

2015 HRS/EHRA/APHRS/SOLAECE Expert Consensus Statement on Optimal Implantable Cardioverter-Defibrillator Programming and Testing

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64 Keywords: Implantable cardioverter-defibrillator, Bradycardia mode and rate, Tachycardia detection,
65 Tachycardia therapy, Defibrillation Testing, Programming.

66 Abbreviations: ICD=Implantable Cardioverter-Defibrillator, DT=Defibrillation Testing, LV=Left Ventricular,
67 RV=Right Ventricular, LVEF=Left Ventricular Ejection Fraction, RCT=Randomized Clinical Trial,
68 CRT=Cardiac Resynchronization Therapy, CRT-D=Cardiac Resynchronization Therapy Defibrillator, ATP=
69 Antitachycardia Pacing, SVT=Supraventricular Tachycardia, VT=Ventricular Tachycardia, EGM=
70 Electrogram, AF=Atrial Fibrillation, S-ICD=Subcutaneous Implantable Cardioverter Defibrillator, PVC=
71 Premature Ventricular Contraction, NYHA=New York Heart Association, NCDR=National Cardiovascular
72 Data Registry.

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74 **Introduction**

75 Implantable cardioverter-defibrillator (ICD) therapy is clearly an effective therapy for selected
76 patients in definable populations. The benefits and risks of ICD therapy are directly impacted by
77 programming and surgical decisions. This flexibility is both a great strength and a weakness, for which
78 there has been no prior official discussion or guidance. It is the consensus of the 4 continental
79 electrophysiology societies that there are 4 important clinical issues for which there are sufficient ICD
80 clinical and trial data to provide evidence-based expert guidance. This document systematically
81 describes the greater than 80% (83%-100%, mean: 96%) required consensus achieved for each
82 recommendation by official balloting in regard to the programming of (1) bradycardia mode and rate, (2)
83 tachycardia detection, (3) tachycardia therapy, and (4) the intraprocedural testing of defibrillation
84 efficacy. Representatives nominated by the Heart Rhythm Society (HRS), European Heart Rhythm
85 Association (EHRA), Asian Pacific Heart Rhythm Society (APHRS) and the *Sociedad Latinoamericana de*
86 *Estimulacion Cardiaca y Electrofisiologia* (SOLAECE-Latin American Society of Cardiac Pacing and
87 Electrophysiology) participated in the project definition, the literature review, the recommendation
88 development, the writing of the document, and its approval. The 32 recommendations were balloted by
89 the 35 writing committee members and were approved by an average of 96%.

90 The classification of the recommendations and the level of evidence follow the recently updated
91 ACC/AHA standard.^{1,2} Class I is a strong recommendation, denoting a benefit greatly exceeding risk.
92 Class IIa is a somewhat weaker recommendation, with a benefit probably exceeding risk, and Class IIb
93 denotes a benefit equivalent to or possibly exceeding risk. Class III is a recommendation against a

94 specific treatment because there is either no net benefit or there is net harm. Level of Evidence A
95 denotes the highest level of evidence from more than 1 high-quality randomized clinical trial (RCT), a
96 meta-analysis of high quality RCTs, or RCTs corroborated by high-quality registry studies. Level of
97 evidence B indicates moderate-quality evidence from either RCTs with a meta-analysis (B-R) or well-
98 executed nonrandomized trials with a meta-analysis (B-NR). Level of evidence C indicates randomized or
99 nonrandomized observational or registry studies with limited data (C-LD) or from expert opinions (C-EO)
100 based on clinical experience in the absence of credible published evidence. These recommendations
101 were also subject to a 1-month public comment period. Each society then officially reviewed,
102 commented, edited, and endorsed the final document and recommendations. All author and peer
103 reviewer disclosure information is provided in Appendix A.

104 The care of individual patients must be provided in context of their specific clinical condition and the
105 data available on that patient. Although the recommendations in this document provide guidance for a
106 strategic approach to ICD programming, as an individual patient's condition changes or progresses and
107 additional clinical considerations become apparent, the programming of their ICDs must reflect those
108 changes. Remote and in-person interrogations of the ICD and clinical monitoring must continue to
109 inform the programming choices made for each patient. The recommendations in this document
110 specifically target adult patients and might not be applicable to pediatric patients, particularly when
111 programming rate criteria.

