarrhythmias in patients at high risk for sudden death: results of the WEARIT/BROAD. *Pacing Clin Electrophysiol.* 2004;27:4-9.

**S7.5-28.** Cooper LT, Okura Y. Idiopathic giant cell myocarditis. *Curr Treat Options Cardiovasc Med.* 2001;3:463-7.

## 7.6. Cardiac Sarcoidosis

**S7.6-1.** Kandolin R, Lehtonen J, Airaksinen J, et al. Cardiac sarcoidosis: epidemiology, characteristics, and outcome over 25 years in a nationwide study. *Circulation.* 2015;131:624-32.

**S7.6-2.** Yazaki Y, Isobe M, Hiroe M, et al. Prognostic determinants of long-term survival in Japanese patients with cardiac sarcoidosis treated with prednisone. *Am J Cardiol.* 2001;88:1006-10.

**S7.6-3.** Kron J, Sauer W, Schuller J, et al. Efficacy and safety of implantable cardiac defibrillators for treatment of ventricular arrhythmias in patients with cardiac sarcoidosis. *Europace.* 2013;15:347-54.

**S7.6-4.** Mohsen A, Jimenez A, Hood RE, et al. Cardiac sarcoidosis: electrophysiological outcomes on long-term follow-up and the role of the implantable cardioverter-defibrillator. *J Cardiovasc Electrophysiol.* 2014;25:171-6.

**S7.6-5.** Schuller JL, Zippe M, Crawford T, et al. Implantable cardioverter defibrillator therapy in patients with cardiac sarcoidosis. *J Cardiovasc Electrophysiol.* 2012;23:925-9.

**S7.6-6.** Coleman GC, Shaw PW, Balfour PC Jr., et al. Prognostic value of myocardial scarring on CMR in patients with cardiac sarcoidosis: a systematic review and meta-analysis. *J Am Coll Cardiol Img.* 2016;10:411-20.

**S7.6-7.** Murtagh G, Laffin LJ, Beshai JF, et al. Prognosis of myocardial damage in sarcoidosis patients with preserved left ventricular ejection fraction: risk stratification using cardiovascular magnetic resonance. *Circ Cardiovasc Imaging.* 2016;9:e003738.

**S7.6-8.** Crawford T, Mueller G, Sarsam S, et al. Magnetic resonance imaging for identifying patients with cardiac sarcoidosis and preserved or mildly reduced left ventricular function at risk of ventricular arrhythmias. *Circ Arrhythm Electrophysiol.* 2014;7:1109-15.

**S7.6-9.** Greulich S, Deluigi CC, Gloekler S, et al. CMR imaging predicts death and other adverse events in suspected cardiac sarcoidosis. *J Am Coll Cardiol Img.* 2013;6:501-11.

**S7.6-10.** Blankstein R, Osborne M, Naya M, et al. Cardiac positron emission tomography enhances prognostic assessments of patients with suspected cardiac sarcoidosis. *J Am Coll Cardiol.* 2014;63:329-36.

**S7.6-11.** Aizer A, Stern EH, Gomes JA, et al. Usefulness of programmed ventricular stimulation in predicting future arrhythmic events in patients with cardiac sarcoidosis. *Am J Cardiol.* 2005;96:276-82.

**S7.6-12.** Mehta RH, Starr AZ, Lopes RD, et al. Incidence of and outcomes associated with ventricular tachycardia or fibrillation in patients undergoing primary percutaneous coronary intervention. *JAMA.* 2009;301:1779-89.

**S7.6-13.** Takaya Y, Kusano KF, Nakamura K, et al. Outcomes in patients with high-degree atrioventricular block as the initial manifestation of cardiac sarcoidosis. *Am J Cardiol.* 2015;115:505-9.

**S7.6-14.** Naruse Y, Sekiguchi Y, Nogami A, et al. Systematic treatment approach to ventricular tachycardia in cardiac sarcoidosis. *Circ Arrhythm Electrophysiol.* 2014;7:407-13.

**S7.6-15.** Segawa M, Fukuda K, Nakano M, et al. Time course and factors correlating with ventricular tachyarrhythmias after introduction of steroid therapy in cardiac sarcoidosis. *Circ Arrhythm Electrophysiol.* 2016;9:e003353.

**S7.6-16.** Yodogawa K, Seino Y, Ohara T, et al. Effect of corticosteroid therapy on ventricular arrhythmias in patients with cardiac sarcoidosis. *Ann Noninvasive Electrocardiol.* 2011;16:140-7.

**S7.6-17.** Birnie DH, Sauer WH, Bogun F, et al. HRS expert consensus statement on the diagnosis and management of arrhythmias associated with cardiac sarcoidosis. *Heart Rhythm.* 2014;11:1305–23.

**S7.6-18.** Mehta D, Mori N, Goldborg SH, et al. Primary prevention of sudden cardiac death in silent cardiac sarcoidosis: role of programmed ventricular stimulation. *Circ Arrhythm Electrophysiol.* 2011;4:43–8.

#### 7.6.1. Other Infiltrative Cardiomyopathies

**S7.6.1-1.** Lubitz SA, Goldborg SH, Mehta D. Sudden cardiac death in infiltrative cardiomyopathies: sarcoidosis, scleroderma, amyloidosis, hemachromatosis. *Prog Cardiovasc Dis.* 2008;51:58–73.

**S7.6.1-2.** Kristen AV, Dengler TJ, Hegenbart U, et al. Prophylactic implantation of cardioverter-defibrillator in patients with severe cardiac amyloidosis and high risk for sudden cardiac death. *Heart Rhythm.* 2008;5:235–40.

**S7.6.1-3.** Sperry BW, Ikram A, Hachamovitch R, et al. Efficacy of chemotherapy for light-chain amyloidosis in patients presenting with symptomatic heart failure. *J Am Coll Cardiol.* 2016;67:2941–8.

**S7.6.1-4.** Varr BC, Zarafshar S, Coakley T, et al. Implantable cardioverter-defibrillator placement in patients with cardiac amyloidosis. *Heart Rhythm.* 2014;11:158–62.

### 7.7. Heart Failure

#### 7.7.1. HF With Reduced Ejection Fraction

**S7.7.1-1.** Frohlich GM, Holzmeister J, Hubler M, et al. Prophylactic implantable cardioverter defibrillator treatment in patients with end-stage heart failure awaiting heart transplantation. *Heart.* 2013;99:1158–65.

**S7.7.1-2.** Kao AC, Krause SW, Handa R, et al. Wearable defibrillator use in heart failure (WIF): results of a prospective registry. *BMC Cardiovasc Disord.* 2012;12:123.

**S7.7.1-3.** Opreanu M, Wan C, Singh V, et al. Wearable cardioverter-defibrillator as a bridge to cardiac transplantation: a national database analysis. *J Heart Lung Transplant.* 2015;34:1305–9.

**S7.7.1-4.** Sandner SE, Wieselthaler G, Zuckermann A, et al. Survival benefit of the implantable cardioverter-defibrillator in patients on the waiting list for cardiac transplantation. *Circulation.* 2001;104:171–6.

**S7.7.1-5.** Vakil K, Duval S, Cogswell R, et al. Impact of implantable cardioverter-defibrillators on waitlist mortality among patients awaiting heart transplantation. An UNOS/OPTN Analysis. *JACC Clin Electrophysiol.* 2017;3:33–40.

**S7.7.1-6.** Packer DL, Prutkin JM, Hellkamp AS, et al. Impact of implantable cardioverter-defibrillator, amiodarone, and placebo on the mode of death in stable patients with heart failure: analysis from the sudden cardiac death in heart failure trial. *Circulation.* 2009;120:2170–6.

**S7.7.1-7.** MERIT-HF Study. Effect of metoprolol CR/XL in chronic heart failure: Metoprolol CR/XL Randomised Intervention Trial in Congestive Heart Failure (MERIT-HF). *Lancet.* 1999;353:2001–7.

**S7.7.1-8.** Luu M, Stevenson WG, Stevenson LW, et al. Diverse mechanisms of unexpected cardiac arrest in advanced heart failure. *Circulation.* 1989;80:1675–80.

**S7.7.1-9.** Narang R, Cleland JG, Erhardt L, et al. Mode of death in chronic heart failure. A request and proposition for more accurate classification. *Eur Heart J.* 1996;17:1390–403.

**S7.7.1-10.** Bonet S, Agusti A, Arnau JM, et al. Beta-adrenergic blocking agents in heart failure: benefits of vasodilating and non-vasodilating agents according to patients' characteristics: a meta-analysis of clinical trials. *Arch Intern Med.* 2000;160:621–7.

**S7.7.1-11.** Pitt B, Remme W, Zannad F, et al. Eplerenone, a selective aldosterone blocker, in patients with left ventricular dysfunction after myocardial infarction. *N Engl J Med.* 2003;348:1309–21.

**S7.7.1-12.** Desai AS, McMurray JJ, Packer M, et al. Effect of the angiotensin-receptor-neprilysin inhibitor LCZ696 compared with enalapril on mode of death in heart failure patients. *Eur Heart J.* 2015;36:1990–7.

**S7.7.1-13.** Cleland JG, Daubert JC, Erdmann E, et al. The effect of cardiac resynchronization on morbidity and mortality in heart failure. *N Engl J Med.* 2005;352:1539–49.

**S7.7.1-14.** Tomaselli GF, Zipes DP. What causes sudden death in heart failure? *Circ Res.* 2004;95:754–63.

**S7.7.1-15.** Friedlander Y, Siscovich DS, Weinmann S, et al. Family history as a risk factor for primary cardiac arrest. *Circulation.* 1998;97:155–60.

**S7.7.1-16.** Gloschat CR, Koppel AC, Aras KK, et al. Arrhythmogenic and metabolic remodelling of failing human heart. *J Physiol.* 2016;594:3963–80.

**S7.7.1-17.** Vakil K, Kazmirczak F, Sathnur N, et al. Implantable cardioverter-defibrillator use in patients with left ventricular assist devices: a systematic review and meta-analysis. *JACC Heart Fail.* 2016;4:772–9.

#### 7.7.2. HF With Preserved Ejection Fraction

**S7.7.2-1.** Owan TE, Hodge DO, Herges RM, et al. Trends in prevalence and outcome of heart failure with preserved ejection fraction. *N Engl J Med.* 2006;355:251–9.

**S7.7.2-2.** Adabag S, Smith LG, Anand IS, et al. Sudden cardiac death in heart failure patients with preserved ejection fraction. *J Card Fail.* 2012;18:749–54.

**S7.7.2-3.** Massie BM, Carson PE, McMurray JJ, et al. Irbesartan in patients with heart failure and preserved ejection fraction. *N Engl J Med.* 2008;359:2456–67.

**S7.7.2-4.** Yusuf S, Pfeffer MA, Swedberg K, et al. Effects of candesartan in patients with chronic heart failure and preserved left-ventricular ejection fraction: the CHARM-Preserved Trial. *Lancet.* 2003;362:777–81.

**S7.7.2-5.** Pitt B, Pfeffer MA, Assmann SF, et al. Spironolactone for heart failure with preserved ejection fraction. *N Engl J Med.* 2014;370:1383–92.

**S7.7.2-6.** Al-Khatib SM, Shaw LK, O'Connor C, et al. Incidence and predictors of sudden cardiac death in patients with diastolic heart failure. *J Cardiovasc Electrophysiol.* 2007;18:1231–5.

**S7.7.2-7.** Gatzoulis MA, Till JA, Somerville J, et al. Mechanoelectrical interaction in tetralogy of Fallot. QRS prolongation relates to right ventricular size and predicts malignant ventricular arrhythmias and sudden death. *Circulation.* 1995;92:231–7.

**S7.7.2-8.** Gatzoulis KA, Tsachiris D, Arsenos P, et al. Prognostic value of programmed ventricular stimulation for sudden death in selected high risk patients

with structural heart disease and preserved systolic function. *Int J Cardiol.* 2014;176:1449–51.

#### 7.7.3. Left Ventricular Assist Device

**S7.7.3-1.** Vakil K, Kazmirczak F, Sathnur N, et al. Implantable cardioverter-defibrillator use in patients with left ventricular assist devices: a systematic review and meta-analysis. *JACC Heart Fail.* 2016;4:772–9.

**S7.7.3-2.** Cantillon DJ, Tarakji KG, Kumbhani DJ, et al. Improved survival among ventricular assist device recipients with a concomitant implantable cardioverter-defibrillator. *Heart Rhythm.* 2010;7:466–71.

**S7.7.3-3.** Raasch H, Jensen BC, Chang PP, et al. Epidemiology, management, and outcomes of sustained ventricular arrhythmias after continuous-flow left ventricular assist device implantation. *Am Heart J.* 2012;164:373–8.

**S7.7.3-4.** Garan AR, Yuzefpolskaya M, Colombo PC, et al. Ventricular arrhythmias and implantable cardioverter-defibrillator therapy in patients with continuous-flow left ventricular assist devices: need for primary prevention? *J Am Coll Cardiol.* 2013;61:2542–50.

**S7.7.3-5.** Refaat MM, Tanaka T, Kormos RL, et al. Survival benefit of implantable cardioverter-defibrillators in left ventricular assist device-supported heart failure patients. *J Card Fail.* 2012;18:140–5.

**S7.7.3-6.** Enriquez AD, Calenda B, Miller MA, et al. The role of implantable cardioverter-defibrillators in patients with continuous flow left ventricular assist devices. *Circ Arrhythm Electrophysiol.* 2013;6:668–74.

**S7.7.3-7.** Lee W, Tay A, Subbiah RN, et al. Impact of implantable cardioverter defibrillators on survival of patients with centrifugal left ventricular assist devices. *Pacing Clin Electrophysiol.* 2015;38:925–33.

**S7.7.3-8.** Jorde UP, Kushwaha SS, Tatooles AJ, et al. Results of the destination therapy post-food and drug administration approval study with a continuous flow left ventricular assist device: a prospective study using the INTERMACS registry (Interagency Registry for Mechanically Assisted Circulatory Support). *J Am Coll Cardiol.* 2014;63:1751–7.

**S7.7.3-9.** Slaughter MS, Meyer AL, Birks EJ. Destination therapy with left ventricular assist devices: patient selection and outcomes. *Curr Opin Cardiol.* 2011;26:232–6.

#### 7.7.4. ICD Use After Heart Transplantation

**S7.7.4-1.** Tsai VW, Cooper J, Garan H, et al. The efficacy of implantable cardioverter-defibrillators in heart transplant recipients: results from a multicenter registry. *Circ Heart Fail.* 2009;2:197–201.

**S7.7.4-2.** McDowell DL, Hauptman PJ. Implantable defibrillators and cardiac resynchronization therapy in heart transplant recipients: results of a national survey. *J Heart Lung Transplant.* 2009;28:847–50.

**S7.7.4-3.** Neylon A, Canniffe C, Parlon B, et al. Implantable cardioverter-defibrillators in a heart transplant population: A single-center experience. *J Heart Lung Transplant.* 2016;35:682–4.

**S7.7.4-4.** Vakil K, Taimeh Z, Sharma A, et al. Incidence, predictors, and temporal trends of sudden cardiac

death after heart transplantation. *Heart Rhythm.* 2014; 11:1684-90.

**S7.7.4-5.** Vaseghi M, Lelouche N, Ritter H, et al. Mode and mechanisms of death after orthotopic heart transplantation. *Heart Rhythm.* 2009;6:503-9.

## 7.8. Neuromuscular Disorders

**S7.8-1.** Bhakta D, Groh MR, Shen C, et al. Increased mortality with left ventricular systolic dysfunction and heart failure in adults with myotonic dystrophy type 1. *Am Heart J.* 2010;160:1137-41, 41.

**S7.8-2.** Merino JL, Carmona JR, Fernandez-Lozano I, et al. Mechanisms of sustained ventricular tachycardia in myotonic dystrophy: implications for catheter ablation. *Circulation.* 1998;98:541-6.

**S7.8-3.** Anselme F, Moubarak G, Savoure A, et al. Implantable cardioverter-defibrillators in lamin A/C mutation carriers with cardiac conduction disorders. *Heart Rhythm.* 2013;10:1492-8.

**S7.8-4.** van Rijssingen IA, Arbustini E, Elliott PM, et al. Risk factors for malignant ventricular arrhythmias in lamin A/C mutation carriers: a European cohort study. *J Am Coll Cardiol.* 2012;59:493-500.

**S7.8-5.** Meune C, Van Berlo JH, Anselme F, et al. Primary prevention of sudden death in patients with lamin A/C gene mutations. *N Engl J Med.* 2006;354:209-10.

**S7.8-6.** Pasotti M, Klersy C, Pilotto A, et al. Long-term outcome and risk stratification in dilated cardiomyopathies. *J Am Coll Cardiol.* 2008;52:1250-60.

**S7.8-7.** Van Berlo JH, de Voogt WG, van der Kooi AJ, et al. Meta-analysis of clinical characteristics of 299 carriers of LMNA gene mutations: do lamin A/C mutations portend a high risk of sudden death? *J Mol Med.* 2005;83:79-83.

**S7.8-8.** Russo V, Nigro G. ICD role in preventing sudden cardiac death in Emery-Dreifuss muscular dystrophy with preserved myocardial function: 2013 ESC guidelines on cardiac pacing and cardiac resynchronization therapy. *Europace.* 2015;17:337.

**S7.8-9.** Groh WJ, Groh MR, Saha C, et al. Electrocardiographic abnormalities and sudden death in myotonic dystrophy type 1. *N Engl J Med.* 2008; 358:2688-97.

**S7.8-10.** Lallemand B, Clementy N, Bernard-Brunet A, et al. The evolution of infrahissian conduction time in myotonic dystrophy patients: clinical implications. *Heart.* 2012;98:291-6.

**S7.8-11.** Nazarian S, Wagner KR, Caffo BS, et al. Clinical predictors of conduction disease progression in type I myotonic muscular dystrophy. *Pacing Clin Electrophysiol.* 2011;34:171-6.

**S7.8-12.** Tanawuttiwat T, Wagner KR, Tomaselli G, et al. Left ventricular dysfunction and conduction disturbances in patients with myotonic muscular dystrophy type I and II. *JAMA Cardiology.* 2017;2:225-8.

**S7.8-13.** Bhakta D, Shen C, Kron J, et al. Pacemaker and implantable cardioverter-defibrillator use in a US myotonic dystrophy type 1 population. *J Cardiovasc Electrophysiol.* 2011;22:1369-75.

**S7.8-14.** Laurent V, Pellieux S, Corcia P, et al. Mortality in myotonic dystrophy patients in the area of prophylactic pacing devices. *Int J Cardiol.* 2011;150:54-8.

**S7.8-15.** Groh WJ. Arrhythmias in the muscular dystrophies. *Heart Rhythm.* 2012;9:1890-5.

**S7.8-16.** Epstein AE, DiMarco JP, Ellenbogen KA, et al. ACC/AHA/HRS 2008 guidelines for device-based therapy of cardiac rhythm abnormalities: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines (Writing Committee to Revise the ACC/AHA/NASPE 2002 Guideline Update for Implantation of Cardiac Pacemakers and Antiarrhythmia Devices) developed in collaboration with the American Association for Thoracic Surgery and Society of Thoracic Surgeons. *J Am Coll Cardiol.* 2008;51:e1-62.

**S7.8-17.** Kumar S, Baldinger SH, Gandjbakhch E, et al. Long-term arrhythmic and nonarrhythmic outcomes of lamin A/C mutation carriers. *J Am Coll Cardiol.* 2016; 68:2299-307.

**S7.8-18.** Bremilla-Perrot B, Luporsi JD, Louis S, et al. Long-term follow-up of patients with myotonic dystrophy: an electrocardiogram every year is not necessary. *Europace.* 2011;13:251-7.

**S7.8-19.** Ha AH, Tarnopolsky MA, Bergstra TG, et al. Predictors of atrio-ventricular conduction disease, long-term outcomes in patients with myotonic dystrophy types I and II. *Pacing Clin Electrophysiol.* 2012; 35:1262-9.

## 7.9. Cardiac Channelopathies

**S7.9-1.** Bai R, Napolitano C, Bloise R, et al. Yield of genetic screening in inherited cardiac channelopathies: how to prioritize access to genetic testing. *Circ Arrhythm Electrophysiol.* 2009;2:6-15.

**S7.9-2.** Goldenberg I, Horr S, Moss AJ, et al. Risk for life-threatening cardiac events in patients with genotype-confirmed long-QT syndrome and normal-range corrected QT intervals. *J Am Coll Cardiol.* 2011; 57:51-9.

**S7.9-3.** Nannenberg EA, Sijbrands EJ, Dijksman LM, et al. Mortality of inherited arrhythmia syndromes: insight into their natural history. *Circ Cardiovasc Genet.* 2012;5:183-9.

**S7.9-4.** Priori SG, Gasparini M, Napolitano C, et al. Risk stratification in Brugada syndrome: results of the PRELUDE (Programmed Electrical stimUlation preDicte valuE) registry. *J Am Coll Cardiol.* 2012; 59:37-45.