112 Please consider that each ICD has specific programmable options that might not be specifically
113 addressed by the 32 distinctive recommendations in this document. Appendix B, published online
114 (<http://www.hrsonline.org/appendix-b>), contains the writing committee's translations specific to each
115 manufacturer and is intended to best approximate the recommended behaviors for each available ICD
116 model.

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122 Bradycardia Mode and Rate Programming**123 Single- or Dual-Chamber Pacing Mode**

124 *Evidence:* Because the ICD is primarily indicated for tachycardia therapy, there might be some
125 uncertainty regarding optimal bradycardia management for ICD patients. Data from clinical studies
126 adequately address only the programmed mode rather than the number of leads implanted, the number
127 of chambers stimulated, or how frequently the patients required bradycardia support. It is of note that
128 most information on pacing modes has been collected from pacemaker patients, and these patients are
129 clinically distinct from ICD recipients. Dual-chamber pacing (atrial and ventricular) has been compared
130 with single-chamber pacing (atrial or ventricular) in patients with bradycardia in 5 multi-center, parallel,
131 randomized trials, in 1 meta-analysis of randomized trials, and in 1 systematic review that also included
132 30 randomized crossover comparisons and 4 economic analyses.³⁻⁹ Meta analyses comparing dual-
133 chamber to single-chamber ICDs did not evaluate pacing modes.^{10,11} Compared with single-chamber
134 pacing, dual chamber pacing results in small but potentially significant benefits in patients with sinus
135 node disease and/or atrioventricular block. No difference in mortality has been observed between
136 ventricular pacing modes and dual-chamber pacing modes. Dual-chamber pacing was associated with a
137 lower rate of atrial fibrillation (AF) and stroke.¹² The benefit in terms of AF prevention was more marked
138 in trials comprised of patients with sinus node disease. Although trends in favor of dual-chamber pacing
139 have been observed in some trials, there was no benefit in terms of heart failure (HF). In patients
140 without symptomatic bradycardia, however, the Dual Chamber and VVI Implantable Defibrillator
141 (DAVID) trial in ICD recipients showed that one specific choice of dual-chamber rate-responsive (DDDR)
142 programming parameters led to poorer outcomes than VVI backup pacing, most likely secondary to
143 unnecessary right ventricular (RV) pacing. The fact that RV stimulation was responsible was reinforced in
144 the DAVID II trial, in which AAI pacing was demonstrated to be noninferior to VVI backup pacing.¹³

145 Approximately a quarter of patients with either sinus node disease or atrioventricular block develop
146 “pacemaker syndrome” with VVI pacing usually associated with retrograde (ventricular to atrial)
147 conduction, which in turn is associated with a reduction in the quality of life.¹⁴ In crossover trials,
148 symptoms of pacemaker syndrome (dyspnea, dizziness, palpitations, pulsations, and chest pain) were
149 reduced by reprogramming to a dual-chamber mode.¹⁴ Dual-chamber pacing is associated with better
150 exercise performance compared with single-chamber VVI pacing without rate adaptation, but produces
151 similar exercise performance when compared with rate-responsive VVIR pacing. Because of the
152 additional lead, dual-chamber devices involve longer implantation times, have a higher risk of

153 complications, and are more expensive. However, because of the additional clinical consequences of
154 pacemaker syndrome and AF (and its sequelae), the overall cost difference between single- and dual-
155 pacing systems is moderated.

156 In patients with persistent sinus bradycardia, atrial rather than ventricular dual-chamber pacing is the
157 pacing mode of choice. There is evidence for superiority of atrial-based pacing over ventricular pacing
158 for patients who require pacing for a significant proportion of the day. The evidence is stronger for
159 patients with sinus node disease, in whom dual-chamber pacing confers a modest reduction in AF and
160 stroke, but not in hospitalization for HF or death compared with ventricular pacing. In patients with
161 acquired atrioventricular block, large randomized parallel trials were unable to demonstrate the
162 superiority of dual-chamber pacing over ventricular pacing with regard to hard clinical endpoints of
163 mortality and morbidity.^{4,6-8} The benefit of dual-chamber over ventricular pacing is primarily due to the
164 avoidance of pacemaker syndrome and to improved exercise capacity.¹⁴ Even if it is a softer endpoint,
165 pacemaker syndrome is associated with a reduction in quality of life that justifies the preference for
166 dual-chamber pacing when reasonable; thus, there is strong evidence for the superiority of dual-
167 chamber pacing over ventricular pacing that is limited to symptom improvement. Conversely, there is
168 strong evidence of nonsuperiority with regard to survival and morbidity. The net result is that the
169 indications for programming the dual-chamber modes are weaker and the choice regarding the pacing
170 mode should be individualized, taking into consideration the increased complication risk and costs of
171 dual-chamber devices. Because ICD patients usually do not require bradycardia support, with the
172 exception of patients who require cardiac resynchronization, programming choices should avoid pacing
173 and in particular avoid single ventricular pacing, if possible.^{15,16}