**S7.9-5.** Priori SG, Napolitano C, Memmi M, et al. Clinical and molecular characterization of patients with catecholaminergic polymorphic ventricular tachycardia. *Circulation.* 2002;106:69-74.

**S7.9-6.** Wilde AA, Moss AJ, Kaufman ES, et al. Clinical aspects of type 3 long-QT syndrome: an international multicenter study. *Circulation.* 2016;134:872-82.

**S7.9-7.** Wedekind H, Burde D, Zumhagen S, et al. QT interval prolongation and risk for cardiac events in genotyped LQTS-index children. *Eur J Pediatr.* 2009; 168:1107-15.

**S7.9-8.** Moss AJ, Zareba W, Hall WJ, et al. Effectiveness and limitations of beta-blocker therapy in congenital long-QT syndrome. *Circulation.* 2000;101: 616-23.

**S7.9-9.** Zareba W, Moss AJ, Daubert JP, et al. Implantable cardioverter defibrillator in high-risk long QT syndrome patients. *J Cardiovasc Electrophysiol.* 2003;14:337-41.

**S7.9-10.** Monnig G, Kobe J, Loher A, et al. Implantable cardioverter-defibrillator therapy in patients with congenital long-QT syndrome: a long-term follow-up. *Heart Rhythm.* 2005;2:497-504.

**S7.9-11.** Hayashi M, Denjoy I, Extramiana F, et al. Incidence and risk factors of arrhythmic events in catecholaminergic polymorphic ventricular tachycardia. *Circulation.* 2009;119:2426-34.

**S7.9-12.** Gehi AK, Duong TD, Metz LD, et al. Risk stratification of individuals with the Brugada electrocardiogram: a meta-analysis. *J Cardiovasc Electrophysiol.* 2006;17:577-83.

**S7.9-13.** Probst V, Veltmann C, Eckardt L, et al. Long-term prognosis of patients diagnosed with Brugada syndrome: results from the FINGER Brugada syndrome registry. *Circulation.* 2010;121:635-43.

**S7.9-14.** Delise P, Allocca G, Marras E, et al. Risk stratification in individuals with the Brugada type 1 ECG pattern without previous cardiac arrest: usefulness of a combined clinical and electrophysiologic approach. *Eur Heart J.* 2011;32:169-76.

**S7.9-15.** Hiraoka M, Takagi M, Yokoyama Y, et al. Prognosis and risk stratification of young adults with Brugada syndrome. *J Electrocardiol.* 2013;46:279-83.

**S7.9-16.** Kaufman ES, McNitt S, Moss AJ, et al. Risk of death in the long QT syndrome when a sibling has died. *Heart Rhythm.* 2008;5:831-6.

**S7.9-17.** Kimbrough J, Moss AJ, Zareba W, et al. Clinical implications for affected parents and siblings of probands with long-QT syndrome. *Circulation.* 2001; 104:557-62.

**S7.9-18.** Priori SG, Napolitano C, Gasparini M, et al. Natural history of Brugada syndrome: insights for risk stratification and management. *Circulation.* 2002;105: 1342-7.

**S7.9-19.** Hobbs JB, Peterson DR, Moss AJ, et al. Risk of aborted cardiac arrest or sudden cardiac death during adolescence in the long-QT syndrome. *JAMA.* 2006; 296:1249-54.

**S7.9-20.** Priori SG, Napolitano C, Schwartz PJ, et al. Association of long QT syndrome loci and cardiac events among patients treated with beta-blockers. *JAMA.* 2004;292:1341-4.

**S7.9-21.** Priori SG, Schwartz PJ, Napolitano C, et al. Risk stratification in the long-QT syndrome. *N Engl J Med.* 2003;348:1866-74.

**S7.9-22.** Spazzolini C, Mullally J, Moss AJ, et al. Clinical implications for patients with long QT syndrome who experience a cardiac event during infancy. *J Am Coll Cardiol.* 2009;54:832-7.

**S7.9-23.** Credible meds. Available at: <http://www.crediblemeds.org>. Accessed December 26, 2016.

**S7.9-24.** Paludan-Muller C, Ahlberg G, Ghose J, et al. Integration of 60,000 exomes and ACMG guidelines question the role of catecholaminergic polymorphic ventricular tachycardia-associated variants. *Clin Genet.* 2017;91:63-72.

**S7.9-25.** Giustetto C, Schimpf R, Mazzanti A, et al. Long-term follow-up of patients with short QT syndrome. *J Am Coll Cardiol.* 2011;58:587-95.

**S7.9-26.** Mahida S, Derval N, Sacher F, et al. Role of electrophysiological studies in predicting risk of ventricular arrhythmias in early repolarization syndrome. *J Am Coll Cardiol.* 2015;65:151-9.

**S7.9-27.** Mazzanti A, Kanthan A, Monteforte N, et al. Novel insight into the natural history of short QT syndrome. *J Am Coll Cardiol.* 2014;63:1300-8.

**S7.9-28.** Siebermair J, Sinner MF, Beckmann BM, et al. Early repolarization pattern is the strongest predictor

of arrhythmia recurrence in patients with idiopathic ventricular fibrillation: results from a single centre long-term follow-up over 20 years. *Europace*. 2016; 18:718–25.

**S7.9-29.** Etheridge SP, Sanatani S, Cohen MI, et al. Long QT syndrome in children in the era of implantable defibrillators. *J Am Coll Cardiol*. 2007;50:1335–40.

**S7.9-30.** Horner JM, Kinoshita M, Webster TL, et al. Implantable cardioverter defibrillator therapy for congenital long QT syndrome: a single-center experience. *Heart Rhythm*. 2010;7:1616–22.

**S7.9-31.** Proclemer A, Ghidina M, Facchini D, et al. Use of implantable cardioverter-defibrillator in inherited arrhythmogenic diseases: data from Italian ICD Registry for the years 2001–6. *Pacing Clin Electrophysiol*. 2009;32:434–45.

**S7.9-32.** Adler A, Sadek MM, Chan AY, et al. Patient outcomes from a specialized inherited arrhythmia clinic. *Circ Arrhythm Electrophysiol*. 2016;9:e003440.

**S7.9-33.** Conte G, Sieira J, Cionte G, et al. Implantable cardioverter-defibrillator therapy in Brugada syndrome: a 20-year single-center experience. *J Am Coll Cardiol*. 2015;65:879–88.

**S7.9-34.** Olde Nordkamp LR, Postema PG, Knops RE, et al. Implantable cardioverter-defibrillator harm in young patients with inherited arrhythmia syndromes: a systematic review and meta-analysis of inappropriate shocks and complications. *Heart Rhythm*. 2016;13: 443–54.

**S7.9-35.** Rodriguez-Manero M, Sacher F, de Asmundis C, et al. Monomorphic ventricular tachycardia in patients with Brugada syndrome: a multi-center retrospective study. *Heart Rhythm*. 2016;13: 669–82.

**S7.9-36.** Rosso R, Glick A, Glikson M, et al. Outcome after implantation of cardioverter defibrillator [corrected] in patients with Brugada syndrome: a multi-center Israeli study (ISRABRU). *Isr Med Assoc J*. 2008; 10:435–9.

**S7.9-37.** Dorostkar PC, Eldar M, Belhassen B, et al. Long-term follow-up of patients with long-QT syndrome treated with beta-blockers and continuous pacing. *Circulation*. 1999;100:2431–6.

**S7.9-38.** Eldar M, Griffin JC, Van Hare GF, et al. Combined use of beta-adrenergic blocking agents and long-term cardiac pacing for patients with the long QT syndrome. *J Am Coll Cardiol*. 1992;20:830–7.

**S7.9-39.** Moss AJ, Liu JE, Gottlieb S, et al. Efficacy of permanent pacing in the management of high-risk patients with long QT syndrome. *Circulation*. 1991; 84:1524–9.

**S7.9-40.** Vincent GM, Schwartz PJ, Denjoy I, et al. High efficacy of beta-blockers in long-QT syndrome type 1: contribution of noncompliance and QT-prolonging drugs to the occurrence of beta-blocker treatment "failures". *Circulation*. 2009;119:215–21.

## 7.9.1. Specific Cardiac Channelopathy Syndromes

### 7.9.1.1. Congenital Long QT Syndrome

**S7.9.1.1-1.** Abu-Zetone A, Peterson DR, Polonsky B, et al. Efficacy of different beta-blockers in the treatment of long QT syndrome. *J Am Coll Cardiol*. 2014; 64:1352–8.

**S7.9.1.1-2.** Goldenberg I, Bradley J, Moss A, et al. Beta-blocker efficacy in high-risk patients with the

congenital long-QT syndrome types 1 and 2: implications for patient management. *J Cardiovasc Electrophysiol*. 2010;21:893–901.

**S7.9.1.1-3.** Moss AJ, Zareba W, Hall WJ, et al. Effectiveness and limitations of beta-blocker therapy in congenital long-QT syndrome. *Circulation*. 2000;101: 616–23.

**S7.9.1.1-4.** Sauer AJ, Moss AJ, McNitt S, et al. Long QT syndrome in adults. *J Am Coll Cardiol*. 2007;49: 329–37.

**S7.9.1.1-5.** Vincent GM, Schwartz PJ, Denjoy I, et al. High efficacy of beta-blockers in long-QT syndrome type 1: contribution of noncompliance and QT-prolonging drugs to the occurrence of beta-blocker treatment "failures". *Circulation*. 2009;119:215–21.

**S7.9.1.1-6.** Chorin E, Hu D, Antzelevitch C, et al. Ranolazine for congenital long-QT syndrome type III: experimental and long-term clinical data. *Circ Arrhythm Electrophysiol*. 2016;9:e004370.

**S7.9.1.1-7.** Chorin E, Taub R, Medina A, et al. Long-term flecainide therapy in type 3 long QT syndrome. *Europace*. 2017; euw439 [Epub ahead of print].

**S7.9.1.1-8.** Hobbs JB, Peterson DR, Moss AJ, et al. Risk of aborted cardiac arrest or sudden cardiac death during adolescence in the long-QT syndrome. *JAMA*. 2006;296:1249–54.

**S7.9.1.1-9.** Jons C, Moss AJ, Goldenberg I, et al. Risk of fatal arrhythmic events in long QT syndrome patients after syncope. *J Am Coll Cardiol*. 2010;55:783–8.

**S7.9.1.1-10.** Mazzanti A, Maragna R, Faragli A, et al. Gene-specific therapy with mexiletine reduces arrhythmic events in patients with long QT syndrome type 3. *J Am Coll Cardiol*. 2016;67:1053–8.

**S7.9.1.1-11.** Nannenberg EA, Sijbrands EJ, Dijkman LM, et al. Mortality of inherited arrhythmia syndromes: insight into their natural history. *Circ Cardiovasc Genet*. 2012;5:183–9.

**S7.9.1.1-12.** Wedekind H, Burde D, Zumhagen S, et al. QT interval prolongation and risk for cardiac events in genotyped LQTS-index children. *Eur J Pediatr*. 2009; 168:1107–15.

**S7.9.1.1-13.** Collura CA, Johnson JN, Moir C, et al. Left cardiac sympathetic denervation for the treatment of long QT syndrome and catecholaminergic polymorphic ventricular tachycardia using video-assisted thoracic surgery. *Heart Rhythm*. 2009;6:752–9.

**S7.9.1.1-14.** Hofferberth SC, Cecchin F, Lobman D, et al. Left thoracoscopic sympathectomy for cardiac denervation in patients with life-threatening ventricular arrhythmias. *J Thorac Cardiovasc Surg*. 2014;147: 404–9.

**S7.9.1.1-15.** Schneider HE, Steinmetz M, Krause U, et al. Left cardiac sympathetic denervation for the management of life-threatening ventricular tachyarrhythmias in young patients with catecholaminergic polymorphic ventricular tachycardia and long QT syndrome. *Clin Res Cardiol*. 2013;102:33–42.

**S7.9.1.1-16.** Schwartz PJ, Priori SG, Cerrone M, et al. Left cardiac sympathetic denervation in the management of high-risk patients affected by the long-QT syndrome. *Circulation*. 2004;109:1826–33.

**S7.9.1.1-17.** Bai R, Napolitano C, Bloise R, et al. Yield of genetic screening in inherited cardiac channelopathies: how to prioritize access to genetic testing. *Circ Arrhythm Electrophysiol*. 2009;2:6–15.

**S7.9.1.1-18.** Costa J, Lopes CM, Barsheshet A, et al. Combined assessment of sex- and mutation-specific information for risk stratification in type 1 long QT syndrome. *Heart Rhythm*. 2012;9:892–8.

**S7.9.1.1-19.** Kim JA, Lopes CM, Moss AJ, et al. Trigger-specific risk factors and response to therapy in long QT syndrome type 2. *Heart Rhythm*. 2010;7:1797–805.

**S7.9.1.1-20.** Migdalovich D, Moss AJ, Lopes CM, et al. Mutation and gender-specific risk in type 2 long QT syndrome: implications for risk stratification for life-threatening cardiac events in patients with long QT syndrome. *Heart Rhythm*. 2011;8:1537–43.

**S7.9.1.1-21.** Tester DJ, Will ML, Haglund CM, et al. Effect of clinical phenotype on yield of long QT syndrome genetic testing. *J Am Coll Cardiol*. 2006;47: 764–8.

**S7.9.1.1-22.** Adler A, van der Werf C, Postema PG, et al. The phenomenon of "QT stunning": The abnormal QT prolongation provoked by standing persists even as the heart rate returns to normal in patients with long QT syndrome. *Heart Rhythm*. 2012;9:901–8.

**S7.9.1.1-23.** Aziz PF, Wieand TS, Ganley J, et al. Genotype- and mutation site-specific QT adaptation during exercise, recovery, and postural changes in children with long-QT syndrome. *Circ Arrhythm Electrophysiol*. 2011;4:867–73.

**S7.9.1.1-24.** Chattha IS, Sy RW, Yee R, et al. Utility of the recovery electrocardiogram after exercise: a novel indicator for the diagnosis and genotyping of long QT syndrome? *Heart Rhythm*. 2010;7:906–11.

**S7.9.1.1-25.** Laksman ZW, Hamilton RM, Chockalingam P, et al. Mutation location effect on severity of phenotype during exercise testing in type 1 long-QT syndrome: impact of transmembrane and C-loop location. *J Cardiovasc Electrophysiol*. 2013;24: 1015–20.

**S7.9.1.1-26.** Moltedo JM, Kim JJ, Friedman RA, et al. Use of a cardioselective beta-blocker for pediatric patients with prolonged QT syndrome. *Pediatr Cardiol*. 2011;32:63–6.

**S7.9.1.1-27.** Sy RW, van der Werf C, Chattha IS, et al. Derivation and validation of a simple exercise-based algorithm for prediction of genetic testing in relatives of LQTS probands. *Circulation*. 2011;124:2187–94.

**S7.9.1.1-28.** Villain E, Denjoy I, Lupoglazoff JM, et al. Low incidence of cardiac events with beta-blocking therapy in children with long QT syndrome. *Eur Heart J*. 2004;25:1405–11.

**S7.9.1.1-29.** Viskin S, Postema PG, Bhuiyan ZA, et al. The response of the QT interval to the brief tachycardia provoked by standing: a bedside test for diagnosing long QT syndrome. *J Am Coll Cardiol*. 2010;55: 1955–61.

**S7.9.1.1-30.** Priori SG, Napolitano C, Schwartz PJ, et al. Association of long QT syndrome loci and cardiac events among patients treated with beta-blockers. *JAMA*. 2004;292:1341–4.

**S7.9.1.1-31.** Goldenberg I, Horr S, Moss AJ, et al. Risk for life-threatening cardiac events in patients with genotype-confirmed long-QT syndrome and normal-range corrected QT intervals. *J Am Coll Cardiol*. 2011; 57:51–9.

**S7.9.1.1-32.** Choy AM, Lang CC, Chomsky DM, et al. Normalization of acquired QT prolongation in humans by intravenous potassium. *Circulation*. 1997; 96:2149–54.

- S7.9.1.1-33.** Kannankeril P, Roden DM, Darbar D. Drug-induced long QT syndrome. *Pharmacol Rev.* 2010;62:760–81.
- S7.9.1.1-34.** Zhang C, Kutyifa V, Moss AJ, et al. Long-QT syndrome and therapy for attention deficit/hyperactivity disorder. *J Cardiovasc Electrophysiol.* 2015;26:1039–44.
- S7.9.1.1-35.** Priori SG, Schwartz PJ, Napolitano C, et al. Risk stratification in the long-QT syndrome. *N Engl J Med.* 2003;348:1866–74.
- S7.9.1.1-36.** Wilde AA, Moss AJ, Kaufman ES, et al. Clinical aspects of type 3 long-QT syndrome: an international multicenter study. *Circulation.* 2016;134:872–82.
- S7.9.1.1-37.** Schwartz PJ, Spazzolini C, Crotti L. All LQT3 patients need an ICD: true or false? *Heart Rhythm.* 2009;6:113–20.
- S7.9.1.1-38.** Garson A Jr, Dick M, Fournier A, et al. The long QT syndrome in children. An international study of 287 patients. *Circulation.* 1993;87:1866–72.
- S7.9.1.1-39.** Priori SG, Bossaert LL, Chamberlain DA, et al. ESC-ERC recommendations for the use of automated external defibrillators (AEDs) in Europe. *Eur Heart J.* 2004;25:437–45.
- S7.9.1.1-40.** Steinberg C, Padfield GJ, Al-Sabiq B, et al. Experience with bisoprolol in long-QT1 and long-QT2 syndrome. *J Interv Card Electrophysiol.* 2016;47:163–70.
- S7.9.1.1-41.** Chockalingam P, Crotti L, Girardengo G, et al. Not all beta-blockers are equal in the management of long QT syndrome types 1 and 2: higher recurrence of events under metoprolol. *J Am Coll Cardiol.* 2012;60:2092–9.
- S7.9.1.1-42.** Khositseth A, Tester DJ, Will ML, et al. Identification of a common genetic substrate underlying postpartum cardiac events in congenital long QT syndrome. *Heart Rhythm.* 2004;1:60–4.
- S7.9.1.1-43.** Rashba EJ, Zareba W, Moss AJ, et al. Influence of pregnancy on the risk for cardiac events in patients with hereditary long QT syndrome. LQTS Investigators. *Circulation.* 1998;97:451–6.
- S7.9.1.1-44.** Dorostkar PC, Eldar M, Belhassen B, et al. Long-term follow-up of patients with long-QT syndrome treated with beta-blockers and continuous pacing. *Circulation.* 1999;100:2431–6.
- S7.9.1.1-45.** Eldar M, Griffin JC, Van Hare GF, et al. Combined use of beta-adrenergic blocking agents and long-term cardiac pacing for patients with the long QT syndrome. *J Am Coll Cardiol.* 1992;20:830–7.
- S7.9.1.1-46.** Moss AJ, Liu JE, Gottlieb S, et al. Efficacy of permanent pacing in the management of high-risk patients with long QT syndrome. *Circulation.* 1991;84:1524–9.
- S7.9.1.1-47.** Viskin S, Glikson M, Fish R, et al. Rate smoothing with cardiac pacing for preventing torsade de pointes. *Am J Cardiol.* 2000;86:111k–5k.
- S7.9.1.1-48.** Bos JM, Bos KM, Johnson JN, et al. Left cardiac sympathetic denervation in long QT syndrome: analysis of therapeutic nonresponders. *Circ Arrhythm Electrophysiol.* 2013;6:705–11.
- S7.9.1.1-49.** Wilde AA, Bhuiyan ZA, Crotti L, et al. Left cardiac sympathetic denervation for catecholaminergic polymorphic ventricular tachycardia. *N Engl J Med.* 2008;358:2024–9.
- S7.9.1.1-50.** Li J, Liu Y, Yang F, et al. Video-assisted thoracoscopic left cardiac sympathetic denervation: a reliable minimally invasive approach for congenital long-QT syndrome. *Ann Thorac Surg.* 2008;86:1955–8.
- S7.9.1.1-51.** De Ferrari GM, Dusi V, Spazzolini C, et al. Clinical management of catecholaminergic polymorphic ventricular tachycardia: the role of left cardiac sympathetic denervation. *Circulation.* 2015;131:2185–93.
- S7.9.1.1-52.** Li C, Hu D, Shang L, et al. Surgical left cardiac sympathetic denervation for long QT syndrome: effects on QT interval and heart rate. *Heart Vessels.* 2005;20:137–41.
- S7.9.1.1-53.** Burns C, Ingles J, Davis AM, et al. Clinical and genetic features of Australian families with long QT syndrome: a registry-based study. *J Arrhythm.* 2016;32:456–61.
- S7.9.1.1-54.** Barsheshet A, Goldenberg I, Uchi J, et al. Mutations in cytoplasmic loops of the KCNQ1 channel and the risk of life-threatening events: implications for mutation-specific response to beta-blocker therapy in type 1 long-QT syndrome. *Circulation.* 2012;125:1988–96.
- S7.9.1.1-55.** Crotti L, Spazzolini C, Schwartz PJ, et al. The common long-QT syndrome mutation KCNQ1/A341V causes unusually severe clinical manifestations in patients with different ethnic backgrounds: toward a mutation-specific risk stratification. *Circulation.* 2007;116:2366–75.
- S7.9.1.1-56.** Zhang C, Kutyifa V, McNitt S, et al. Identification of low-risk adult congenital LQTS patients. *J Cardiovasc Electrophysiol.* 2015;26:853–8.
- S7.9.1.1-57.** Gaita F, Giustetto C, Bianchi F, et al. Short QT syndrome: pharmacological treatment. *J Am Coll Cardiol.* 2004;43:1494–9.
- S7.9.1.1-58.** Spazzolini C, Mullally J, Moss AJ, et al. Clinical implications for patients with long QT syndrome who experience a cardiac event during infancy. *J Am Coll Cardiol.* 2009;54:832–7.
- S7.9.1.1-59.** Credible meds. Available at: <http://www.crediblemeds.org>. Accessed December 26, 2016.
- S7.9.1.1-60.** Rohatgi RK, Bos JM, Ackerman MJ. Stimulant therapy in children with attention-deficit/hyperactivity disorder and concomitant long QT syndrome: a safe combination? *Heart Rhythm.* 2015;12:1807–12.
- S7.9.1.1-61.** Amin AS, Klemens CA, Verkerk AO, et al. Fever-triggered ventricular arrhythmias in Brugada syndrome and type 2 long-QT syndrome. *Neth Heart J.* 2010;18:165–9.
- S7.9.1.1-62.** Roden DM. Drug-induced prolongation of the QT interval. *N Engl J Med.* 2004;350:1013–22.
- 7.9.1.2. Catecholaminergic Polymorphic Ventricular Tachycardia**
- S7.9.1.2-1.** Hayashi M, Denjoy I, Extramiana F, et al. Incidence and risk factors of arrhythmic events in catecholaminergic polymorphic ventricular tachycardia. *Circulation.* 2009;119:2426–34.
- S7.9.1.2-2.** Roston TM, Vinocur JM, Maginot KR, et al. Catecholaminergic polymorphic ventricular tachycardia in children: analysis of therapeutic strategies and outcomes from an international multicenter registry. *Circ Arrhythm Electrophysiol.* 2015;8:633–42.
- S7.9.1.2-3.** Collura CA, Johnson JN, Moir C, et al. Left cardiac sympathetic denervation for the treatment of long QT syndrome and catecholaminergic polymorphic ventricular tachycardia using video-assisted thoracic surgery. *Heart Rhythm.* 2009;6:752–9.
- S7.9.1.2-4.** Hofferberth SC, Cecchin F, Lobeman D, et al. Left thoracoscopic sympathectomy for cardiac denervation in patients with life-threatening ventricular arrhythmias. *J Thorac Cardiovasc Surg.* 2014;147:404–9.
- S7.9.1.2-5.** Schneider HE, Steinmetz M, Krause U, et al. Left cardiac sympathetic denervation for the management of life-threatening ventricular tachyarrhythmias in young patients with catecholaminergic polymorphic ventricular tachycardia and long QT syndrome. *Clin Res Cardiol.* 2013;102:33–42.
- S7.9.1.2-6.** van der Werf C, Kannankeril PJ, Sacher F, et al. Flecainide therapy reduces exercise-induced ventricular arrhythmias in patients with catecholaminergic polymorphic ventricular tachycardia. *J Am Coll Cardiol.* 2011;57:2244–54.
- S7.9.1.2-7.** Priori SG, Napolitano C, Memmi M, et al. Clinical and molecular characterization of patients with catecholaminergic polymorphic ventricular tachycardia. *Circulation.* 2002;106:69–74.
- S7.9.1.2-8.** Leenhardt A, Lucet V, Denjoy I, et al. Catecholaminergic polymorphic ventricular tachycardia in children. A 7-year follow-up of 21 patients. *Circulation.* 1995;91:1512–9.
- S7.9.1.2-9.** Rosso R, Kalman JM, Rogowski O, et al. Calcium channel blockers and beta-blockers versus beta-blockers alone for preventing exercise-induced arrhythmias in catecholaminergic polymorphic ventricular tachycardia. *Heart Rhythm.* 2007;4:1149–54.
- S7.9.1.2-10.** Swan H, Laitinen P, Kontula K, et al. Calcium channel antagonism reduces exercise-induced ventricular arrhythmias in catecholaminergic polymorphic ventricular tachycardia patients with RyR2 mutations. *J Cardiovasc Electrophysiol.* 2005;16:162–6.
- S7.9.1.2-11.** Watanabe H, van der Werf C, Roses-Noguer F, et al. Effects of flecainide on exercise-induced ventricular arrhythmias and recurrences in genotype-negative patients with catecholaminergic polymorphic ventricular tachycardia. *Heart Rhythm.* 2013;10:542–7.
- S7.9.1.2-12.** Hwang HS, Hasdemir C, Laver D, et al. Inhibition of cardiac Ca<sup>2+</sup> release channels (RyR2) determines efficacy of class I antiarrhythmic drugs in catecholaminergic polymorphic ventricular tachycardia. *Circ Arrhythm Electrophysiol.* 2011;4:128–35.
- S7.9.1.2-13.** Adler A, Sadek MM, Chan AY, et al. Patient outcomes from a specialized inherited arrhythmia clinic. *Circ Arrhythm Electrophysiol.* 2016;9:e003440.
- S7.9.1.2-14.** Olde Nordkamp LR, Postema PG, Knops RE, et al. Implantable cardioverter-defibrillator harm in young patients with inherited arrhythmia syndromes: a systematic review and meta-analysis of inappropriate shocks and complications. *Heart Rhythm.* 2016;13:443–54.
- S7.9.1.2-15.** Roses-Noguer F, Jarman JW, Clague JR, et al. Outcomes of defibrillator therapy in catecholaminergic polymorphic ventricular tachycardia. *Heart Rhythm.* 2014;11:58–66.
- S7.9.1.2-16.** Sy RW, Gollob MH, Klein GJ, et al. Arrhythmia characterization and long-term outcomes

in catecholaminergic polymorphic ventricular tachycardia. *Heart Rhythm.* 2011;8:864–71.

**S7.9.1.2-17.** De Ferrari GM, Dusi V, Spazzolini C, et al. Clinical management of catecholaminergic polymorphic ventricular tachycardia: the role of left cardiac sympathetic denervation. *Circulation.* 2015;131:2185–93.

**S7.9.1.2-18.** Wilde AA, Bhuiyan ZA, Crotti L, et al. Left cardiac sympathetic denervation for catecholaminergic polymorphic ventricular tachycardia. *N Engl J Med.* 2008;358:2024–9.

**S7.9.1.2-19.** Zhang C, Kutyifa V, Moss AJ, et al. Long-QT syndrome and therapy for attention deficit/hyperactivity disorder. *J Cardiovasc Electrophysiol.* 2015;26:1039–44.

**S7.9.1.2-20.** Jabbari J, Jabbari R, Nielsen MW, et al. New exome data question the pathogenicity of genetic variants previously associated with catecholaminergic polymorphic ventricular tachycardia. *Circ Cardiovasc Genetics.* 2013;6:481–9.

**S7.9.1.2-21.** Crotti L, Johnson CN, Graf E, et al. Calmodulin mutations associated with recurrent cardiac arrest in infants. *Circulation.* 2013;127:1009–17.

**S7.9.1.2-22.** Makita N, Yagihara N, Crotti L, et al. Novel calmodulin mutations associated with congenital arrhythmia susceptibility. *Circ Cardiovasc Genet.* 2014;7:466–74.

**S7.9.1.2-23.** Marsman RF, Barc J, Beekman L, et al. A mutation in CALM1 encoding calmodulin in familial idiopathic ventricular fibrillation in childhood and adolescence. *J Am Coll Cardiol.* 2014;63:259–66.

**S7.9.1.2-24.** Nyegaard M, Overgaard MT, Sondergaard MT, et al. Mutations in calmodulin cause ventricular tachycardia and sudden cardiac death. *Am J Hum Genet.* 2012;91:703–12.

**S7.9.1.2-25.** Paludan-Muller C, Ahlberg G, Ghose J, et al. Integration of 60,000 exomes and acmg guidelines question the role of catecholaminergic polymorphic ventricular tachycardia-associated variants. *Clin Genet.* 2017;91:63–72.

#### 7.9.1.3. Brugada Syndrome

**S7.9.1.3-1.** Casado-Arroyo R, Berne P, Rao JY, et al. Long-term trends in newly diagnosed Brugada syndrome: implications for risk stratification. *J Am Coll Cardiol.* 2016;68:614–23.

**S7.9.1.3-2.** Gehi AK, Duong TD, Metz LD, et al. Risk stratification of individuals with the Brugada electrocardiogram: a meta-analysis. *J Cardiovasc Electrophysiol.* 2006;17:577–83.

**S7.9.1.3-3.** Hiraoka M, Takagi M, Yokoyama Y, et al. Prognosis and risk stratification of young adults with Brugada syndrome. *J Electrocardiol.* 2013;46:279–83.

**S7.9.1.3-4.** Priori SG, Gasparini M, Napolitano C, et al. Risk stratification in Brugada syndrome: results of the PRELUDE (PRogrammed Electrical stimUlation preDicte valueE) registry. *J Am Coll Cardiol.* 2012;59:37–45.

**S7.9.1.3-5.** Probst V, Veltmann C, Eckardt L, et al. Long-term prognosis of patients diagnosed with Brugada syndrome: results from the FINGER Brugada syndrome registry. *Circulation.* 2010;121:635–43.

**S7.9.1.3-6.** Sroubek J, Probst V, Mazzanti A, et al. Programmed ventricular stimulation for risk stratification in the Brugada syndrome: a pooled analysis. *Circulation.* 2016;133:622–30.

**S7.9.1.3-7.** Belhassen B, Rahkovich M, Michowitz Y, et al. Management of Brugada syndrome: thirty-three-year experience using electrophysiologically guided therapy with class 1A antiarrhythmic drugs. *Circ Arrhythm Electrophysiol.* 2015;8:1393–402.

**S7.9.1.3-8.** Brugada J, Pappone C, Berrezzo A, et al. Brugada syndrome phenotype elimination by epicardial substrate ablation. *Circ Arrhythm Electrophysiol.* 2015;8:1373–81.

**S7.9.1.3-9.** Nademanee K, Veerakul G, Chandana-mattha P, et al. Prevention of ventricular fibrillation episodes in Brugada syndrome by catheter ablation over the anterior right ventricular outflow tract epicardium. *Circulation.* 2011;123:I270–9.

**S7.9.1.3-10.** Sunsaneewitayakul B, Yao Y, Thamaree S, et al. Endocardial mapping and catheter ablation for ventricular fibrillation prevention in Brugada syndrome. *J Cardiovasc Electrophysiol.* 2012;23 suppl 1:S10–6.

**S7.9.1.3-11.** Zhang P, Tung R, Zhang Z, et al. Characterization of the epicardial substrate for catheter ablation of Brugada syndrome. *Heart Rhythm.* 2016;13:2151–8.

**S7.9.1.3-12.** Antzelevitch C, Yan GX, Ackerman MJ, et al. J-wave syndromes expert consensus conference report: emerging concepts and gaps in knowledge. *Europace.* 2017;19:665–94.

**S7.9.1.3-13.** Delise P, Allocca G, Marras E, et al. Risk stratification in individuals with the Brugada type 1 ECG pattern without previous cardiac arrest: usefulness of a combined clinical and electrophysiologic approach. *Eur Heart J.* 2011;32:169–76.

**S7.9.1.3-14.** Somani R, Krahn AD, Healey JS, et al. Procainamide infusion in the evaluation of unexplained cardiac arrest: from the Cardiac Arrest Survivors with Preserved Ejection Fraction Registry (CASPER). *Heart Rhythm.* 2014;11:1047–54.

**S7.9.1.3-15.** Kusumoto FM, Bailey KR, Chaouki AS, et al. Systematic review for the 2017 AHA/ACC/HRS guideline for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2018;72:1653–76.

**S7.9.1.3-16.** Sieira J, Cionte G, Conte G, et al. Asymptomatic Brugada syndrome: clinical characterization and long-term prognosis. *Circ Arrhythm Electrophysiol.* 2015;8:1144–50.

**S7.9.1.3-17.** Sieira J, Conte G, Cionte G, et al. Prognostic value of programmed electrical stimulation in Brugada syndrome: 20 years experience. *Circ Arrhythm Electrophysiol.* 2015;8:777–84.

**S7.9.1.3-18.** Crotti L, Marcou CA, Tester DJ, et al. Spectrum and prevalence of mutations involving BrS1-through BrS12-susceptibility genes in a cohort of unrelated patients referred for Brugada syndrome genetic testing: implications for genetic testing. *J Am Coll Cardiol.* 2012;60:1410–8.

**S7.9.1.3-19.** Probst V, Wilde AA, Barc J, et al. SCN5A mutations and the role of genetic background in the pathophysiology of Brugada syndrome. *Circ Cardiovasc Genet.* 2009;2:552–7.

**S7.9.1.3-20.** Risgaard B, Jabbari R, Refsgaard L, et al. High prevalence of genetic variants previously

associated with Brugada syndrome in new exome data. *Clinical Genetics.* 2013;84:489–95.

**S7.9.1.3-21.** Brugada drugs. Available at: <http://www.brugadadrugs.org>. Accessed October 6, 2016.

**S7.9.1.3-22.** Postema PG, Wolpert C, Amin AS, et al. Drugs and Brugada syndrome patients: review of the literature, recommendations, and an up-to-date website ([www.brugadadrugs.org](http://www.brugadadrugs.org)). *Heart Rhythm.* 2009;6:1335–41.

**S7.9.1.3-23.** Andorin A, Behr ER, Denjoy I, et al. Impact of clinical and genetic findings on the management of young patients with Brugada syndrome. *Heart Rhythm.* 2016;13:1274–82.

**S7.9.1.3-24.** Priori SG, Wilde AA, Horie M, et al. HRS/EHRA/APHRS expert consensus statement on the diagnosis and management of patients with inherited primary arrhythmia syndromes. *Heart Rhythm.* 2013;10:1932–63.

**S7.9.1.3-25.** McNamara DA, Goldberger JJ, Berendsen MA, et al. Implantable defibrillators versus medical therapy for cardiac channelopathies. *Cochrane Database Syst Rev.* 2015;CD011168.

**S7.9.1.3-26.** Nademanee K, Veerakul G, Mower M, et al. Defibrillator versus beta-blockers for unexplained death in Thailand (DEBUT): a randomized clinical trial. *Circulation.* 2003;107:2221–6.

**S7.9.1.3-27.** Rodriguez-Manero M, Sacher F, de AC, et al. Monomorphic ventricular tachycardia in patients with Brugada syndrome: a multicenter retrospective study. *Heart Rhythm.* 2016;13:669–82.

**S7.9.1.3-28.** Brugada R, Brugada J, Antzelevitch C, et al. Sodium channel blockers identify risk for sudden death in patients with ST-segment elevation and right bundle branch block but structurally normal hearts. *Circulation.* 2000;101:510–5.

**S7.9.1.3-29.** Shimono K, Takagi M, Maeda K, et al. Usefulness of multichannel Holter ECG recording in the third intercostal space for detecting type 1 Brugada ECG: comparison with repeated 12-lead ECGs. *J Cardiovasc Electrophysiol.* 2009;20:1026–31.

**S7.9.1.3-30.** Conte G, Sieira J, Cionte G, et al. Implantable cardioverter-defibrillator therapy in Brugada syndrome: a 20-year single-center experience. *J Am Coll Cardiol.* 2015;65:879–88.

**S7.9.1.3-31.** Kapplinger JD, Tester DJ, Alders M, et al. An international compendium of mutations in the SCN5A-encoded cardiac sodium channel in patients referred for Brugada syndrome genetic testing. *Heart Rhythm.* 2010;7:33–46.

**S7.9.1.3-32.** Adler A, Rosso R, Chorin E, et al. Risk stratification in Brugada syndrome: clinical characteristics, electrocardiographic parameters, and auxiliary testing. *Heart Rhythm.* 2016;13:299–310.

**S7.9.1.3-33.** Priori SG, Napolitano C, Gasparini M, et al. Natural history of Brugada syndrome: insights for risk stratification and management. *Circulation.* 2002;105:1342–7.

#### 7.9.1.4. Early Repolarization “J-wave” Syndrome

**S7.9.1.4-1.** Rosso R, Kogan E, Belhassen B, et al. J-point elevation in survivors of primary ventricular fibrillation and matched control subjects: incidence and clinical significance. *J Am Coll Cardiol.* 2008;52:1231–8.

**S7.9.1.4-2.** Adhikarla C, Boga M, Wood AD, et al. Natural history of the electrocardiographic pattern of early repolarization in ambulatory patients. *Am J Cardiol.* 2011;108:1831-5.

**S7.9.1.4-3.** Haissaguerre M, Sacher F, Nogami A, et al. Characteristics of recurrent ventricular fibrillation associated with inferolateral early repolarization role of drug therapy. *J Am Coll Cardiol.* 2009;53:612-9.

**S7.9.1.4-4.** Siebermair J, Sinner MF, Beckmann BM, et al. Early repolarization pattern is the strongest predictor of arrhythmia recurrence in patients with idiopathic ventricular fibrillation: results from a single centre long-term follow-up over 20 years. *Europace.* 2016;18:718-25.

**S7.9.1.4-5.** Sinner MF, Porthan K, Noseworthy PA, et al. A meta-analysis of genome-wide association studies of the electrocardiographic early repolarization pattern. *Heart Rhythm.* 2012;9:1627-34.

**S7.9.1.4-6.** Tikkainen JT, Anttonen O, Junnila MJ, et al. Long-term outcome associated with early repolarization on electrocardiography. *N Engl J Med.* 2009;361:2529-37.

**S7.9.1.4-7.** Tikkainen JT, Wichmann V, Junnila MJ, et al. Association of early repolarization and sudden cardiac death during an acute coronary event. *Circ Arrhythm Electrophysiol.* 2012;5:714-8.

**S7.9.1.4-8.** Junnila MJ, Tikkainen JT, Kentta T, et al. Early repolarization as a predictor of arrhythmic and nonrhythmic cardiac events in middle-aged subjects. *Heart Rhythm.* 2014;11:1701-6.

**S7.9.1.4-9.** Cheng YJ, Lin XX, Ji CC, et al. Role of early repolarization pattern in increasing risk of death. *J Am Heart Assoc.* 2016;5:e003375.

#### 7.9.1.5. Short QT Syndrome

**S7.9.1.5-1.** Dhutia H, Malhotra A, Parpia S, et al. The prevalence and significance of a short QT interval in 18,825 low-risk individuals including athletes. *Br J Sports Med.* 2016;50:124-9.

**S7.9.1.5-2.** Gollob MH, Redpath CJ, Roberts JD. The short QT syndrome: proposed diagnostic criteria. *J Am Coll Cardiol.* 2011;57:802-12.

**S7.9.1.5-3.** Giustetto C, Schimpf R, Mazzanti A, et al. Long-term follow-up of patients with short QT syndrome. *J Am Coll Cardiol.* 2011;58:587-95.

**S7.9.1.5-4.** Mazzanti A, Kanthan A, Monteforte N, et al. Novel insight into the natural history of short QT syndrome. *J Am Coll Cardiol.* 2014;63:1300-8.

**S7.9.1.5-5.** Villafane J, Atallah J, Gollob MH, et al. Long-term follow-up of a pediatric cohort with short QT syndrome. *J Am Coll Cardiol.* 2013;61:1183-91.

**S7.9.1.5-6.** Giustetto C, Di MF, Wolpert C, et al. Short QT syndrome: clinical findings and diagnostic-therapeutic implications. *Eur Heart J.* 2006;27:2440-7.

**S7.9.1.5-7.** Bun SS, Maury P, Giustetto C, et al. Electrical storm in short-QT syndrome successfully treated with isoproterenol. *J Cardiovasc Electrophysiol.* 2012;23:1028-30.

**S7.9.1.5-8.** Guerrier K, Kwiatkowski D, Czosek RJ, et al. Short QT interval prevalence and clinical outcomes in a pediatric population. *Circ Arrhythm Electrophysiol.* 2015;8:1460-4.

**S7.9.1.5-9.** Iribarren C, Round AD, Peng JA, et al. Short QT in a cohort of 1.7 million persons: prevalence,

correlates, and prognosis. *Ann Noninvasive Electrocardiol.* 2014;19:490-500.

**S7.9.1.5-10.** Gollob MH, Blier L, Brugada R, et al. Recommendations for the use of genetic testing in the clinical evaluation of inherited cardiac arrhythmias associated with sudden cardiac death: Canadian Cardiovascular Society/Canadian Heart Rhythm Society joint position paper. *Can J Cardiol.* 2011;27:232-45.

#### 8. VA IN THE STRUCTURALLY NORMAL HEART

**S8-1.** Gill JS, Blaszyk K, Ward DE, et al. Verapamil for the suppression of idiopathic ventricular tachycardia of left bundle branch block-like morphology. *Am Heart J.* 1993;126:1126-33.