174 **Programming of Rate Modulation**

175 The benefit of rate response programming has been evaluated in patients with bradycardia in 5 multi-
176 center, randomized trials and in 1 systematic review that also included 7 single-center studies.¹⁷⁻²² Most
177 of these data were obtained from pacemaker studies and must be interpreted in that light.

178 Although there is evidence of the superiority of VVIR pacing compared with VVI pacing in improving
179 quality of life and exercise capacity, improvements in exercise capacity with DDDR compared with DDD
180 have been inconsistent. In 2 small studies on patients with chronotropic incompetence comparing DDD
181 and DDDR pacing, the latter had improved quality of life and exercise capacity; however, a larger,
182 multicenter randomized trial (Advanced Elements of Pacing Randomized Controlled Trial [ADEPT]) failed

183 to show a difference in patients with a modest blunted heart rate response to exercise.¹⁷⁻¹⁹ In addition,
184 DDDR programming in CRT patients has the potential to impair AV synchrony and timing. It should be
185 noted that trials evaluating CRT generally did not use rate responsive pacing, and many in fact avoided
186 atrial stimulation using atrial sensed and ventricular paced pacing modes with a lower base rate.
187 However, the Pacing Evaluation-Atrial Support Study in Cardiac Resynchronization Therapy (PEGASUS
188 CRT) trial is the exception and did not demonstrate adverse impact on mortality and HF events.²³

189

190 **Sinus Node Disease**

191 In patients with persistent or intermittent sinus node dysfunction or chronotropic incompetence, the
192 first choice is DDDR with algorithms responding to intermittent atrioventricular conduction. There is
193 sufficient evidence for the superiority of VVIR compared with VVI in improving quality of life and
194 exercise capacity. The evidence is much weaker in dual-chamber pacing (DDDR vs DDD).

195 Although only an issue when there is some concomitant AV block, the upper rate limit should be
196 programmed higher than the fastest spontaneous sinus rhythm to avoid upper rate limit behavior. To
197 avoid symptomatic bradycardia, the lower rate should be programmed on an individual basis, according
198 to the clinical characteristics and the underlying cardiac substrate of the patient.

199 **Atrial Fibrillation and Atrioventricular Block**

200 Patients with permanent atrial fibrillation and either spontaneous or AV junctional ablation-induced
201 high-degree atrioventricular block have little to no chronotropic response to exercise, thus VVIR pacing
202 is associated with better exercise performance, improved daily activities, improved quality of life, and
203 decreased symptoms of shortness of breath, chest pain, and heart palpitations, compared with VVI.²⁰⁻
204 ^{22,24-26} Therefore, rate-adaptive pacing is the first choice of pacing mode; fixed-rate VVI pacing should be
205 abandoned in patients with permanent AF and atrioventricular block. It is the experts' opinion that the
206 minimum rate can be programmed higher (e.g., 70 bpm) than for sinus rhythm patients, in an attempt
207 to compensate for the loss of active atrial filling. In addition, the maximum sensor rate should be
208 programmed restrictively (e.g., 110–120 bpm) to avoid "overpacing" (i.e. pacing with a heart rate faster
209 than necessary), which can be symptomatic, particularly in patients with coronary artery disease. In a
210 small study, however, it was found that rate-responsive pacing could be safe and effective in patients
211 with angina pectoris, without an increase in subjective or objective signs of ischemia.²⁵ The lower rate

212 should be programmed on an individual basis, according to the clinical characteristics and the underlying
213 cardiac substrate of the patient. The clinical benefit of programming a lower resting rate at night based
214 on internal clocks has not been evaluated in ICD patients. There is some concern that atrioventricular
215 junction ablation and permanent ventricular pacing might predispose the patient to an increased risk of
216 sudden cardiac death related to a bradycardia-dependent prolongation of the QT interval. This risk
217 might be overcome by setting the ventricular pacing rate to a minimum of 80 or 90 bpm for the first 1–2
218 months following the atrioventricular junction ablation, then reducing it to a conventional 60–70
219 bpm.^{27,28} Not all patients with AF and milder forms of atrioventricular block will require a high
220 percentage of ventricular pacing or have a wide QRS. Physicians should consider the risk of increasing
221 pre-existing left ventricular dysfunction with RV pacing versus improved chronotropic responsiveness
222 and the potential value of cardiac resynchronization therapy.