**S8-2.** Gill JS, Ward DE, Camm AJ. Comparison of verapamil and diltiazem in the suppression of idiopathic ventricular tachycardia. *Pacing Clin Electrophysiol.* 1992;15:2122-6.

**S8-3.** Kontos MC, Diercks DB, Ho PM, et al. Treatment and outcomes in patients with myocardial infarction treated with acute beta-blocker therapy: results from the American College of Cardiology's NCDR(R). *Am Heart J.* 2011;161:864-70.

**S8-4.** Levine JH, Massumi A, Scheinman MM, et al. Intravenous amiodarone for recurrent sustained hypotensive ventricular tachyarrhythmias. Intravenous Amiodarone Multicenter Trial Group. *J Am Coll Cardiol.* 1996;27:67-75.

**S8-5.** Noda T, Shimizu W, Taguchi A, et al. Malignant entity of idiopathic ventricular fibrillation and polymorphic ventricular tachycardia initiated by premature extrasystoles originating from the right ventricular outflow tract. *J Am Coll Cardiol.* 2005;46:1288-94.

**S8-6.** Viskin S. Idiopathic polymorphic ventricular tachycardia: a "benign disease" with a touch of bad luck? *Korean Circ J.* 2017;47:299-306.

**S8-7.** Viskin S, Rosso R, Rogowski O, et al. The "short-coupled" variant of right ventricular outflow ventricular tachycardia: a not-so-benign form of benign ventricular tachycardia? *J Cardiovasc Electrophysiol.* 2005;16:912-6.

**S8-8.** Kittayaphong R, Bhuripanyo K, Punlee K, et al. Effect of atenolol on symptomatic ventricular arrhythmia without structural heart disease: a randomized placebo-controlled study. *Am Heart J.* 2002;144:e10.

**S8-9.** Ling Z, Liu Z, Su L, et al. Radiofrequency ablation versus antiarrhythmic medication for treatment of ventricular premature beats from the right ventricular outflow tract: prospective randomized study. *Circ Arrhythm Electrophysiol.* 2014;7:237-43.

**S8-10.** Hohnloser SH, Meinertz T, Stubbs P, et al. Efficacy and safety of d-sotalol, a pure class III antiarrhythmic compound, in patients with symptomatic complex ventricular ectopy. Results of a multicenter, randomized, double-blind, placebo-controlled dose-finding study. The d-Sotalol PVC Study Group. *Circulation.* 1995;92:1517-25.

**S8-11.** Badhwar N, Scheinman MM. Idiopathic ventricular tachycardia: diagnosis and management. *Curr Probl Cardiol.* 2007;32:7-43.

**S8-12.** Connolly SJ. Meta-analysis of antiarrhythmic drug trials. *Am J Cardiol.* 1999;84:90R-3R.

#### 8.1. Outflow Tract and Atrioventricular Annular VA

**S8.1-1.** Tada H, Ito S, Naito S, et al. Idiopathic ventricular arrhythmia arising from the mitral annulus: a distinct subgroup of idiopathic ventricular arrhythmias. *J Am Coll Cardiol.* 2005;45:877-86.

**S8.1-2.** Yamada T, Litovsky SH, Kay GN. The left ventricular ostium: an anatomic concept relevant to idiopathic ventricular arrhythmias. *Circ Arrhythm Electrophysiol.* 2008;1:396-404.

**S8.1-3.** Yamada T, Maddox WR, McElderry HT, et al. Radiofrequency catheter ablation of idiopathic ventricular arrhythmias originating from intramural foci in the left ventricular outflow tract: efficacy of sequential versus simultaneous unipolar catheter ablation. *Circ Arrhythm Electrophysiol.* 2015;8:344-52.

**S8.1-4.** Ling Z, Liu Z, Su L, et al. Radiofrequency ablation versus antiarrhythmic medication for treatment of ventricular premature beats from the right ventricular outflow tract: prospective randomized study. *Circ Arrhythm Electrophysiol.* 2014;7:237-43.

**S8.1-5.** Carballera Pol L, Deyell MW, Frankel DS, et al. Ventricular premature depolarization QRS duration as a new marker of risk for the development of ventricular premature depolarization-induced cardiomyopathy. *Heart Rhythm.* 2014;11:299-306.

**S8.1-6.** Kamioka M, Mathew S, Lin T, et al. Electrophysiological and electrocardiographic predictors of ventricular arrhythmias originating from the left ventricular outflow tract within and below the coronary sinus cusps. *Clin Res Cardiol.* 2015;104:544-54.

**S8.1-7.** Konstantinidou M, Koeketurk B, Wissner E, et al. Catheter ablation of right ventricular outflow tract tachycardia: a simplified remote-controlled approach. *Europace.* 2011;13:696-700.

**S8.1-8.** Latchamsetty R, Yokokawa M, Morady F, et al. Multicenter outcomes for catheter ablation of idiopathic premature ventricular complexes. *JACC Clin Electrophysiol.* 2015;1:116-23.

**S8.1-9.** Liao Z, Zhan X, Wu S, et al. Idiopathic ventricular arrhythmias originating from the pulmonary sinus cusp: prevalence, electrocardiographic/electrophysiological characteristics, and catheter ablation. *J Am Coll Cardiol.* 2015;66:2633-44.

**S8.1-10.** Morady F, Kadish AH, DiCarlo L, et al. Long-term results of catheter ablation of idiopathic right ventricular tachycardia. *Circulation.* 1990;82:2093-9.

**S8.1-11.** Ouyang F, Fotuhi P, Ho SY, et al. Repetitive monomorphic ventricular tachycardia originating from the aortic sinus cusp: electrocardiographic characterization for guiding catheter ablation. *J Am Coll Cardiol.* 2002;39:500-8.

**S8.1-12.** Pedersen CT, Kay GN, Kalman J, et al. EHRA/HRS/APHRS expert consensus on ventricular arrhythmias. *Heart Rhythm.* 2014;11:e166-96.

**S8.1-13.** Tada H, Tadokoro K, Miyaji K, et al. Idiopathic ventricular arrhythmias arising from the pulmonary artery: prevalence, characteristics, and topography of the arrhythmia origin. *Heart Rhythm.* 2008;5:419-26.

**S8.1-14.** Yamada T, McElderry HT, Doppalapudi H, et al. Idiopathic ventricular arrhythmias originating from the aortic root prevalence, electrocardiographic and electrophysiologic characteristics, and results of radiofrequency catheter ablation. *J Am Coll Cardiol.* 2008;52:139-47.

- S8.1-15.** Yamada T, McElderry HT, Doppalapudi H, et al. Idiopathic ventricular arrhythmias originating from the left ventricular summit: anatomic concepts relevant to ablation. *Circ Arrhythm Electrophysiol*. 2010;3:616–23.
- S8.1-16.** Yamada T, McElderry HT, Okada T, et al. Idiopathic left ventricular arrhythmias originating adjacent to the left aortic sinus of valsalva: electrophysiological rationale for the surface electrocardiogram. *J Cardiovasc Electrophysiol*. 2010;21:170–6.
- S8.1-17.** Mountantonakis SE, Frankel DS, Tschabrunn CM, et al. Ventricular arrhythmias from the coronary venous system: prevalence, mapping, and ablation. *Heart Rhythm*. 2015;12:1145–53.
- S8.1-18.** Nagashima K, Choi EK, Lin KY, et al. Ventricular arrhythmias near the distal great cardiac vein: challenging arrhythmia for ablation. *Circ Arrhythm Electrophysiol*. 2014;7:906–12.
- S8.1-19.** Hai JJ, Chahal AA, Friedman PA, et al. Electrophysiologic characteristics of ventricular arrhythmias arising from the aortic mitral continuity–potential role of the conduction system. *J Cardiovasc Electrophysiol*. 2015;26:158–63.
- S8.1-20.** Tada H, Tadokoro K, Ito S, et al. Idiopathic ventricular arrhythmias originating from the tricuspid annulus: prevalence, electrocardiographic characteristics, and results of radiofrequency catheter ablation. *Heart Rhythm*. 2007;4:7–16.
- 8.2. Papillary Muscle VA**
- S8.2-1.** Ban JE, Lee HS, Lee DI, et al. Electrophysiological characteristics related to outcome after catheter ablation of idiopathic ventricular arrhythmias originating from the papillary muscle in the left ventricle. *Korean Circ J*. 2013;43:811–8.
- S8.2-2.** Crawford T, Mueller G, Good E, et al. Ventricular arrhythmias originating from papillary muscles in the right ventricle. *Heart Rhythm*. 2010;7:725–30.
- S8.2-3.** Doppalapudi H, Yamada T, McElderry HT, et al. Ventricular tachycardia originating from the posterior papillary muscle in the left ventricle: a distinct clinical syndrome. *Circ Arrhythm Electrophysiol*. 2008;1:23–9.
- S8.2-4.** Yamada T, Doppalapudi H, McElderry HT, et al. Electrocardiographic and electrophysiological characteristics in idiopathic ventricular arrhythmias originating from the papillary muscles in the left ventricle: relevance for catheter ablation. *Circ Arrhythm Electrophysiol*. 2010;3:324–31.
- S8.2-5.** Yokokawa M, Good E, Desjardins B, et al. Predictors of successful catheter ablation of ventricular arrhythmias arising from the papillary muscles. *Heart Rhythm*. 2010;7:1654–9.
- 8.3. Interfascicular Reentrant VT (Belhassen Tachycardia)**
- S8.3-1.** Lin D, Hsia HH, Gerstenfeld EP, et al. Idiopathic fascicular left ventricular tachycardia: linear ablation lesion strategy for noninducible or nonsustained tachycardia. *Heart Rhythm*. 2005;2:934–9.
- S8.3-2.** Liu Y, Fang Z, Yang B, et al. Catheter ablation of fascicular ventricular tachycardia: long-term clinical outcomes and mechanisms of recurrence. *Circ Arrhythm Electrophysiol*. 2015;8:1443–51.
- S8.3-3.** Nogami A, Naito S, Tada H, et al. Demonstration of diastolic and presystolic Purkinje potentials as critical potentials in a macroreentry circuit of verapamil-sensitive idiopathic left ventricular tachycardia. *J Am Coll Cardiol*. 2000;36:811–23.
- S8.3-4.** Belhassen B, Rotmansch HH, Laniado S. Response of recurrent sustained ventricular tachycardia to verapamil. *Br Heart J*. 1981;46:679–82.
- S8.3-5.** German LD, Packer DL, Bardy GH, et al. Ventricular tachycardia induced by atrial stimulation in patients without symptomatic cardiac disease. *Am J Cardiol*. 1983;52:1202–7.
- S8.3-6.** Tsuchiya T, Okumura K, Honda T, et al. Effects of verapamil and lidocaine on two components of the re-entry circuit of verapamil-sensitive idiopathic left ventricular tachycardia. *J Am Coll Cardiol*. 2001;37:1415–21.
- S8.3-7.** Anderson JH, Tester DJ, Will ML, et al. Whole-exome molecular autopsy after exertion-related sudden unexplained death in the young. *Circ Cardiovasc Genet*. 2016;9:259–65.
- S8.3-8.** Ohe T, Shimomura K, Aihara N, et al. Idiopathic sustained left ventricular tachycardia: clinical and electrophysiologic characteristics. *Circulation*. 1988;77:560–8.
- S8.3-9.** Snyder C, Bishara J, Darling R, et al. Verapamil-sensitive ventricular tachycardia in an infant. *Congenit Heart Dis*. 2006;1:124–6.
- S8.3-10.** Wang JD, Fu YC, Jan SL, et al. Verapamil sensitive idiopathic ventricular tachycardia in an infant. *Jpn Heart J*. 2003;44:667–71.
- 8.4. Idiopathic Polymorphic VT/VF**
- S8.4-1.** Anderson JH, Tester DJ, Will ML, et al. Whole-exome molecular autopsy after exertion-related sudden unexplained death in the young. *Circ Cardiovasc Genet*. 2016;9:259–65.
- S8.4-2.** Dalal A, Czosek RJ, Kovach J, et al. Clinical presentation of pediatric patients at risk for sudden cardiac arrest. *J Pediatr*. 2016.
- S8.4-3.** Kumar S, Peters S, Thompson T, et al. Familial cardiological and targeted genetic evaluation: low yield in sudden unexplained death and high yield in unexplained cardiac arrest syndromes. *Heart Rhythm*. 2013;10:1653–60.
- S8.4-4.** Linzer M, Pritchett EL, Pontinen M, et al. Incremental diagnostic yield of loop electrocardiographic recorders in unexplained syncope. *Am J Cardiol*. 1990;66:214–9.
- S8.4-5.** Solomon SD, Zelenkofske S, McMurray JJ, et al. Sudden death in patients with myocardial infarction and left ventricular dysfunction, heart failure, or both. *N Engl J Med*. 2005;352:2581–8.
- S8.4-6.** Tester DJ, Medeiros-Domingo A, Will ML, et al. Unexplained drownings and the cardiac channelopathies: a molecular autopsy series. *Mayo Clin Proc*. 2011;86:941–7.
- S8.4-7.** Tzimas I, Zingraff JC, Bajanowski T, et al. The role of known variants of KCNQ1, KCNH2, KCNE1, SCNSA, and NOS1AP in water-related deaths. *Int J Legal Med*. 2016;1575–9.
- S8.4-8.** Wang D, Shah KR, Um SY, et al. Cardiac channelopathy testing in 274 ethnically diverse sudden unexplained deaths. *Forensic Sci Int*. 2014;237:90–9.
- S8.4-9.** Conte G, Caputo ML, Regoli F, et al. True idiopathic ventricular fibrillation in out-of-hospital cardiac arrest survivors in the Swiss Canton Ticino: prevalence, clinical features, and long-term follow-up. *Europace*. 2017;19:259–66.
- S8.4-10.** Frommeyer G, Decherling DG, Kochhauser S, et al. Long-time “real-life” performance of the subcutaneous ICD in patients with electrical heart disease or idiopathic ventricular fibrillation. *J Interv Card Electrophysiol*. 2016;47:185–8.
- S8.4-11.** Haissaguerre M, Shoda M, Jais P, et al. Mapping and ablation of idiopathic ventricular fibrillation. *Circulation*. 2002;106:962–7.
- S8.4-12.** Knecht S, Sacher F, Wright M, et al. Long-term follow-up of idiopathic ventricular fibrillation ablation: a multicenter study. *J Am Coll Cardiol*. 2009;54:522–8.
- S8.4-13.** Lehnhardt A, Glaser E, Burguera M, et al. Short-coupled variant of torsade de pointes. A new electrocardiographic entity in the spectrum of idiopathic ventricular tachyarrhythmias. *Circulation*. 1994;89:206–15.
- S8.4-14.** Haissaguerre M, Shah DC, Jais P, et al. Role of Purkinje conducting system in triggering of idiopathic ventricular fibrillation. *Lancet*. 2002;359:677–8.
- S8.4-15.** Albertella L, Crawford J, Skinner JR. Presentation and outcome of water-related events in children with long QT syndrome. *Arch Dis Child*. 2011;96:704–7.
- S8.4-16.** Miyake CY, Motonaga KS, Fischer-Colbrie ME, et al. Risk of cardiac disease and observations on lack of potential predictors by clinical history among children presenting for cardiac evaluation of mid-exertional syncope. *Cardiol Young*. 2016;26:894–900.
- S8.4-17.** Gula LJ, Klein GJ, Hellkamp AS, et al. Ejection fraction assessment and survival: an analysis of the Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT). *Am Heart J*. 2008;156:1196–200.
- S8.4-18.** Winkel BG, Yuan L, Olesen MS, et al. The role of the sodium current complex in a nonreferred nationwide cohort of sudden infant death syndrome. *Heart Rhythm*. 2015;12:1241–9.
- S8.4-19.** Noda T, Shimizu W, Taguchi A, et al. Malignant entity of idiopathic ventricular fibrillation and polymorphic ventricular tachycardia initiated by premature extrasystoles originating from the right ventricular outflow tract. *J Am Coll Cardiol*. 2005;46:1288–94.
- S8.4-20.** Sadek MM, Benhayon D, Sureddi R, et al. Idiopathic ventricular arrhythmias originating from the moderator band: electrocardiographic characteristics and treatment by catheter ablation. *Heart Rhythm*. 2015;12:67–75.
- S8.4-21.** Van HH, Zado ES, Haqqani H, et al. Catheter ablation of ventricular fibrillation: importance of left ventricular outflow tract and papillary muscle triggers. *Heart Rhythm*. 2014;11:566–73.
- S8.4-22.** Viskin S, Belhassen B. Idiopathic ventricular fibrillation. *Am Heart J*. 1990;120:661–71.
- 9. PVC-INDUCED CARDIOMYOPATHY**
- S9-1.** Haissaguerre M, Shah DC, Jais P, et al. Role of Purkinje conducting system in triggering of idiopathic ventricular fibrillation. *Lancet*. 2002;359:677–8.
- S9-2.** Haissaguerre M, Shoda M, Jais P, et al. Mapping and ablation of idiopathic ventricular fibrillation. *Circulation*. 2002;106:962–7.
- S9-3.** Lee GK, Klarich KW, Grogan M, et al. Premature ventricular contraction-induced cardiomyopathy: a

treatable condition. *Circ Arrhythm Electrophysiol.* 2012;5:229–36.

**S9-4.** Singh SN, Fletcher RD, Fisher SG, et al. Amiodarone in patients with congestive heart failure and asymptomatic ventricular arrhythmia. Survival Trial of Antiarrhythmic Therapy in Congestive Heart Failure. *N Engl J Med.* 1995;333:77–82.

**S9-5.** Baman TS, Lange DC, Ilg KJ, et al. Relationship between burden of premature ventricular complexes and left ventricular function. *Heart Rhythm.* 2010;7:865–9.

**S9-6.** Ban JE, Park HC, Park JS, et al. Electrocardiographic and electrophysiological characteristics of premature ventricular complexes associated with left ventricular dysfunction in patients without structural heart disease. *Europace.* 2013;15:735–41.

**S9-7.** Bogun F, Crawford T, Reich S, et al. Radiofrequency ablation of frequent, idiopathic premature ventricular complexes: comparison with a control group without intervention. *Heart Rhythm.* 2007;4:863–7.

**S9-8.** Carballera Pol L, Deyell MW, Frankel DS, et al. Ventricular premature depolarization QRS duration as a new marker of risk for the development of ventricular premature depolarization-induced cardiomyopathy. *Heart Rhythm.* 2014;11:299–306.

**S9-9.** Del Carpio Munoz F, Syed FF, Noheria A, et al. Characteristics of premature ventricular complexes as correlates of reduced left ventricular systolic function: study of the burden, duration, coupling interval, morphology and site of origin of PVCs. *J Cardiovasc Electrophysiol.* 2011;22:791–8.

**S9-10.** Deyell MW, Park KM, Han Y, et al. Predictors of recovery of left ventricular dysfunction after ablation of frequent ventricular premature depolarizations. *Heart Rhythm.* 2012;9:1465–72.

**S9-11.** Hamon D, Blaye-Felice MS, Bradfield JS, et al. A new combined parameter to predict premature ventricular complexes induced cardiomyopathy: impact and recognition of epicardial origin. *J Cardiovasc Electrophysiol.* 2016;27:709–17.

**S9-12.** Hasdemir C, Ulucan C, Yavuzgil O, et al. Tachycardia-induced cardiomyopathy in patients with idiopathic ventricular arrhythmias: the incidence, clinical and electrophysiologic characteristics, and the predictors. *J Cardiovasc Electrophysiol.* 2011;22:663–8.

**S9-13.** Kanei Y, Friedman M, Ogawa N, et al. Frequent premature ventricular complexes originating from the right ventricular outflow tract are associated with left ventricular dysfunction. *Ann Noninvasive Electrocardiol.* 2008;13:81–5.

**S9-14.** Kawamura M, Badhwar N, Vedantham V, et al. Coupling interval dispersion and body mass index are independent predictors of idiopathic premature ventricular complex-induced cardiomyopathy. *J Cardiovasc Electrophysiol.* 2014;25: 756–2.

**S9-15.** Niwano S, Wakisaka Y, Niwano H, et al. Prognostic significance of frequent premature ventricular contractions originating from the ventricular outflow tract in patients with normal left ventricular function. *Heart.* 2009;95:1230–7.

**S9-16.** Olgun H, Yokokawa M, Baman T, et al. The role of interpolation in PVC-induced cardiomyopathy. *Heart Rhythm.* 2011;8:1046–9.

**S9-17.** Yokokawa M, Good E, Crawford T, et al. Recovery from left ventricular dysfunction after ablation of frequent premature ventricular complexes. *Heart Rhythm.* 2013;10:172–5.

**S9-18.** Zhong L, Lee YH, Huang XM, et al. Relative efficacy of catheter ablation vs antiarrhythmic drugs in treating premature ventricular contractions: a single-center retrospective study. *Heart Rhythm.* 2014;11:187–93.

**S9-19.** Deyell MW, Park KM, Han Y, et al. Predictors of recovery of left ventricular dysfunction after ablation of frequent ventricular premature depolarizations. *Heart Rhythm.* 2012;9:1465–72.

**S9-20.** Kubac G, Klinke WP, Grace M. Randomized double blind trial comparing sotalol and propranolol in chronic ventricular arrhythmia. *Can J Cardiol.* 1988;4:355–9.

## 10. VA AND SCD RELATED TO SPECIFIC POPULATIONS

### 10.1. Athletes

**S10.1-1.** Zipes DP, Link MS, Ackerman MJ, et al. Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities: Task Force 9: arrhythmias and conduction defects: a scientific statement from the American Heart Association and American College of Cardiology. *Circulation.* 2015;132:e315–25.

**S10.1-2.** Biffi A, Pelliccia A, Verdile L, et al. Long-term clinical significance of frequent and complex ventricular tachyarrhythmias in trained athletes. *J Am Coll Cardiol.* 2002;40:446–52.

**S10.1-3.** Verdile L, Maron BJ, Pelliccia A, et al. Clinical significance of exercise-induced ventricular tachyarrhythmias in trained athletes without cardiovascular abnormalities. *Heart Rhythm.* 2015;12:78–85.

**S10.1-4.** Harmon KG, Asif IM, Maleszewski JJ, et al. Incidence, cause, and comparative frequency of sudden cardiac death in national collegiate athletic association athletes: a decade in review. *Circulation.* 2015;132:10–9.

**S10.1-5.** Maron BJ, Haas TS, Ahluwalia A, et al. Incidence of cardiovascular sudden deaths in Minnesota high school athletes. *Heart Rhythm.* 2013;10:374–7.

**S10.1-6.** Harmon KG, Drezner JA, Wilson MG, et al. Incidence of sudden cardiac death in athletes: a state-of-the-art review. *Br J Sports Med.* 2014;48:1185–92.

**S10.1-7.** Marijon E, Tafflet M, Celermajer DS, et al. Sports-related sudden death in the general population. *Circulation.* 2011;124:672–81.

**S10.1-8.** Meyer L, Stubbs B, Fahrenbruch C, et al. Incidence, causes, and survival trends from cardiovascular-related sudden cardiac arrest in children and young adults 0 to 35 years of age: a 30-year review. *Circulation.* 2012;126:1363–72.

**S10.1-9.** Marijon E, Bouguoin W, Perier MC, et al. Incidence of sports-related sudden death in France by specific sports and sex. *JAMA.* 2013;310:642–3.

**S10.1-10.** Anderson JH, Tester DJ, Will ML, et al. Whole-exome molecular autopsy after exertion-related sudden unexplained death in the young. *Circ Cardiovasc Genet.* 2016;9:259–65.

**S10.1-11.** Bagnall RD, Weintraub RG, Ingles J, et al. A prospective study of sudden cardiac death among children and young adults. *N Engl J Med.* 2016;374:2441–52.

**S10.1-12.** Semsarian C, Ingles J, Wilde AA. Sudden cardiac death in the young: the molecular autopsy and a practical approach to surviving relatives. *Eur Heart J.* 2015;36:1290–6.

**S10.1-13.** Maron BJ, Haas TS, Ahluwalia A, et al. Demographics and epidemiology of sudden deaths in young competitive athletes: from the United States national registry. *Am J Med.* 2016;129:1170–7.

**S10.1-14.** Yankelson L, Sadeh B, Gershovitz L, et al. Life-threatening events during endurance sports: is heat stroke more prevalent than arrhythmic death? *J Am Coll Cardiol.* 2014;64:463–9.

**S10.1-15.** Maron BJ, Haas TS, Murphy CJ, et al. Incidence and causes of sudden death in U.S. college athletes. *J Am Coll Cardiol.* 2014;63:1636–43.

**S10.1-16.** Link MS, Myerburg RJ, Estes NA 3rd. Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities: Task Force 12: emergency action plans, resuscitation, cardiopulmonary resuscitation, and automated external defibrillators: a scientific statement from the American Heart Association and American College of Cardiology. *Circulation.* 2015;132:e334–8.

**S10.1-17.** Brothers JA, Frommelt MA, Jaquiss RDB, et al. Expert consensus guideline: anomalous aortic origin of a coronary artery – American Association for Thoracic Surgery Clinical Practice Guidelines. *J Thorac Cardiovasc Surg.* 2017;153:1440–57.

**S10.1-18.** Van Hare GF, Ackerman MJ, Evangelista JA, et al. Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities: Task Force 4: congenital heart disease: a scientific statement from the American Heart Association and American College of Cardiology. *Circulation.* 2015;132:e281–91.

**S10.1-19.** Ackerman MJ, Zipes DP, Kovacs RJ, et al. Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities: Task Force 10: the cardiac channelopathies: a scientific statement from the American Heart Association and American College of Cardiology. *Circulation.* 2015;132:e326–9.

**S10.1-20.** Maron BJ, Udelson JE, Bonow RO, et al. Eligibility and disqualification recommendations for competitive athletes with cardiovascular abnormalities: Task Force 3: Hypertrophic cardiomyopathy, arrhythmogenic right ventricular cardiomyopathy and other cardiomyopathies, and myocarditis: a scientific statement from the American Heart Association and American College of Cardiology. *Circulation.* 2015;132:e273–80.

### 10.2. Pregnancy

**S10.2-1.** Seth R, Moss AJ, McNitt S, et al. Long QT syndrome and pregnancy. *J Am Coll Cardiol.* 2007;49:1092–8.

**S10.2-2.** Jejeebhoy FM, Zelop CM, Lipman S, et al. Cardiac arrest in pregnancy: a scientific statement from the American Heart Association. *Circulation.* 2015;132:1747–73.

**S10.2-3.** Vanden Hoek TL, Morrison LJ, Shuster M, et al. Part 12: cardiac arrest in special situations: 2010

American Heart Association guidelines for cardiopulmonary resuscitation and emergency cardiovascular care. *Circulation.* 2010;122 suppl 3:S829–61.

**S10.2-4.** Natale A, Davidson T, Geiger MJ, et al. Implantable cardioverter-defibrillators and pregnancy: a safe combination? *Circulation.* 1997;96:2808–12.

**S10.2-5.** Colletti PM, Lee KH, Elkayam U. Cardiac imaging of the pregnant patient. *Am J Roentgenol.* 2013;200:515–21.

**S10.2-6.** Khositseth A, Tester DJ, Will ML, et al. Identification of a common genetic substrate underlying postpartum cardiac events in congenital long QT syndrome. *Heart Rhythm.* 2004;1:60–4.

**S10.2-7.** Rashba EJ, Zareba W, Moss AJ, et al. Influence of pregnancy on the risk for cardiac events in patients with hereditary long QT syndrome. LQTS Investigators. *Circulation.* 1998;97:451–6.

**S10.2-8.** Heradien MJ, Goosen A, Crotti L, et al. Does pregnancy increase cardiac risk for LQT1 patients with the KCNQ1-A34IV mutation? *J Am Coll Cardiol.* 2006;48:1410–5.

**S10.2-9.** Davis RL, Eastman D, McPhillips H, et al. Risks of congenital malformations and perinatal events among infants exposed to calcium channel and beta-blockers during pregnancy. *Pharmacoepidemiol Drug Saf.* 2011;20:138–45.

**S10.2-10.** Ruys TP, Maggioni A, Johnson MR, et al. Cardiac medication during pregnancy, data from the ROPAC. *Int J Cardiol.* 2014;177:124–8.

**S10.2-11.** Beinder E, Grancay T, Menendez T, et al. Fetal sinus bradycardia and the long QT syndrome. *Am J Obstet Gynecol.* 2001;185:743–7.

**S10.2-12.** Hofbeck M, Ulmer H, Beinder E, et al. Prenatal findings in patients with prolonged QT interval in the neonatal period. *Heart.* 1997;77:198–204.

**S10.2-13.** Cuneo BF, Etheridge SP, Horigome H, et al. Arrhythmia phenotype during fetal life suggests long-QT syndrome genotype: risk stratification of perinatal long-QT syndrome. *Circ Arrhythm Electrophysiol.* 2013;6:946–51.

**S10.2-14.** Cuneo BF, Strasburger JF, Wakai RT. The natural history of fetal long QT syndrome. *J Electrocardiol.* 2016;49:807–13.

**S10.2-15.** Damilakis J, Theοcharopoulos N, Perisinakis K, et al. Conceptus radiation dose and risk from cardiac catheter ablation procedures. *Circulation.* 2001;104:893–7.

**S10.2-16.** Duncker D, Haghikia A, Konig T, et al. Risk for ventricular fibrillation in peripartum cardiomyopathy with severely reduced left ventricular function—value of the wearable cardioverter/defibrillator. *Eur J Heart Fail.* 2014;16:1331–6.

### 10.3. Older Patients With Comorbidities

**S10.3-1.** Kusumoto FM, Bailey KR, Chaouki AS, et al. Systematic review for the 2017 AHA/ACC/HRS guideline for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2018;72:1653–76.

**S10.3-2.** American Geriatrics Society Expert Panel on the Care of Older Adults with M. Guiding principles for the care of older adults with multimorbidity: an

approach for clinicians. *J Am Geriatr Soc.* 2012;60:E1–25.

### 10.4. Chronic Kidney Disease

**S10.4-1.** Bardy GH, Lee KL, Mark DB, et al. Amiodarone or an implantable cardioverter-defibrillator for congestive heart failure. *N Engl J Med.* 2005;352:225–37.

**S10.4-2.** Buxton AE, Lee KL, Fisher JD, et al. A randomized study of the prevention of sudden death in patients with coronary artery disease. Multicenter Unsustained Tachycardia Trial Investigators. *N Engl J Med.* 1999;341:1882–90.

**S10.4-3.** Moss AJ, Hall WJ, Cannom DS, et al. Improved survival with an implanted defibrillator in patients with coronary disease at high risk for ventricular arrhythmia. Multicenter Automatic Defibrillator Implantation Trial Investigators. *N Engl J Med.* 1996;335:1933–40.

**S10.4-4.** Moss AJ, Zareba W, Hall WJ, et al. Prophylactic implantation of a defibrillator in patients with myocardial infarction and reduced ejection fraction. *N Engl J Med.* 2002;346:877–83.

**S10.4-5.** Pun PH, Al-Khatib SM, Han JY, et al. Implantable cardioverter-defibrillators for primary prevention of sudden cardiac death in CKD: a meta-analysis of patient-level data from 3 randomized trials. *Am J Kidney Dis.* 2014;64:32–9.

**S10.4-6.** Charytan DM, Patrick AR, Liu J, et al. Trends in the use and outcomes of implantable cardioverter-defibrillators in patients undergoing dialysis in the United States. *Am J Kidney Dis.* 2011;58:409–17.

**S10.4-7.** Wong MC, Kalman JM, Pedagogos E, et al. Bradycardia and asystole is the predominant mechanism of sudden cardiac death in patients with chronic kidney disease. *J Am Coll Cardiol.* 2015;65:1263–5.

**S10.4-8.** Kusumoto FM, Bailey KR, Chaouki AS, et al. Systematic review for the 2017 AHA/ACC/HRS guideline for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines and the Heart Rhythm Society. *J Am Coll Cardiol.* 2018;72:1653–76.

**S10.4-9.** Pun PH, Hellkamp AS, Sanders GD, et al. Primary prevention implantable cardioverter defibrillators in end-stage kidney disease patients on dialysis: a matched cohort study. *Nephrol Dial Transplant.* 2015;30:829–35.

### 10.5. Valvular Heart Disease

**S10.6-1.** Nishimura RA, Otto CM, Bonow RO, et al. 2014 AHA/ACC guideline for the management of patients with valvular heart disease: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;63:e57–185.

**S10.6-2.** Genereux P, Stone GW, O'Gara PT, et al. Natural history, diagnostic approaches, and therapeutic strategies for patients with asymptomatic severe aortic stenosis. *J Am Coll Cardiol.* 2016;67:2263–88.

**S10.6-3.** Avierinos JF, Gersh BJ, Melton LJ 3rd, et al. Natural history of asymptomatic mitral valve prolapse in the community. *Circulation.* 2002;106:1355–61.

**S10.6-4.** Grigioni F, Enriquez-Sarano M, Ling LH, et al. Sudden death in mitral regurgitation due to flail leaflet. *J Am Coll Cardiol.* 1999;34:2078–85.

**S10.6-5.** Nordhues BD, Sontis KC, Scott CG, et al. Bileaflet mitral valve prolapse and risk of ventricular dysrhythmias and death. *J Cardiovasc Electrophysiol.* 2016;27:463–8.

**S10.6-6.** Basso C, Perazzolo MM, Rizzo S, et al. Arrhythmic mitral valve prolapse and sudden cardiac death. *Circulation.* 2015;132:556–66.

**S10.6-7.** Siram CS, Syed FF, Ferguson ME, et al. Malignant bileaflet mitral valve prolapse syndrome in patients with otherwise idiopathic out-of-hospital cardiac arrest. *J Am Coll Cardiol.* 2013;62:222–30.

**S10.6-8.** Albert CM, Chae CU, Grodstein F, et al. Prospective study of sudden cardiac death among women in the United States. *Circulation.* 2003;107:2096–101.

**S10.6-9.** Kannel WB, Wilson PW, D'Agostino RB, et al. Sudden coronary death in women. *Am Heart J.* 1998;136:205–12.

**S10.6-10.** Bogle BM, Ning H, Mehrotra S, et al. Lifetime risk for sudden cardiac death in the community. *J Am Heart Assoc.* 2016;5:e002398.

**S10.6-11.** Kim C, Fahrenbruch CE, Cobb LA, et al. Out-of-hospital cardiac arrest in men and women. *Circulation.* 2001;104:2699–703.

**S10.6-12.** Teodorescu C, Reinier K, Uy-Evanado A, et al. Survival advantage from ventricular fibrillation and pulseless electrical activity in women compared to men: the Oregon Sudden Unexpected Death Study. *J Interv Card Electrophysiol.* 2012;34:219–25.

**S10.6-13.** Wissenberg M, Hansen CM, Folke F, et al. Survival after out-of-hospital cardiac arrest in relation to sex: a nationwide registry-based study. *Resuscitation.* 2014;85:1212–8.

**S10.6-14.** Bray JE, Stub D, Bernard S, et al. Exploring gender differences and the "oestrogen effect" in an Australian out-of-hospital cardiac arrest population. *Resuscitation.* 2013;84:957–63.

**S10.6-15.** Ng YY, Wah W, Liu N, et al. Associations between gender and cardiac arrest outcomes in Pan-Asian out-of-hospital cardiac arrest patients. *Resuscitation.* 2016;102:116–21.

**S10.6-16.** Hagihara A, Onozuka D, Hasegawa M, et al. Resuscitation outcomes of reproductive-age females who experienced out-of-hospital cardiac arrest. *Eur Heart J Acute Cardiovasc Care.* 2017;6:121–9.

**S10.6-17.** Safdar B, Stolz U, Stiell IG, et al. Differential survival for men and women from out-of-hospital cardiac arrest varies by age: results from the OPALS study. *Acad Emerg Med.* 2014;21:1503–11.

**S10.6-18.** Albert CM, McGovern BA, Newell JB, et al. Sex differences in cardiac arrest survivors. *Circulation.* 1996;93:1170–6.

**S10.6-19.** Chugh SS, Uy-Evanado A, Teodorescu C, et al. Women have a lower prevalence of structural heart disease as a precursor to sudden cardiac arrest: The Ore-SUDS (Oregon Sudden Unexpected Death Study). *J Am Coll Cardiol.* 2009;54:2006–11.

**S10.6-20.** Kim MJ, Shin SD, McClellan WM, et al. Neurological prognostication by gender in out-of-hospital cardiac arrest patients receiving hypothermia treatment. *Resuscitation.* 2014;85:1732–8.

**S10.6-21.** Marijon E, Bouguin W, Perier MC, et al. Incidence of sports-related sudden death in France by specific sports and sex. *JAMA.* 2013;310:642–3.

**S10.6-22.** Makkar RR, Fromm BS, Steinman RT, et al. Female gender as a risk factor for torsades de pointes

associated with cardiovascular drugs. *JAMA*. 1993;270:2590-7.

**S10.6-23.** Higgins AY, Waks JW, Josephson ME. Influence of gender on the tolerability, safety, and efficacy of quinidine used for treatment of supraventricular and ventricular arrhythmias. *Am J Cardiol*. 2015;116:1845-51.

**S10.6-24.** Tanaka Y, Tada H, Ito S, et al. Gender and age differences in candidates for radiofrequency catheter ablation of idiopathic ventricular arrhythmias. *Circ J*. 2011;75:1585-91.

#### 10.7. Medication-Induced Arrhythmias

**S10.7-1.** Antman EM, Wenger TL, Butler VP Jr., et al. Treatment of 150 cases of life-threatening digitalis intoxication with digoxin-specific Fab antibody fragments. Final report of a multicenter study. *Circulation*. 1990;81:1744-52.

**S10.7-2.** Chan BS, Buckley NA. Digoxin-specific antibody fragments in the treatment of digoxin toxicity. *Clin Toxicol*. 2014;52:824-36.

**S10.7-3.** Keren A, Tzivoni D, Gavish D, et al. Etiology, warning signs and therapy of torsade de pointes. A study of 10 patients. *Circulation*. 1981;64:1167-74.

**S10.7-4.** Tzivoni D, Banai S, Schuger C, et al. Treatment of torsade de pointes with magnesium sulfate. *Circulation*. 1988;77:392-7.

**S10.7-5.** Kannankeril P, Roden DM, Darbar D. Drug-induced long QT syndrome. *Pharmacol Rev*. 2010;62:760-81.

**S10.7-6.** Choy AM, Lang CC, Chomsky DM, et al. Normalization of acquired QT prolongation in humans by intravenous potassium. *Circulation*. 1997;96:2149-54.

**S10.7-7.** Yang T, Roden DM. Extracellular potassium modulation of drug block of IKr. Implications for torsade de pointes and reverse use-dependence. *Circulation*. 1996;93:407-11.

**S10.7-8.** Hellestrand KJ, Burnett PJ, Milne JR, et al. Effect of the antiarrhythmic agent flecainide acetate on acute and chronic pacing thresholds. *Pacing Clin Electrophysiol*. 1983;6:892-9.

**S10.7-9.** Echt DS, Black JN, Barbey JT, et al. Evaluation of antiarrhythmic drugs on defibrillation energy requirements in dogs. Sodium channel block and action potential prolongation. *Circulation*. 1989;79:1106-17.

**S10.7-10.** Schwartz PJ, Woosley RL. Predicting the unpredictable: drug-induced QT prolongation and torsades de pointes. *J Am Coll Cardiol*. 2016;67:1639-50.

**S10.7-11.** Hauptman PJ, Kelly RA. Digitalis. *Circulation*. 1999;99:1265-70.

**S10.7-12.** Kelly RA, Smith TW. Recognition and management of digitalis toxicity. *Am J Cardiol*. 1992;69:108G-18G.

**S10.7-13.** Osmonov D, Erdinler I, Ozcan KS, et al. Management of patients with drug-induced atrioventricular block. *Pacing Clin Electrophysiol*. 2012;35:804-10.

**S10.7-14.** Zipes DP, Camm AJ, Borggrefe M, et al. ACC/AHA/ESC 2006 guidelines for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death: a report of the American College of Cardiology/American Heart Association Task Force and the European Society of

Cardiology Committee for Practice Guidelines. *J Am Coll Cardiol*. 2006;48:e247-346.

**S10.7-15.** Behr ER, Ritchie MD, Tanaka T, et al. Genome wide analysis of drug-induced torsades de pointes: lack of common variants with large effect sizes. *PLoS One*. 2013;8:e78511.

**S10.7-16.** Kaab S, Crawford DC, Sinner MF, et al. A large candidate gene survey identifies the KCNE1 D85N polymorphism as a possible modulator of drug-induced torsades de pointes. *Circ Cardiovasc Genet*. 2012;5:91-9.

**S10.7-17.** Strauss DG, Vicente J, Johannessen L, et al. A common genetic variant risk score is associated with drug-induced QT prolongation and torsade de pointes risk: a pilot study. *Circulation*. 2017;135:1300-10.

**S10.7-18.** Yang P, Kanki H, Drolet B, et al. Allelic variants in long-QT disease genes in patients with drug-associated torsades de pointes. *Circulation*. 2002;105:1943-8.

**S10.7-19.** Weeke P, Mosley JD, Hanna D, et al. Exome sequencing implicates an increased burden of rare potassium channel variants in the risk of drug-induced long QT interval syndrome. *J Am Coll Cardiol*. 2014;63:1430-7.

**S10.7-20.** Echt DS, Liebson PR, Mitchell LB, et al. Mortality and morbidity in patients receiving encainide, flecainide, or placebo. The Cardiac Arrhythmia Suppression Trial. *N Engl J Med*. 1991;324:781-8.

**S10.7-21.** Chamberlain DA, Jewitt DE, Julian DG, et al. Oral mexiletine in high-risk patients after myocardial infarction. *Lancet*. 1980;2:1324-7.