223

224 **Intact Atrioventricular Conduction**

225 **Right-Ventricular Pacing**

226 The results of a number of large-scale, prospective randomized trials demonstrated a significant
227 reduction in AF in pacemaker patients with atrial-based pacing (AAI or DDD) compared with patients
228 with ventricular-based pacing.^{4,8,29} In the Mode Selection Trial, which enrolled 2010 patients with sick
229 sinus syndrome, the risk of AF increased linearly with the increasing percentage of RV pacing.³⁰ At the
230 same time, deleterious effects of right ventricular (RV) pacing in patients with left ventricular (LV)
231 dysfunction (LVEF \leq 40%) implanted with dual-chamber ICD systems were observed in the Dual Chamber
232 and VVI Implantable Defibrillator (DAVID) trial, which included 506 ICD patients without indications for
233 bradycardia pacing. Patients within the DDDR-70 group (with paced and sensed atrioventricular delays
234 of 170 and 150 ms, respectively, in most of the DDDR group patients) showed a trend toward higher
235 mortality and an increased incidence of HF compared with the patients programmed to ventricular
236 backup pacing—the VVI-40 group. Within the DDDR-70 group, there were more cardiac events when the
237 percentage of ventricular pacing exceeded 40% ($P = .09$) compared with patients with <40% of RV
238 pacing, although almost all the patients had >95% RV stimulation (DDDR-70) or <5% RV stimulation (VVI-
239 40).^{31,32} However, a more detailed post-hoc analysis of the Inhibition of Unnecessary RV Pacing With
240 Atrial-Ventricular Search Hysteresis in ICDs (INTRINSIC RV) trial revealed that the most favorable clinical
241 results were not in the VVI groups with the least percentage of RV pacing, but in the subgroup that had

242 DDD pacing with longer atrioventricular delays and 11%–19% of ventricular pacing. This parameter
 243 selection probably helped patients to avoid exceedingly low heart rates while preserving intrinsic
 244 atrioventricular conduction most of the time.^{31,33} In the Second Multicenter Automated Defibrillator
 245 Implantation Trial (MADIT II), a higher risk of HF was observed in patients who had a greater than 50%
 246 burden of RV pacing.³⁴ In another large observational study of 456 ICD patients without HF at baseline, a
 247 high RV pacing burden (RV pacing more than 50% of the time) was associated with an increased risk of
 248 HF events and appropriate ICD shocks.³⁵ Optimally, right ventricular stimulation should be avoided but
 249 the precise tradeoff between the percentage of ventricular pacing and atrioventricular timing is unclear
 250 in non-CRT patients.

251 **Non-CRT Devices: Algorithms to Reduce Right Ventricular Stimulation**

252 The importance of reducing or avoiding right ventricular pacing in ICD patients with left ventricular
 253 dysfunction was illustrated in the DAVID trial.³¹ The feasibility of algorithms designed to decrease the
 254 burden of unnecessary ventricular pacing has been demonstrated in patients with dual-chamber
 255 pacemakers.³⁶⁻³⁸ These algorithms usually provide functional AAI pacing with monitoring of
 256 atrioventricular conduction and an automatic mode switch from AAI to DDD during episodes of
 257 atrioventricular block. Some studies directly compared various algorithms to decrease ventricular
 258 pacing, showing that a “managed ventricular pacing” (MVP) algorithm resulted in greater ventricular
 259 pacing reduction than an “atrioventricular search” algorithm,^{39,40} however, no randomized studies
 260 comparing these two algorithms with respect to important cardiovascular endpoints (e.g., HF, cardiac
 261 death) have been performed. The results of the studies on these pacing algorithms are summarized in
 262 Table 1.

263 **Table 1.** Influence of pacing modes and algorithms on clinical endpoints.