**S10.7-22.** UK Rythmodan Multicentre Study Group. Oral disopyramide after admission to hospital with suspected acute myocardial infarction. U. K. Rythmodan Multicentre Study Group. *Postgrad Med J*. 1984;60:98-107.

**S10.7-23.** Crijns HJ, Van Gelder IC, Lie Kl. Supraventricular tachycardia mimicking ventricular tachycardia during flecainide treatment. *Am J Cardiol*. 1988;62:1303-6.

**S10.7-24.** Alings M, Wilde A. "Brugada" syndrome: clinical data and suggested pathophysiological mechanism. *Circulation*. 1999;99:666-73.

**S10.7-25.** Konigstein M, Rosso R, Topaz G, et al. Drug-induced Brugada syndrome: clinical characteristics and risk factors. *Heart Rhythm*. 2016;13:1083-7.

**S10.7-26.** Tada H, Sticherling C, Oral H, et al. Brugada syndrome mimicked by tricyclic antidepressant overdose. *J Cardiovasc Electrophysiol*. 2001;12:275.

**S10.7-27.** Littmann L, Monroe MH, Svenson RH. Brugada-type electrocardiographic pattern induced by cocaine. *Mayo Clin Proc*. 2000;75:845-9.

**S10.7-28.** Brugada drugs. Available at: <http://www.brugadadrugs.org>. Accessed October 6, 2016.

**S10.7-29.** Bajaj AK, Woosley RL, Roden DM. Acute electrophysiologic effects of sodium administration in dogs treated with O-desmethyl encainide. *Circulation*. 1989;80:994-1002.

**S10.7-30.** Myerburg RJ, Kessler KM, Cox MM, et al. Reversal of proarrhythmic effects of flecainide acetate and encainide hydrochloride by propranolol. *Circulation*. 1989;80:1571-9.

**S10.7-31.** Brumfield E, Bernard KR, Kabrhel C. Life-threatening flecainide overdose treated with intralipid

and extracorporeal membrane oxygenation. *Am J Emerg Med*. 2015;33:1840-5.

**S10.7-32.** Credible meds. Available at: <http://www.crediblemeds.org>. Accessed December 26, 2016.

#### 10.8. Adult Congenital Heart Disease

**S10.8-1.** Diller GP, Kempny A, Lioudakis E, et al. Left ventricular longitudinal function predicts life-threatening ventricular arrhythmia and death in adults with repaired tetralogy of fallot. *Circulation*. 2012;125:2440-6.

**S10.8-2.** Gatzoulis MA, Till JA, Somerville J, et al. Mechanoelectrical interaction in tetralogy of Fallot. QRS prolongation relates to right ventricular size and predicts malignant ventricular arrhythmias and sudden death. *Circulation*. 1995;92:231-7.

**S10.8-3.** Harrison DA, Harris L, Siu SC, et al. Sustained ventricular tachycardia in adult patients late after repair of tetralogy of Fallot. *J Am Coll Cardiol*. 1997;30:1368-73.

**S10.8-4.** Knauth AL, Gauvreau K, Powell AJ, et al. Ventricular size and function assessed by cardiac MRI predict major adverse clinical outcomes late after tetralogy of Fallot repair. *Heart*. 2008;94:211-6.

**S10.8-5.** Koyak Z, de Groot JR, Bouma BJ, et al. Sudden cardiac death in adult congenital heart disease: can the unpredictable be foreseen? *Europac*. 2017;19:401-6.

**S10.8-6.** Koyak Z, Harris L, de Groot JR, et al. Sudden cardiac death in adult congenital heart disease. *Circulation*. 2012;126:1944-54.

**S10.8-7.** Adamson L, Vohra HA, Haw MP. Does pulmonary valve replacement post repair of tetralogy of Fallot improve right ventricular function? *Interact Cardiovasc Thorac Surg*. 2009;9:520-7.

**S10.8-8.** Deal BJ, Scagliotti D, Miller SM, et al. Electrophysiologic drug testing in symptomatic ventricular arrhythmias after repair of tetralogy of Fallot. *Am J Cardiol*. 1987;59:1380-5.

**S10.8-9.** Khairy P, Van Hare GF, Balaji S, et al. PACES/HRS Expert consensus statement on the recognition and management of arrhythmias in adult congenital heart disease: developed in partnership between the Pediatric and Congenital Electrophysiology Society (PACES) and the Heart Rhythm Society (HRS). *Heart Rhythm*. 2014;11:e102-65.

**S10.8-10.** Sabate Rotes A, Connolly HM, Warnes CA, et al. Ventricular arrhythmia risk stratification in patients with tetralogy of Fallot at the time of pulmonary valve replacement. *Circ Arrhythm Electrophysiol*. 2015;8:110-6.

**S10.8-11.** Therrien J, Provost Y, Merchant N, et al. Optimal timing for pulmonary valve replacement in adults after tetralogy of Fallot repair. *Am J Cardiol*. 2005;95:779-82.

**S10.8-12.** Therrien J, Siu SC, Harris L, et al. Impact of pulmonary valve replacement on arrhythmia propensity late after repair of tetralogy of Fallot. *Circulation*. 2001;103:2489-94.

**S10.8-13.** Kella DK, Merchant FM, Veledar E, et al. Lesion-specific differences for implantable cardioverter defibrillator therapies in adults with congenital heart disease. *Pacing Clin Electrophysiol*. 2014;37:1492-8.

**S10.8-14.** Koyak Z, de Groot JR, Van Gelder IC, et al. Implantable cardioverter defibrillator therapy in adults

- S10.8-15.** Santharam S, Hudsmith L, Thorne S, et al. Long-term follow-up of implantable cardioverter-defibrillators in adult congenital heart disease patients: indications and outcomes. *Europace.* 2017;19:407-13.
- S10.8-16.** Vehmeijer JT, Brouwer TF, Limpens J, et al. Implantable cardioverter-defibrillators in adults with congenital heart disease: a systematic review and meta-analysis. *Eur Heart J.* 2016;37:1439-48.
- S10.8-17.** Yap SC, Roos-Hesselink JW, Hoendermis ES, et al. Outcome of implantable cardioverter defibrillators in adults with congenital heart disease: a multi-centre study. *Eur Heart J.* 2007;28:1854-61.
- S10.8-18.** Chandar JS, Wolff GS, Garson A Jr, et al. Ventricular arrhythmias in postoperative tetralogy of Fallot. *Am J Cardiol.* 1990;65:655-61.
- S10.8-19.** Khairy P, Landzberg MJ, Gatzoulis MA, et al. Value of programmed ventricular stimulation after tetralogy of Fallot repair: a multicenter study. *Circulation.* 2004;109:1994-2000.
- S10.8-20.** Khairy P, Harris L, Landzberg MJ, et al. Implantable cardioverter-defibrillators in tetralogy of Fallot. *Circulation.* 2008;117:363-70.
- S10.8-21.** Kapel GF, Reichlin T, Wijnmaalen AP, et al. Re-entry using anatomically determined isthmuses: a curable ventricular tachycardia in repaired congenital heart disease. *Circ Arrhythm Electrophysiol.* 2015;8:102-9.
- S10.8-22.** Kapel GF, Reichlin T, Wijnmaalen AP, et al. Left-sided ablation of ventricular tachycardia in adults with repaired tetralogy of Fallot: a case series. *Circ Arrhythm Electrophysiol.* 2014;7:889-97.
- S10.8-23.** Kapel GF, Sacher F, Dekkers OM, et al. Arrhythmogenic anatomical isthmuses identified by electroanatomical mapping are the substrate for ventricular tachycardia in repaired tetralogy of Fallot. *Eur Heart J.* 2017;38:268-76.
- S10.8-24.** van Zyl M, Kapa S, Padmanabhan D, et al. Mechanism and outcomes of catheter ablation for ventricular tachycardia in adults with repaired congenital heart disease. *Heart Rhythm.* 2016;13:1449-54.
- S10.8-25.** Zeppenfeld K, Schalij MJ, Bartelings MM, et al. Catheter ablation of ventricular tachycardia after repair of congenital heart disease: electroanatomic identification of the critical right ventricular isthmus. *Circulation.* 2007;116:2241-52.
- S10.8-26.** Khairy P, Harris L, Landzberg MJ, et al. Sudden death and defibrillators in transposition of the great arteries with intra-atrial baffles: a multicenter study. *Circ Arrhythm Electrophysiol.* 2008;1:250-7.
- S10.8-27.** Engelings CC, Helm PC, Abdul-Khalil H, et al. Cause of death in adults with congenital heart disease - an analysis of the German National Register for Congenital Heart Defects. *Int J Cardiol.* 2016;211:31-6.
- S10.8-28.** Gallego P, Gonzalez AE, Sanchez-Recalde A, et al. Incidence and predictors of sudden cardiac arrest in adults with congenital heart defects repaired before adult life. *Am J Cardiol.* 2012;110:109-17.
- S10.8-29.** Tutarel O, Kempny A, Alonso-Gonzalez R, et al. Congenital heart disease beyond the age of 60: emergence of a new population with high resource utilization, high morbidity, and high mortality. *Eur Heart J.* 2014;35:725-32.
- S10.8-30.** Fish FA, Gillette PC, Benson DW Jr. Proarrhythmia, cardiac arrest and death in young patients receiving encainide and flecainide. The Pediatric Electrophysiology Group. *J Am Coll Cardiol.* 1991;18:356-65.
- S10.8-31.** Stan MN, Sathananthan M, Warnes C, et al. Amiodarone-induced thyrotoxicosis in adults with congenital heart disease-clinical presentation and response to therapy. *Endocr Pract.* 2014;21:33-40.
- S10.8-32.** Thorne SA, Barnes I, Cullinan P, et al. Amiodarone-associated thyroid dysfunction: risk factors in adults with congenital heart disease. *Circulation.* 1999;100:149-54.
- S10.8-33.** Garson A Jr, Porter CB, Gillette PC, et al. Induction of ventricular tachycardia during electrophysiologic study after repair of tetralogy of Fallot. *J Am Coll Cardiol.* 1983;1:1493-502.
- S10.8-34.** Khairy P, Aboulhosn J, Gurvitz MZ, et al. Arrhythmia burden in adults with surgically repaired tetralogy of Fallot: a multi-institutional study. *Circulation.* 2010;122:868-75.
- S10.8-35.** Tsai SF, Chan DP, Ro PS, et al. Rate of inducible ventricular arrhythmia in adults with congenital heart disease. *Am J Cardiol.* 2010;106:730-6.
- S10.8-36.** Valente AM, Gauvreau K, Assenza GE, et al. Contemporary predictors of death and sustained ventricular tachycardia in patients with repaired tetralogy of Fallot enrolled in the INDICATOR cohort. *Heart.* 2014;100:247-53.
- S10.8-37.** Sakamoto T, Nagashima M, Hiramatsu T, et al. Fontan circulation over 30 years. What should we learn from those patients? *Asian Cardiovasc Thorac Ann.* 2016;24:765-71.
- S10.8-38.** Harrild DM, Berul CI, Cecchin F, et al. Pulmonary valve replacement in tetralogy of Fallot: impact on survival and ventricular tachycardia. *Circulation.* 2009;119:445-51.
- S10.8-39.** Berul CI, Van Hare GF, Kertesz NJ, et al. Results of a multicenter retrospective implantable cardioverter-defibrillator registry of pediatric and congenital heart disease patients. *J Am Coll Cardiol.* 2008;51:1685-91.
- S10.8-40.** Khanna AD, Warnes CA, Phillips SD, et al. Single-center experience with implantable cardioverter-defibrillators in adults with complex congenital heart disease. *Am J Cardiol.* 2011;108:729-34.
- S10.8-41.** Moore JP, Mondesert B, Lloyd MS, et al. Clinical experience with the subcutaneous implantable cardioverter-defibrillator in adults with congenital heart disease. *Circ Arrhythm Electrophysiol.* 2016;9:e004338.
- S10.8-42.** Okamura H, McLeod CJ, DeSimone CV, et al. Right parasternal lead placement increases eligibility for subcutaneous implantable cardioverter defibrillator therapy in adults with congenital heart disease. *Circ J.* 2016;80:1328-35.
- S10.8-43.** Witte KK, Pepper CB, Cowan JC, et al. Implantable cardioverter-defibrillator therapy in adult patients with tetralogy of Fallot. *Europace.* 2008;10:926-30.
- S10.8-44.** Raissadati A, Nieminen H, Haukka J, et al. Late causes of death after pediatric cardiac surgery: a 60-year population-based study. *J Am Coll Cardiol.* 2016;68:487-98.
- S10.8-45.** Diller GP, Kempny A, Alonso-Gonzalez R, et al. Survival prospects and circumstances of death in contemporary adult congenital heart disease patients under follow-up at a large tertiary centre. *Circulation.* 2015;132:2118-25.
- S10.8-46.** Nieminen HP, Jokinen EV, Sairanen HI. Causes of late deaths after pediatric cardiac surgery: a population-based study. *J Am Coll Cardiol.* 2007;50:1263-71.
- S10.8-47.** Zomer AC, Vaartjes I, Uiterwaal CS, et al. Circumstances of death in adult congenital heart disease. *Int J Cardiol.* 2012;154:168-72.
- S10.8-48.** Lange R, Horer J, Kostolny M, et al. Presence of a ventricular septal defect and the Mustard operation are risk factors for late mortality after the atrial switch operation: thirty years of follow-up in 417 patients at a single center. *Circulation.* 2006;114:1905-13.
- S10.8-49.** Lubiszewska B, Gosiewska E, Hoffman P, et al. Myocardial perfusion and function of the systemic right ventricle in patients after atrial switch procedure for complete transposition: long-term follow-up. *J Am Coll Cardiol.* 2000;36:1365-70.
- S10.8-50.** Millane T, Bernard EJ, Jaeggi E, et al. Role of ischemia and infarction in late right ventricular dysfunction after atrial repair of transposition of the great arteries. *J Am Coll Cardiol.* 2000;35:1661-8.
- S10.8-51.** Scherzmann M, Salehian O, Harris L, et al. Ventricular arrhythmias and sudden death in adults after a Mustard operation for transposition of the great arteries. *Eur Heart J.* 2009;30:1873-9.
- S10.8-52.** Buber J, Ackley TJ, Daniels CJ, et al. Outcomes following the implantation of cardioverter-defibrillator for primary prevention in transposition of the great arteries after intra-atrial baffle repair: a single-centre experience. *Europace.* 2016;18:1016-22.
- S10.8-53.** Backhoff D, Kerst G, Peters A, et al. Internal cardioverter defibrillator indications and therapies after atrial baffle procedure for d-transposition of the great arteries: a multicenter analysis. *Pacing Clin Electrophysiol.* 2016;39:1070-6.
- S10.8-54.** Oechslin EN, Harrison DA, Connelly MS, et al. Mode of death in adults with congenital heart disease. *Am J Cardiol.* 2000;86:1111-6.
- S10.8-55.** Verheugt CL, Uiterwaal CS, van der Velde ET, et al. Mortality in adult congenital heart disease. *Eur Heart J.* 2010;31:1220-9.
- S10.8-56.** Silka MJ, Hardy BG, Menashe VD, et al. A population-based prospective evaluation of risk of sudden cardiac death after operation for common congenital heart defects. *J Am Coll Cardiol.* 1998;32:245-51.
- S10.8-57.** Abou Hassan OK, Fahed AC, Batrawi M, et al. NKX2-5 mutations in an inbred consanguineous population: genetic and phenotypic diversity. *Sci Rep.* 2015;5:8848.
- S10.8-58.** El Malti R, Liu H, Doray B, et al. A systematic variant screening in familial cases of congenital heart defects demonstrates the usefulness of molecular genetics in this field. *Eur J Human Genet.* 2016;24:228-36.
- S10.8-59.** Ellesoe SG, Johansen MM, Bjerre JV, et al. Familial atrial septal defect and sudden cardiac death:

identification of a novel NKX2-5 mutation and a review of the literature. *Congenit Heart Dis.* 2016;11:283-90.

**S10.8-60.** Cuypers JA, Opic P, Menting ME, et al. The unnatural history of an atrial septal defect: longitudinal 35 year follow up after surgical closure at young age. *Heart.* 2013;99:1346-52.

**S10.8-61.** Kuijpers JM, van der Bom T, van Riel AC, et al. Secundum atrial septal defect is associated with reduced survival in adult men. *Eur Heart J.* 2015;36:2079-86.

**S10.8-62.** Verheugt CL, Uiterwaal CS, Grobbee DE, et al. Long-term prognosis of congenital heart defects: a systematic review. *Int J Cardiol.* 2008;131:25-32.

**S10.8-63.** Engelfriet P, Boersma E, Oechslin E, et al. The spectrum of adult congenital heart disease in Europe: morbidity and mortality in a 5 year follow-up period. The Euro Heart Survey on adult congenital heart disease. *Eur Heart J.* 2005;26:2325-33.

**S10.8-64.** Arya S, Kovach J, Singh H, et al. Arrhythmias and sudden death among older children and young adults following tetralogy of Fallot repair in the current era: are previously reported risk factors still applicable? *Congenit Heart Dis.* 2014;9:407-14.

**S10.8-65.** Wu MH, Lu CW, Chen HC, et al. Arrhythmic burdens in patients with tetralogy of Fallot: a national database study. *Heart Rhythm.* 2015;12:604-9.

**S10.8-66.** Barona F, Valente AM, Porayette P, et al. Coronary arteries in childhood heart disease: implications for management of young adults. *J Clin Exp Cardiol.* 2012; suppl 8:006.

**S10.8-67.** Naimo PS, Fricke TA, Yong MS, et al. Outcomes of truncus arteriosus repair in children: 35 years of experience from a single institution. *Semin Thorac Cardiovasc Surg.* 2016;28:500-11.

**S10.8-68.** Pundi KN, Pundi KN, Johnson JN, et al. Sudden cardiac death and late arrhythmias after the Fontan operation. *Congenit Heart Dis.* 2017;12:17-23.

## 11. DEFIBRILLATORS OTHER THAN TRANSVENOUS ICDs

### 11.1. Subcutaneous Implantable Cardioverter-Defibrillator

**S11.1-1.** Bardy GH, Smith WM, Hood MA, et al. An entirely subcutaneous implantable cardioverter-defibrillator. *N Engl J Med.* 2010;363:36-44.

**S11.1-2.** Weiss R, Knight BP, Gold MR, et al. Safety and efficacy of a totally subcutaneous implantable cardioverter defibrillator. *Circulation.* 2013;128:944-53.

**S11.1-3.** Lambiase PD, Barr C, Theuns DA, et al. Worldwide experience with a totally subcutaneous implantable defibrillator: early results from the EFFORTLESS S-ICD Registry. *Eur Heart J.* 2014;35:1657-65.

**S11.1-4.** Burke MC, Gold MR, Knight BP, et al. Safety and efficacy of the totally subcutaneous implantable defibrillator: 2-year results from a pooled analysis of the IDE study and EFFORTLESS registry. *J Am Coll Cardiol.* 2015;65:1605-15.

**S11.1-5.** El-Chami MF, Levy M, Kelli HM, et al. Outcome of subcutaneous implantable cardioverter defibrillator implantation in patients with end-stage

renal disease on dialysis. *J Cardiovasc Electrophysiol.* 2015;26:900-4.

**S11.1-6.** de Bie MK, Thijssen J, van Rees JB, et al. Suitability for subcutaneous defibrillator implantation: results based on data from routine clinical practice. *Heart.* 2013;99:1018-23.

**S11.1-7.** Olde Nordkamp LR, Dabiri AL, Boersma LV, et al. The entirely subcutaneous implantable cardioverter-defibrillator: initial clinical experience in a large Dutch cohort. *J Am Coll Cardiol.* 2012;60:1933-9.

**S11.1-8.** Köbe J, Reinke F, Meyer C, et al. Implantation and follow-up of totally subcutaneous versus conventional implantable cardioverter-defibrillators: a multicenter case-control study. *Heart Rhythm.* 2013;10:29-36.

**S11.1-9.** Groh CA, Sharma S, Pelchovitz DJ, et al. Use of an electrocardiographic screening tool to determine candidacy for a subcutaneous implantable cardioverter-defibrillator. *Heart Rhythm.* 2014;11:1361-6.

**S11.1-10.** Olde Nordkamp LR, Warnaars JL, Kooiman KM, et al. Which patients are not suitable for a subcutaneous ICD: incidence and predictors of failed QRS-T-wave morphology screening. *J Cardiovasc Electrophysiol.* 2014;25:494-9.

**S11.1-11.** Randles DA, Hawkins NM, Shaw M, et al. How many patients fulfil the surface electrocardiogram criteria for subcutaneous implantable cardioverter-defibrillator implantation? *Europace.* 2014;16:1015-21.

**S11.1-12.** Al-Khatib SM, Greiner MA, Peterson ED, et al. Patient and implanting physician factors associated with mortality and complications after implantable cardioverter-defibrillator implantation, 2002-2005. *Circ Arrhythm Electrophysiol.* 2008;1:240-9.