Study	Patients (PM/ICD)	Results and remarks
SAVE PACe, randomized multicenter (2007) ⁴¹	1065 (PM)	40% relative risk reduction of AF in the MVP group compared with DDD pacing (4.8% absolute risk reduction).
MVP, randomized multicenter (2011) ⁴²	1030 (ICD)	No superiority of MVP over VVI-40 in terms of AF, VT/VF, quality of life, HF.
Steinbach et al, retrospective single-	102 (PM)	In patients over 75 years of age, MVP showed lower rates of HF episodes and all-cause mortality than conventional

center (2011) ⁴³		DDD pacing
long-MinVPACE, randomized single-center (2011) ⁴⁴	66 (PM)	Less RV pacing, less AF burden in MinVP group patients compared with DDDR (mean 12.8 vs 47.6%). Chosen AV/PV delay (150/130 ms) was probably too short in the DDDR (control) group.
Generation MVP, observational multicenter (2012) ⁴⁵	220 (PM)	Significantly fewer atrial arrhythmias when programmed to MVP compared with DDD.
PreFER MVP, randomized multicenter (2014) ⁴⁶	605 (556 PM, 49 ICD)	No difference between cardiovascular hospitalization, AF, and the composite of death and hospitalization between the MVP and DDD groups. The authors stated that “patients were enrolled upon elective replacement of the device, and were healthy enough to survive the first device without experiencing a significant decrease in LV function.”
MINERVA, randomized multicenter (2014) ⁴⁷	1300 (PM)	AF burden: no superiority of MVP pacing compared with the DDDR mode (AV/PV delay >180/210 ms in greater than 60% of patients, 53% of RV pacing). MVP in combination with atrial antitachycardia pacing was superior to both DDDR and MVP-only.
COMPARE, randomized multicenter (2014) ⁴⁸	385 (PM)	Lower percentage of ventricular pacing (%VP) in the MVP group compared with the SearchAV+ group. A trend in the correlation between %VP and AT/AF burden.

264 AT=atrial tachycardia, PM=pacemaker, HF=heart failure, MVP=Managed Ventricular Pacing

265 Unnecessary RV pacing should be minimized by using specific algorithms or programming longer
 266 atrioventricular delays, and this process is more important for patients with a higher risk of AF or who
 267 already have poorer LV function.⁴⁹ Patients with longer baseline PR intervals have a higher risk of AF
 268 regardless of the percentage of ventricular pacing or the length of the programmed atrioventricular
 269 interval.⁵⁰ Use of the AAIR pacing mode with exceedingly long atrioventricular conduction times can lead
 270 to “AAIR pacemaker syndrome” and actually increases the risk of AF compared with the DDDR mode, as
 271 was shown in the Danish Multicenter Randomized Trial on Single Lead Atrial versus Dual-Chamber
 272 Pacing in Sick Sinus Syndrome (DANPACE).^{3,51} Therefore, excessively long atrioventricular delays

273 resulting in non-physiologic atrioventricular contraction patterns should be avoided. The potential harm
274 of atrial pacing with a prolonged atrioventricular delay was also demonstrated in the MVP trial. In the
275 MVP trial, dual-chamber pacing with the MVP algorithm was not superior to ventricular backup pacing
276 (VVI 40 bpm) with respect to HF events. After a follow-up of 2.4 years, there was an apparent increase in
277 HF events that was limited primarily to patients with a baseline PR interval of >230 ms (mean PR of 255
278 to 260 ms).⁴² Long atrioventricular intervals also predispose the patient to repetitive atrioventricular
279 reentrant rhythms, “repetitive nonreentrant VA synchrony,” or “atrioventricular desynchronization
280 arrhythmia,” which manifest as mode switching but which also cause sustained episodes with poor
281 hemodynamics.⁵² Thus, based on the available data, it appears that atrial pacing with excessively long
282 atrioventricular delays should be avoided.

283 Algorithms that minimize ventricular pacing sometimes lead to inadvertent bradycardia or spontaneous
284 premature, beat-related short-long-short RR interval sequences with proarrhythmic potential.⁵³⁻⁵⁵
285 However, in a study retrospectively analyzing the onset of ventricular tachycardia (VT) in ICD patients,
286 the MVP mode was less frequently associated with the onset of VT compared with the DDD and VVI
287 modes.⁵⁴ Atrioventricular decoupling (greater than 40% of atrioventricular intervals exceeding 300 ms)
288 was observed in 14% of the ICD patients in the Marquis ICD MVP study, which might have a negative
289 effect on ventricular filling.⁵⁶

290 In ICD patients with structural heart disease, spontaneous atrioventricular conduction can become
291 prolonged instead of shortening, with increased atrial paced heart rates.³³ This outcome frequently
292 leads to a higher percentage of ventricular paced complexes. In view of the results of the ADEPT trial,
293 which failed to demonstrate the clinical superiority of combined rate modulation and DDD pacing, the
294 need for and aggressiveness of sensor-driven rate responses should be individualized or eliminated.¹⁹
295 Rate-dependent shortening of atrioventricular delay could have the same effect and should usually be
296 avoided.