**S11.1-13.** Olde Nordkamp LR, Knops RE, Bardy GH, et al. Rationale and design of the PRAETORIAN trial: a Prospective, RANdomizEd comparison of subcuTaneOus and tRansvenous ImplANTable cardioverteR-defibrillator therapy. *Am Heart J.* 2012;163:753-60.

### 11.2. Wearable Cardioverter-Defibrillator

**S11.2-1.** Chung MK. The role of the wearable cardioverter defibrillator in clinical practice. *Cardiol Clin.* 2014;32:253-70.

**S11.2-2.** Chung MK, Szymkiewicz SJ, Shao M, et al. Aggregate national experience with the wearable cardioverter-defibrillator: event rates, compliance, and survival. *J Am Coll Cardiol.* 2010;56:194-203.

**S11.2-3.** Klein HU, Meltendorf U, Reek S, et al. Bridging a temporary high risk of sudden arrhythmic death. Experience with the wearable cardioverter defibrillator (WCD). *Pacing Clin Electrophysiol.* 2010;33:353-67.

**S11.2-4.** Piccini JP Sr., Allen LA, Kudenchuk PJ, et al. Wearable cardioverter-defibrillator therapy for the prevention of sudden cardiac death: a science advisory from the American Heart Association. *Circulation.* 2016;133:1715-27.

**S11.2-5.** Ellenbogen KA, Koneru JN, Sharma PS, et al. Benefit of the wearable cardioverter-defibrillator in protecting patients after implantable-cardioverter defibrillator explant. Results from the National Registry. *JACC Clin Electrophysiol.* 2017;3:243-50.

**S11.2-6.** LifeVest Wearable Cardioverter Defibrillator. Available at: <http://www.fda.gov/MedicalDevices/ProductsandMedicalProcedures/DeviceApprovalsandClearances/Recently-ApprovedDevices/ucm458494.htm>. Accessed August 26, 2016.

Clearances/Recently-ApprovedDevices/ucm458494.htm. Accessed August 26, 2016.

### 11.3. Automated External Defibrillator

**S11.3-1.** Marengo JP, Wang PJ, Link MS, et al. Improving survival from sudden cardiac arrest: the role of the automated external defibrillator. *JAMA.* 2001;285:1193-200.

**S11.3-2.** Priori SG, Bossaert LL, Chamberlain DA, et al. ESC-ERC recommendations for the use of automated external defibrillators (AEDs) in Europe. *Eur Heart J.* 2004;25:437-5

**S11.3-3.** Blom MT, Beesems SG, Homma PC, et al. Improved survival after out-of-hospital cardiac arrest and use of automated external defibrillators. *Circulation.* 2014;130:1868-75.

**S11.3-4.** Koster RW. Automatic external defibrillator: key link in the chain of survival. *J Cardiovasc Electrophysiol.* 2002;13:S92-5.

**S11.3-5.** Valenzuela TD, Roe DJ, Nichol G, et al. Outcomes of rapid defibrillation by security officers after cardiac arrest in casinos. *N Engl J Med.* 2000;343:1206-9.

**S11.3-6.** Caffrey SL, Willoughby PJ, Pepe PE, et al. Public use of automated external defibrillators. *N Engl J Med.* 2002;347:1242-7.

**S11.3-7.** Page RL, Joglar JA, Kowal RC, et al. Use of automated external defibrillators by a U.S. airline. *N Engl J Med.* 2000;343:1210-6.

**S11.3-8.** Page RL, Husain S, White LY, et al. Cardiac arrest at exercise facilities: implications for placement of automated external defibrillators. *J Am Coll Cardiol.* 2013;62:2102-9.

**S11.3-9.** Weisfeldt ML, Sitrani CM, Ornato JP, et al. Survival after application of automatic external defibrillators before arrival of the emergency medical system: evaluation in the resuscitation outcomes consortium population of 21 million. *J Am Coll Cardiol.* 2010;55:1713-20.

**S11.3-10.** Hansen CM, Lippert FK, Wissenberg M, et al. Temporal trends in coverage of historical cardiac arrests using a volunteer-based network of automated external defibrillators accessible to laypersons and emergency dispatch centers. *Circulation.* 2014;130:1859-67.

**S11.3-11.** Weisfeldt ML, Everson-Stewart S, Sitrani C, et al. Ventricular tachyarrhythmias after cardiac arrest in public versus at home. *N Engl J Med.* 2011;364:313-21.

**S11.3-12.** Bardy GH, Lee KL, Mark DB, et al. Home use of automated external defibrillators for sudden cardiac arrest. *N Engl J Med.* 2008;358:1793-804.

**S11.3-13.** Travers AH, Perkins GD, Berg RA, et al. Part 3: adult basic life support and automated external defibrillation: 2015 International Consensus on Cardiopulmonary Resuscitation and Emergency Cardiovascular Care Science With Treatment Recommendations. *Circulation.* 2015;132:S51-83.

## 12. SPECIAL CONSIDERATIONS FOR CATHETER ABLATION

**S12-1.** Blanck Z, Dhala A, Deshpande S, et al. Bundle branch reentrant ventricular tachycardia: cumulative

experience in 48 patients. *J Cardiovasc Electrophysiol.* 1993;4:253–62.

**S12-2.** Lopera G, Stevenson WG, Soejima K, et al. Identification and ablation of three types of ventricular tachycardia involving the his-purkinje system in patients with heart disease. *J Cardiovasc Electrophysiol.* 2004;15:52–8.

**S12-3.** Mehdrad AA, Keim S, Rist K, et al. Long-term clinical outcome of right bundle branch radiofrequency catheter ablation for treatment of bundle branch reentrant ventricular tachycardia. *Pacing Clin Electrophysiol.* 1995;18:2135–43.

**S12-4.** Dinov B, Fiedler L, Schonbauer R, et al. Outcomes in catheter ablation of ventricular tachycardia in dilated nonischemic cardiomyopathy compared with ischemic cardiomyopathy: results from the Prospective Heart Centre of Leipzig VT (HELP-VT) study. *Circulation.* 2014;129:728–36.

**S12-5.** Tanner H, Hindricks G, Volkmer M, et al. Catheter ablation of recurrent scar-related ventricular tachycardia using electroanatomical mapping and irrigated ablation technology: results of the prospective multicenter Euro-VT-study. *J Cardiovasc Electrophysiol.* 2010;21:47–53.

**S12-6.** Marchlinski FE, Haffajee CI, Beshai JF, et al. Long-term success of irrigated radiofrequency catheter ablation of sustained ventricular tachycardia: post-approval THERMOCOOL VT trial. *J Am Coll Cardiol.* 2016;67:674–83.

**S12-7.** Assomull RG, Prasad SK, Lyne J, et al. Cardiovascular magnetic resonance, fibrosis, and prognosis in dilated cardiomyopathy. *J Am Coll Cardiol.* 2006;48:1977–85.

**S12-8.** Bogun FM, Desjardins B, Good E, et al. Delayed-enhanced magnetic resonance imaging in nonischemic cardiomyopathy: utility for identifying the ventricular arrhythmia substrate. *J Am Coll Cardiol.* 2009;53:1138–45.

**S12-9.** Roberts WC, Siegel RJ, McManus BM. Idiopathic dilated cardiomyopathy: analysis of 152 necropsy patients. *Am J Cardiol.* 1987;60:1340–55.

**S12-10.** Piers SR, Tao Q, van Huls van Taxis CFB, et al. Contrast-enhanced MRI-derived scar patterns and associated ventricular tachycardias in nonischemic cardiomyopathy: implications for the ablation strategy. *Circ Arrhythm Electrophysiol.* 2013;6:875–83.

**S12-11.** Sacher F, Roberts-Thomson K, Maury P, et al. Epicardial ventricular tachycardia ablation a multicenter safety study. *J Am Coll Cardiol.* 2010;55:2366–72.

**S12-12.** Della Bella P, Brugada J, Zeppenfeld K, et al. Epicardial ablation for ventricular tachycardia: a European multicenter study. *Circ Arrhythm Electrophysiol.* 2011;4:653–9.

### 13. POSTMORTEM EVALUATION OF SCD

**S13-1.** Basso C, Burke M, Fornes P, et al. Guidelines for autopsy investigation of sudden cardiac death. *Virchows Arch.* 2008;452:11–8.

**S13-2.** de Noronha SV, Behr ER, Papadakis M, et al. The importance of specialist cardiac histopathological examination in the investigation of young sudden cardiac deaths. *Europace.* 2014;16:899–907.

**S13-3.** Wilde AA, Behr ER. Genetic testing for inherited cardiac disease. *Nature reviews. Cardiology.* 2013;10:571–83.

**S13-4.** Tester DJ, Medeiros-Domingo A, Will ML, et al. Cardiac channel molecular autopsy: insights from 173 consecutive cases of autopsy-negative sudden unexplained death referred for postmortem genetic testing. *Mayo Clin Proc.* 2012;87:524–39.

**S13-5.** Tang Y, Stahl-Herz J, Sampson BA. Molecular diagnostics of cardiovascular diseases in sudden unexplained death. *Cardiovasc Pathol.* 2014;23:1–4.

**S13-6.** Bagnall RD, Das KJ, Duflou J, et al. Exome analysis-based molecular autopsy in cases of sudden unexplained death in the young. *Heart Rhythm.* 2014;11:655–62.

**S13-7.** Bagnall RD, Weintraub RG, Ingles J, et al. A prospective study of sudden cardiac death among children and young adults. *N Engl J Med.* 2016;374:2441–52.

**S13-8.** Semsarian C, Ingles J. Molecular autopsy in victims of inherited arrhythmias. *J Arrhythm.* 2016;32:359–65.

**S13-9.** Vassalini M, Verzeletti A, Restori M, et al. An autopsy study of sudden cardiac death in persons aged 1–40 years in Brescia (Italy). *J Cardiovasc Med.* 2016;17:446–53.

**S13-10.** Wu Q, Zhang L, Zheng J, et al. Forensic pathological study of 1656 cases of sudden cardiac death in southern China. *Medicine.* 2016;95:e2707.

**S13-11.** Giudici V, Spanaki A, Hendry J, et al. Sudden arrhythmic death syndrome: diagnostic yield of comprehensive clinical evaluation of pediatric first-degree relatives. *Pacing Clin Electrophysiol.* 2014;37:1681–5.

**S13-12.** Steinberg C, Padfield GJ, Champagne J, et al. Cardiac abnormalities in first-degree relatives of unexplained cardiac arrest victims: a report from the Cardiac Arrest Survivors With Preserved Ejection Fraction Registry. *Circ Arrhythm Electrophysiol.* 2016;9:e004274.

**S13-13.** Wong LC, Roses-Noguer F, Till JA, et al. Cardiac evaluation of pediatric relatives in sudden arrhythmic death syndrome: a 2-center experience. *Circ Arrhythm Electrophysiol.* 2014;7:800–6.

**S13-14.** van der Werf C, Stiekema L, Tan HL, et al. Low rate of cardiac events in first-degree relatives of diagnosis-negative young sudden unexplained death syndrome victims during follow-up. *Heart Rhythm.* 2014;11:1728–32.

**S13-15.** Papadakis M, Raju H, Behr ER, et al. Sudden cardiac death with autopsy findings of uncertain significance: potential for erroneous interpretation. *Circ Arrhythm Electrophysiol.* 2013;6:588–96.

**S13-16.** Priori SG, Wilde AA, Horie M, et al. HRS/EHRA/APHRS expert consensus statement on the diagnosis and management of patients with inherited primary arrhythmia syndromes. *Heart Rhythm.* 2013;10:1932–63.

### 14. TERMINAL CARE

**S14-1.** Berger JT, Gorski M, Cohen T. Advance health planning and treatment preferences among recipients of implantable cardioverter defibrillators: an exploratory study. *J Clin Ethics.* 2006;17:72–8.

**S14-2.** Lewis KB, Stacey D, Matlock DD. Making decisions about implantable cardioverter-defibrillators from implantation to end of life: an integrative review of patients' perspectives. *Patient.* 2014;7:243–60.

**S14-3.** Hill L, McIlpatrick S, Taylor B, et al. Patients' perception of implantable cardioverter defibrillator deactivation at the end of life. *Palliat Med.* 2015;29:310–23.

**S14-4.** Hauptman PJ, Chibnall JT, Guild C, et al. Patient perceptions, physician communication, and the implantable cardioverter-defibrillator. *JAMA Intern Med.* 2013;173:571–7.

**S14-5.** Matlock DD, Nowels CT, Masoudi FA, et al. Patient and cardiologist perceptions on decision making for implantable cardioverter-defibrillators: a qualitative study. *Pacing Clin Electrophysiol.* 2011;34:1634–44.

**S14-6.** Buchhalter LC, Ottenberg AL, Webster TL, et al. Features and outcomes of patients who underwent cardiac device deactivation. *JAMA Intern Med.* 2014;174:80–5.

**S14-7.** Dodson JA, Fried TR, Van Ness PH, et al. Patient preferences for deactivation of implantable cardioverter-defibrillators. *JAMA Intern Med.* 2013;173:377–9.

**S14-8.** Goldstein NE, Lampert R, Bradley E, et al. Management of implantable cardioverter defibrillators in end-of-life care. *Ann Intern Med.* 2004;141:835–8.

**S14-9.** Goldstein N, Carlson M, Livote E, et al. Brief communication: management of implantable cardioverter-defibrillators in hospice: a nationwide survey. *Ann Intern Med.* 2010;152:296–9.

**S14-10.** Kirkpatrick JN, Gottlieb M, Sehgal P, et al. Deactivation of implantable cardioverter defibrillators in terminal illness and end of life care. *Am J Cardiol.* 2012;109:91–4.

**S14-11.** Kelley AS, Reid MC, Miller DH, et al. Implantable cardioverter-defibrillator deactivation at the end of life: a physician survey. *Am Heart J.* 2009;157:702–8.

**S14-12.** Kramer DB, Kesselheim AS, Brock DW, et al. Ethical and legal views of physicians regarding deactivation of cardiac implantable electrical devices: a quantitative assessment. *Heart Rhythm.* 2010;7:1537–42.

**S14-13.** Mueller PS, Jenkins SM, Bramstedt KA, et al. Deactivating implanted cardiac devices in terminally ill patients: practices and attitudes. *Pacing Clin Electrophysiol.* 2008;31:560–8.

### 15. SHARED DECISION-MAKING

**S15-1.** Lewis KB, Stacey D, Matlock DD. Making decisions about implantable cardioverter-defibrillators from implantation to end of life: an integrative review of patients' perspectives. *Patient.* 2014;7:243–60.

**S15-2.** Stewart GC, Weintraub JR, Pratibhu PP, et al. Patient expectations from implantable defibrillators to prevent death in heart failure. *J Card Fail.* 2010;16:106–13.

**S15-3.** Hauptman PJ, Chibnall JT, Guild C, et al. Patient perceptions, physician communication, and the implantable cardioverter-defibrillator. *JAMA Intern Med.* 2013;173:571–7.

**S15-4.** Ottenberg AL, Mueller PS, Topazian RJ, et al. "It's not broke, so let's not try to fix it": why patients decline a cardiovascular implantable electronic device. *Pacing Clin Electrophysiol.* 2014;37:1306-14.

**S15-5.** Yuhas J, Mattocks K, Gravelin L, et al. Patients' attitudes and perceptions of implantable cardioverter-defibrillators: potential barriers to appropriate primary prophylaxis. *Pacing Clin Electrophysiol.* 2012;35:1179-87.

**S15-6.** Vig EK, Pearlman RA. Good and bad dying from the perspective of terminally ill men. *Arch Intern Med.* 2004;164:977-81.

**S15-7.** Charles C, Gafni A, Whelan T. Shared decision-making in the medical encounter: what does it mean? (or it takes at least two to tango). *Soc Sci Med.* 1997; 44:681-92.

**S15-8.** Quill TE, Brody H. Physician recommendations and patient autonomy: finding a balance between physician power and patient choice. *Ann Intern Med.* 1996;125:763-9.

## 16. COST AND VALUE CONSIDERATIONS

**S16-1.** Anderson JL, Heidenreich PA, Barnett PG, et al. ACC/AHA statement on cost/value methodology in clinical practice guidelines and performance measures: a report of the American College of Cardiology/American Heart Association Task Force on Performance Measures and Task Force on Practice Guidelines. *J Am Coll Cardiol.* 2014;63:2304-22.

**S16-2.** Sanders GD, Neumann PJ, Basu A, et al. Recommendations for conduct, methodological practices, and reporting of cost-effectiveness analyses: Second Panel on Cost-Effectiveness in Health and Medicine. *JAMA.* 2016;316:1093-103.

**S16-3.** Moshlin AI, Hall J, Zwanziger J, et al. The cost-effectiveness of automatic implantable cardiac defibrillators. Results from MADIT. *Circulation.* 1998;97: 2129-35.

**S16-4.** O'Brien BJ, Connolly SJ, Goeree R, et al. Cost-effectiveness of the implantable cardioverter-defibrillator. Results from the Canadian Implantable Defibrillator Study (CIDS). *Circulation.* 2001;103: 1416-21.

**S16-5.** Larson G, Hallstrom A, McAnulty J, et al. Cost-effectiveness of the implantable cardioverter-defibrillator versus antiarrhythmic drugs in survivors of serious ventricular tachyarrhythmias. Results of the Antiarrhythmics Versus Implantable Defibrillators (AVID) Economic Analysis Substudy. *Circulation.* 2002; 105:2049-57.

**S16-6.** Zwanziger J, Hall WJ, Dick AW, et al. The cost-effectiveness of implantable cardioverter-defibrillators. Results from the Multicenter Automatic Defibrillator Implantation Trial (MADIT)-II. *J Am Coll Cardiol.* 2006;47:2310-8.

**S16-7.** Mark DB, Nelson CL, Anstrom KJ, et al. Cost-effectiveness of defibrillator therapy or amiodarone in chronic stable heart failure. Results from the Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT). *Circulation.* 2006;114:135-42.

**S16-8.** Weiss JP, Saynina O, McDonald KM, et al. Effectiveness and cost-effectiveness of implantable cardioverter defibrillators in the treatment of ventricular arrhythmias among Medicare beneficiaries. *Am J Med.* 2002;112:519-27.

**S16-9.** Al-Khatib SM, Anstrom KJ, Eisenstein EL, et al. Clinical and economic implications of the Multicenter Automatic Defibrillator Implantation Trial-II. *Ann Intern Med.* 2005;142:593-600.

**S16-10.** Buxton M, Caine N, Chase D, et al. A review of the evidence on the effects and costs of implantable cardioverter defibrillator therapy in different patient groups, and modelling of cost-effectiveness and cost-utility for these groups in a UK context. *Health Technol Assess.* 2006;10:iii-iv, ix-xi, 1-164.

**S16-11.** Sanders GD, Hlatky MA, Owens DK. Cost effectiveness of implantable cardioverter-defibrillators. *N Engl J Med.* 2005;353:1471-80.

**S16-12.** Smith T, Jordans L, Theuns DAMJ, et al. The cost-effectiveness of primary prophylactic implantable defibrillator therapy in patients with ischaemic or non-ischaemic heart disease: A European analysis. *Eur Heart J.* 2013;34:211-9.

**S16-13.** Cowie MR, Marshall D, Drummond M, et al. Lifetime cost-effectiveness of prophylactic implantation of cardioverter defibrillator in patients with reduced left ventricular systolic function: Results of Markov modelling in a European population. *Europace.* 2008;11:716-26.

**S16-14.** Goldenberg I, Moss AJ, Maron BJ, et al. Cost-effectiveness of implanted defibrillators in young people with inherited cardiac arrhythmias. *Ann Noninvasive Electrocardiol.* 2005;10 suppl:67-83.

**S16-15.** Bigger JT. Prophylactic use of implanted cardiac defibrillators in patients at high risk for ventricular arrhythmias after coronary-artery bypass graft surgery. *N Engl J Med.* 1997;337:1569-75.

**S16-16.** Hohnloser SH, Kuck KH, Dorian P, et al. Prophylactic use of an implantable cardioverter-defibrillator after acute myocardial infarction. *N Engl J Med.* 2004;351:2481-8.

**S16-17.** Steinbeck G, Andresen D, Seidl K, et al. Defibrillator implantation early after myocardial infarction. *N Engl J Med.* 2009;361:1427-36.

**S16-18.** Connolly SJ, Hallstrom AP, Cappato R, et al. Meta-analysis of the implantable cardioverter defibrillator secondary prevention trials. *Eur Heart J.* 2000;21:2071-8.

**S16-19.** Owens DK, Sanders GD, Heidenreich PA, et al. Effect of risk stratification on cost-effectiveness of the implantable cardioverter defibrillator. *Am Heart J.* 2002;144:440-8.

**S16-20.** Ezekowitz JA, Rowe BH, Dryden DM, et al. Systematic review: implantable cardioverter defibrillators for adults with left ventricular systolic dysfunction. *Ann Intern Med.* 2007;147:251-62.

## 17. QUALITY OF LIFE

**S17-1.** Mark DB, Anstrom KJ, Sun JL, et al. Quality of life with defibrillator therapy or amiodarone in heart failure. *N Engl J Med.* 2008;359:999-1008.

**S17-2.** Noyes K, Corona E, Veazie P, et al. Examination of the effect of implantable cardioverter-defibrillators on health-related quality of life: based on results from the Multicenter Automatic Defibrillator Trial-II. *American journal of cardiovascular drugs: drugs, devices, and other interventions.* 2009;9:393-400.

**S17-3.** Passman R, Subacius H, Ruo B, et al. Implantable cardioverter defibrillators and quality of life: results from the defibrillators in nonischemic cardiomyopathy treatment evaluation study. *Archives of internal medicine.* 2007;167:2226-32.

**KEY WORDS** ACC/AHA Clinical Practice Guidelines, acute coronary syndrome, ambulatory ECG monitoring, antiarrhythmic drug therapy, arrhythmogenic cardiomyopathy, athletes, cardiac electrophysiology, cardiac resynchronization therapy, cardiomyopathy, catheter ablation, congenital heart disease, CT imaging, ECG, echocardiography, electrophysiological testing, genetic arrhythmias, Guidelines, heart failure, imaging, implantable cardioverter-defibrillator, implantable and external cardioverter devices, medication-induced arrhythmias, MR imaging, myocardial infarction, premature ventricular beats, resuscitation, sarcoidosis, specific pathology (e.g., congenital heart disease, myocarditis, renal failure), stable coronary artery disease, sudden cardiac arrest, sudden cardiac death, torsades de pointes, ventricular fibrillation, ventricular tachycardia

**APPENDIX 1. AUTHOR RELATIONSHIPS WITH INDUSTRY AND OTHER ENTITIES (RELEVANT)—  
2017 AHA/ACC/HRS GUIDELINE FOR MANAGEMENT OF PATIENTS WITH VENTRICULAR ARRHYTHMIAS  
AND THE PREVENTION OF SUDDEN CARDIAC DEATH (OCTOBER 2017)**

Committee Member	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness	Voting Recusals by Section*
Sana M. Al-Khatib (Chair)	Duke Clinical Research Institute; Duke University—Professor of Medicine	None	None	None	None	None	None	None
William G. Stevenson (Vice Chair)	Vanderbilt University Medical Center—Professor; Brigham and Women's Hospital—Director of Clinical Cardiac EP	■ St. Jude Medical ■ Boston Scientific ■ Biosense Webster‡			None	None	None	4.1, 4.2.2, 4.2.3, 5, 10.1, 5.4, 5.6, 6, 7, 8, 9 (except 9.7), 13, 15
Michael J. Ackerman	Mayo Clinic—Professor of Medicine, Pediatrics, and Pharmacology; Long QT Syndrome/Genetic Heart Rhythm Clinic and the Mayo Clinic Windland Smith Rice Sudden Death Genomics Laboratory—Director	■ Audentes Therapeutics ■ Boston Scientific ■ Gilead Sciences ■ Invitae ■ Medtronic ■ MyoKardia ■ St. Jude Medical	None	None	None	■ Transgenomic (Familion)† ■ Blue Ox Health Corporation‡ ■ AliveCor‡ ■ StemoniX‡	None	4.1, 4.2.2, 4.2.3, 5 (except 5.1.5.2, 5.5), 6, 7, 8, 9 (except 9.7), 10 (except 10.2) 11, 13, 15
William J. Bryant	Dominick Feld Hyde—Attorney at Law	None	None	None	None	None	None	None
David J. Callans	University of Pennsylvania Health System—Professor of Medicine; Associate Director of EP	■ Biosense Webster† ■ Biotronik ■ Boston Scientific† ■ Medtronic ■ St. Jude Medical	None	None	■ Biosense Webster (PI)‡ ■ Endosense (PI)‡	■ Acutus	None	4.1, 4.2.2, 4.2.3, 5.3, 5.4, 5.5.1, 5.6, 6, 7, 8, 9 (expect 9.7), 10 (except 10.3), 13, 15
Anne B. Curtis	University at Buffalo—SUNY Distinguished Professor; Charles and Mary Bauer Professor and Chair		None	None	None	None	None	4.1, 4.2.2, 4.2.3, 5.1.1, 5.1.2, 5.1.3, 5.1.4, 5.2, 5.4, 5.6, 6, 7, 8, 9, 10, 12, 13, 15
Barbara J. Deal	Getz Professor of Cardiology Feinberg School of Medicine Northwestern University	None	None	None	None	None	None	None
Timm Dickfeld	University of Maryland—Associate Professor of Medicine	■ Biosense ■ St. Jude Medical ■ Siemens	None	None	■ Biosense† ■ General Electric† ■ Siemens†	■ Impulse Dynamics‡ ■ Siemens†	None	4.1, 4.2 (except 4.2.6), 4.3, 5.3, 5.4, 5.6, 6, 7, 8, 9 (except 9.7), 10.1, 11, 13, 15
Anne M. Gillis	University of Calgary—Professor of Medicine	None	None	None	■ Medtronic	None	None	4.2, 5.2.2, 5.3.2, 6.4.1, 6.4.2, 6.4.4, 6.5, 6.7, 7, 8, 9, 10, 11 (except 11.7), 13, 15

continued on the next page

**APPENDIX 1. CONTINUED**

Committee Member	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness	Voting Recusals by Section*
Christopher B. Granger	Duke Clinical Research Institute; Duke University—Professor of Medicine; Director, Cardiac Care Unit	■ AstraZeneca† ■ Gilead Sciences† ■ GlaxoSmithKline† ■ Janssen Pharmaceuticals† ■ Medtronic† ■ Pfizer† ■ Sanofi-aventis†	None	None	■ AstraZeneca† ■ GlaxoSmithKline ■ Janssen Pharmaceuticals† ■ Medtronic† ■ Pfizer ■ Sanofi-aventis†	■ GE Healthcare† ■ Medtronic† ■ ZOLL Medical† ■ Spacelabs† ■ Phillips†	None	4, 5.1 (except 5.1.5), 5.2, 5.3, 5.4, 5.6, 6, 7, 8, 9, 12, 13, 15
Mark A. Hlatky	Stanford University School of Medicine—Professor of Health and Research Policy, and of Cardiovascular Medicine	None	None	None	None	None	None	None
Stephen C. Hammill	Mayo Clinic—Professor Emeritus of Medicine	None	None	None	None	None	None	None
José A. Joglar	UT Southwestern Medical Center—Professor of Internal Medicine; Clinical Cardiac EP—Fellowship Program Director	None	None	None	None	None	None	None
G. Neal Kay	University of Alabama at Birmingham—Professor Emeritus	None	None	None	None	None	None	None
Michael E. Field	University of Wisconsin School of Medicine and Public Health—Director, Clinical EP and Cardiac Arrhythmia Service, Associate Professor of Medicine	None	None	None	None	None	None	None
Gregg C. Fonarow	Ahmanson-UCLA Cardiomyopathy Center—Director; UCLA Division of Cardiology—Co-Chief	■ Amgen ■ Janssen Pharmaceuticals ■ Medtronic ■ ZS Pharma	None	None	■ Medtronic—IMPROVE-HF (Steering Committee)‡ ■ Medtronic†	None	None	4.1, 4.2.2, 4.2.3, 5.1 (except 5.1.5.1), 5.2, 5.3, 5.4, 5.6, 6, 7, 8, 9, 10, 12, 13, 15
Daniel D. Matlock	University of Colorado School of Medicine—Associate Professor of Medicine	None	None	None	None	None	None	None
Robert J. Myerburg	University of Miami Miller School of Medicine—Professor of Medicine and Physiology	None	None	None	None	None	None	None
Richard L. Page	University of Wisconsin Hospital and Clinics—Chair, Department of Medicine	None	None	None	None	None	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 relationships were reviewed and updated in conjunction with all meetings and/or conference calls of the writing committee during the document development process. The table does not necessarily reflect relationships with industry at the time of publication. A person is deemed to have a significant interest in a business if the interest represents ownership of ≥5% of the voting stock or share of the business entity, or ownership of ≥\$5,000 of the fair market value of the business entity; or if funds received by the person from the business entity exceed 5% of the person's gross income for the previous year. Relationships that exist with no financial benefit are also included for the purpose of transparency. Relationships in this table are modest unless otherwise noted.

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

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

†Significant relationship.

‡No financial benefit.

ACC indicates American College of Cardiology; AHA, American Heart Association; EP, Electrophysiology; HRS, Heart Rhythm Society; IMPROVE-HF, Improve the Use of Evidence-Based Heart Failure Therapies in the Outpatient Setting; PI, principle investigator; SUNY, State University of New York; and UT, University of Texas.

**APPENDIX 2. REVIEWER RELATIONSHIPS WITH INDUSTRY AND OTHER ENTITIES (COMPREHENSIVE)—2017 AHA/ACC/HRS GUIDELINE FOR MANAGEMENT OF PATIENTS WITH VENTRICULAR ARRHYTHMIAS AND THE PREVENTION OF SUDDEN CARDIAC DEATH (JULY 2017)**

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Salary	Expert Witness
Alfred E. Buxton	Content Reviewer	Professor of Medicine—Harvard Medical School—Beth Israel Deaconess Medical Center	None	None	None	■ NHLBI (DSMB)†	■ Medtronic† ■ Biosense Webster†	None	None
Andrew E. Epstein	Content Reviewer	Professor of Medicine—Cardiovascular Division University of Pennsylvania—Chief of Cardiology Section—Philadelphia VA Medical Center	■ Zoll*	None	None	■ Biotronik* ■ Boston Scientific* ■ Boston Scientific (DSMB)* ■ Medtronic* ■ Medtronic (DSMB) ■ St Jude Medical/Abbott* ■ St Jude Medical/Abbott (DSMB)*	None	None	■ Defendant, Amiodarone pulmonary toxicity, 2016 ■ Defendant, Appropriateness of pacemaker implantation, 2016*
Brian Olshansky	Content Reviewer	Adjunct Professor of Medicine—Des Moines University—Professor Emeritus—University of Iowa	■ Boehringer Ingelheim ■ Lundbeck Inc* ■ On-X/Cryolife	■ Lundbeck Inc* ■ On-X/Cryolife	None	■ Amarin (DSMB)*	None	None	■ Plaintiff, Long QT sudden death, 2017
Bulent Gorenек	Content Reviewer—ACC EP Council		None	None	None	None	None	None	None
Charles I. Berul	Content Reviewer	Division Chief of Pediatric Cardiology—Children's National Medical Center	None	None	None	None	■ Circulation*	None	None
Darren Sudman	Content Reviewer	Executive Director—Simon's Fund	None	None	None	None	None	None	None
George J. Klein	Content Reviewer	Chief of Cardiology—London Health Sciences Center	■ Biotronik ■ Boston Scientific ■ Medtronic*	None	None	None	None	None	None
Glenn N. Levine	Content Reviewer—ACC/AHA Task Force on Clinical Practice Guidelines	Professor of Medicine—Baylor College of Medicine—Director—Cardiac Care Unit—Michael E. DeBakey Medical Center	None	None	None	None	None	None	■ Defendant, Catheterization Laboratory Procedure, 2016 ■ Defendant, Out of hospital death, 2016
Gurushen S. Panjwani	Content Reviewer—ACC Heart Failure and Transplant Council	Director Heart Failure and Mechanical Support Program—George Washington University	■ Amgen Inc.*	None	None	None	■ BEAT-HF‡ ■ ENDEAVOUR‡	None	None

Continued on the next page

**APPENDIX 2. CONTINUED**

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Salary	Expert Witness
James P. Daubert	Official Reviewer—AHA	Duke University Medical Center	■ Biosense Webster ■ Boston Scientific ■ CardioFocus ■ Gilead ■ Heart Metabolics ■ Medtronic* ■ St. Jude Medical ■ Zoll	None	None	■ ARCA biopharma ■ Biosense Webster* ■ Boston Scientific* ■ Gilead* ■ Gilead (DSMB) ■ Medtronic* ■ NHLBI* ■ NHLBI (DSMB) ■ Northwestern University ■ St. Jude Medical (DSMB) ■ VytronUS (DSMB)	■ Biosense* ■ Biotronik* ■ Boston Scientific* ■ Gilead Sciences, Inc.* ■ Medtronic* ■ St. Jude Medical*	■ ACC	None
James Tisdale	Content Reviewer—ACC EP Council	Professor—College of Pharmacy Purdue University—Adjunct Professor—School of Medicine Indiana University	None	None	None	■ AHA* ■ HRS* ■ Indiana Clinical Translational Sciences Institute/Strategic Research Initiative*	■ ACC† ■ AHA† ■ AZCert† ■ QT drugs list, credible meds.org†	None	■ Plaintiff, Drug-induced torsades de pointes, 2017*
John L. Sapp	Official Reviewer—HRS	Interim Head—Division of Cardiology QEII Health Sciences Centre—Professor of Medicine—Dalhousie University	■ Biosense Webster* ■ Medtronic ■ St. Jude	None	None	■ Biosense Webster* ■ Canadian Institute of Health Research* ■ DSMB† ■ Phillips healthcare* ■ St. Jude Medical*	■ ARTESIA‡ ■ Medtronic‡ ■ Optisure Registry‡ ■ St. Jude‡	None	None
Joseph Edward Marine	Official Reviewer—ACC	Associate Professor of Medicine—Johns Hopkins University School of Medicine	None	None	None	None	■ UpToDate	None	None
Kathleen T. Hickey	Official Reviewer—AHA	Professor of Nursing—Columbia University Medical Center	None	None	None	None	None	None	None
Kenneth A. Ellenbogen	Content Reviewer	Chief of Cardiology—Virginia Commonwealth University Medical Center	■ AHA ■ AtriCure* ■ Biosense Webster* ■ Biotronik* ■ Boston Science* ■ Capricor ■ HRS ■ Janssen ■ Medtronic* ■ Pfizer* ■ Sentra heart ■ St. Jude Medical*	None	None	■ AtriCure* ■ Biosense Webster* ■ Boston Science* ■ Daiichi Sankyo ■ Medtronic* ■ Medtronic (DSMB)* ■ NIH* ■ Pfizer*	■ Biosense Webster* ■ Boston Science* ■ Circulation† ■ Heart Rhythm† ■ JACC† ■ Medtronic* ■ PACE† ■ Sanofi Aventis	None	None

*Continued on the next page*

## APPENDIX 2. CONTINUED

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Salary	Expert Witness
Kim K. Birtcher	Content Reviewer—ACC/AHA Task Force on Clinical Practice Guidelines	University of Houston—College of Pharmacology	■ Jones and Bartlett Learning	None	None	None	■ Accreditation Council for Clinical Lipidology	■ University of Houston College of Pharmacology* ■ Walgreens*	None
Kristen B. Campbell	Content Reviewer	Duke University Hospital	None	None	None	None	None	None	None
Kristen K. Patton	Content Reviewer	Professor of Medicine—University of Washington	None	None	None	None	■ ABIM ■ ACGME† ■ AHA† ■ FDA ■ HRS†	None	None
L. Brent Mitchell	Content Reviewer	Professor—Department of Cardiac Sciences—Libin Cardiovascular Institute of Alberta—University of Calgary—Alberta Health Services	■ Boehringer Ingelheim* ■ Forest Pharmaceuticals ■ Guidant Canada* ■ Medtronic Canada* ■ Medtronic Inc* ■ Merck ■ Pfizer* ■ Servier Canada*	None	None	■ Boston Scientific*	■ ARTESIA‡ ■ Health Protection Branch, Government of Canada	None	None
Martin Borggrefe	Content Reviewer	I Medizinische KlinikKlinikum Mannheim GmbHUniversitätsklinikum	■ Bayer Health Care ■ Boehringer Ingelheim ■ Impulse Dynamics ■ Sanofi Aventis ■ St. Jude Medical	None	None	■ German Centre for Cardiovascular Research*	None	None	None
Mathew D. Hutchinson	Official Reviewer—HRS	Professor of Medicine—University of Arizona College of Medicine—Tucson	■ St. Jude Medical	None	None	None	None	None	None
Matthew W. Martinez	Content Reviewer—Sports and Exercise EP Council	Lehigh Valley Health Network	None	None	None	None	None	None	None
Melissa R. Robinson	Content Reviewer	Director—Complex Ablation Program—University of Washington	■ Medtronic* ■ Abbott* ■ Boston Scientific*	None	None	None	None	None	None

Continued on the next page

**APPENDIX 2. CONTINUED**

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Salary	Expert Witness
Michael J. Silka	Content Reviewer	Children's Hospital Los Angeles	None	None	None	None	None	None	■ Defendant, ICD implantation, 2017
Miguel A. Quinones	Content Reviewer	Methodist DeBakey Heart and Vascular Center	None	None	None	None	■ Houston Methodist Hospital*	None	None
Mitchell T. Saltzberg	Organizational Reviewer—HFSA	Jefferson Medical College—Christiania Care Health System	None	None	■ Nephroceuticals* ■ Stem Cell Theranostics*	None	None	None	None
N.A. Mark Estes III	Content Reviewer	Professor of Medicine—Tufts University School of Medicine	■ Boston Scientific* ■ Medtronic* ■ St. Jude Medical*	None	None	■ Boston Scientific* ■ International Board of Heart Rhythm Examiners† ■ Medtronic* ■ St. Jude Medical*	None	None	None
Norma M. Keller	Official Reviewer—ACC	New York University Medical Center	None	None	None	None	None	None	None
Peter Leong-Sit	Content Reviewer—HRS	Associate Professor of Medicine—Western University—London Health Sciences Centre	■ Medtronic Canada	■ Bayer Healthcare Pharmaceuticals ■ Biosense Webster ■ Johnson and Johnson	None	None	None	■ Bayer Healthcare Pharmaceuticals*	None
Rachel J. Lampert	Content Reviewer	Yale University School of Medicine—Section of Cardiology	■ Medtronic*	None	None	■ Boston Scientific* ■ GE Medical* ■ Medtronic, Inc.* ■ St. Jude Medical*	None	None	None
Sami Viskin	Content Reviewer	Tel Aviv Medical Center—Department of Cardiology	■ Boston Scientific European Strategy Advisory Board	None	None	None	None	None	None
Samuel S. Gidding	Content Reviewer—ACC/AHA Task Force on Clinical Practice Guidelines	Dupont Hospital for Children—Nemours Cardiac Center	■ Familial Hypercholesterolemia Foundation† ■ Regenxbio	None	None	■ Familial Hypercholesterolemia Foundation† ■ NIH Grants*	■ Cardiology Division Hea †No financial benefit.	None	None

*Continued on the next page*

## APPENDIX 2. CONTINUED

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Salary	Expert Witness
Silvia G. Priori	Content Reviewer	Professore Ordinario di Cardiologia—Università di Pavia—Direttore Scientifico—Istituti Clinici Scientifici Maugeri—Pavia, Italia	<ul style="list-style-type: none"> <li>■ Ambry Genetics</li> <li>■ Boston Scientific</li> <li>■ Medtronic</li> <li>■ Medtronic, Inc.</li> </ul>	None	<ul style="list-style-type: none"> <li>■ Audentes Therapeutics Inc*</li> </ul>	<ul style="list-style-type: none"> <li>■ Gilead Sciences*</li> </ul>	<p>#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 ACC/AHA Disclosure Policy for Writing Committees.<sup>d†</sup></p> <ul style="list-style-type: none"> <li>■ HRS</li> <li>■ GS-US-372-1234‡</li> </ul>	None	None
Susan Strong	Official Reviewer—AHA	Sabin Middle School	None	None	None	None	None	None	None
Win-Kuang Shen	Content Reviewer	Professor of Medicine—Consultant—Mayo Clinic Arizona, Phoenix Campus	None	None	None	None	None	None	None
Zachary D. Goldberger	Official Reviewer—ACC/AHA Task Force on Clinical Practice Guidelines Lead Reviewer	Assistant Professor of Medicine—Division of Cardiology—Harborview Medical Center—University of Washington School of Medicine	<ul style="list-style-type: none"> <li>■ RubiconMD</li> </ul>	None	None	None	None	None	None

This table represents the relationships of reviewers with industry and other entities that were disclosed at the time of peer review, including those not deemed to be relevant to this document, at the time this document was under review. The table does not necessarily reflect relationships with industry at the time of publication. A person is deemed to have a significant interest in a business if the interest represents ownership of  $\geq 5\%$  of the voting stock or share of the business entity, or ownership of  $\geq \$5000$  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. Names are listed in alphabetical order within each category of review. Please refer to <http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/relationships-with-industry-policy> for definitions of disclosure categories or additional information about the ACC/AHA Disclosure Policy for Writing Committees.

\*Significant relationship.

ABIM indicates American Board of Internal Medicine; ACC, American College of Cardiology; ACGME, Accreditation Council for Graduate Medical Education; AHA, American Heart Association; ARTESiA, Apixaban for the Reduction of Thrombo-Embolism in Patients With Device-Detected Sub-Clinical Atrial Fibrillation; BEAT-HF, Better Effectiveness After Transition-Heart Failure DSMB, data safety monitoring board; ENDEAVOUR, carfilzomib and dexamethasone versus bortezomib and dexamethasone for