297 Patients with hypertrophic cardiomyopathy represent a small but intricate subset of the ICD population
298 for whom pacing has not been demonstrated to be a consistently effective treatment for outflow tract
299 obstruction. However, according to the 2011 ACCF/AHA Hypertrophic Cardiomyopathy Guideline, dual-
300 chamber ICDs are reasonable for patients with resting LV outflow tract gradients more than 50 mm Hg,
301 and who have indications for ICD implantation to reduce mortality.⁵⁷ In these patients, atrioventricular
302 delays should be individually programmed to be short enough to achieve RV pre-excitation and decrease
303 LV outflow tract gradient, but not too short, which would impair LV filling; usually in the ranges of 60–

304 150 ms.^{58,59} There are few studies of pacing modes in these patients, and they are limited by small
305 numbers and the failure to quantify important cardiac outcomes.

306 In conclusion, atrioventricular interval programming and choosing between DDDR and MVP or other
307 atrioventricular interval management modes should be performed on an individual basis. The goal is to
308 minimize the percentage of RV pacing and to avoid atrial-based pacing with atrioventricular intervals
309 exceeding 250–300 ms leading to atrioventricular uncoupling. In patients with prolonged PR intervals
310 and impaired LV function, biventricular pacing can be considered.

311 **Cardiac Resynchronization Therapy: Consistent Delivery of Ventricular Pacing**

312 Cardiac resynchronization therapy in combination with a defibrillator device (CRT-D) improves survival
313 and cardiac function in patients with LV systolic dysfunction, prolonged QRS duration, and mild-to-
314 severe HF.⁶⁰⁻⁶² The beneficial effect of CRT-D compared with ICD is likely to be derived from biventricular
315 pacing, with a decrease in dyssynchrony and an improvement in cardiac function. The percentage of
316 biventricular pacing capture in the ventricles can be negatively influenced by a number of factors,
317 including atrial tachyarrhythmias, premature ventricular complexes, and programming of the
318 atrioventricular delay, giving way to the intrinsic conduction of the patient and a reduced percentage of
319 biventricular pacing. Some large observational studies have investigated the optimal level of
320 biventricular pacing percentage and found a higher percentage to be associated with more pronounced
321 CRT benefits. An optimal CRT benefit was observed with a biventricular pacing percentage as close to
322 100% as possible.⁶³⁻⁶⁶

323 In the analysis of the LBBB population in the MADIT-CRT trial, those patients with less than 90%
324 biventricular pacing had similar rates of HF and death compared with the patients randomized to no
325 CRT. By contrast, biventricular pacing exceeding 90% was associated with a benefit of CRT-D in terms of
326 HF or death compared with ICD patients and no CRT. Biventricular pacing 97% and greater was
327 associated with a further reduction in HF or death and a significant reduction in death alone.
328 Consistently, every 1% increase in biventricular pacing percentage was associated with a 6% risk
329 reduction in HF or death, a 10% risk reduction in death alone, and an increase in LV reverse
330 remodeling.⁶⁷ Therefore, in ICD patients with biventricular pacing, it can be beneficial to adjust the
331 therapy to produce the highest achievable percentage of ventricular pacing, preferably above 98%, to
332 improve survival and reduce HF hospitalization. Approaches to increasing the percentage of

333 biventricular pacing include programming shorter but hemodynamically appropriate atrioventricular
 334 delays and minimizing atrial and ventricular ectopic activity and tachyarrhythmias.

335 Optimizing the location of ventricular pacing sites and the timing of the pacing pulses can significantly
 336 improve cardiac hemodynamics in CRT patients. Echocardiographic optimization of atrioventricular
 337 delays in CRT patients can alleviate HF symptoms and increase exercise capacity compared with nominal
 338 programming, particularly when approaching nonresponding populations.⁶⁸ However,
 339 echocardiographic optimization in the PROSPECT study did not support this approach in a randomized
 340 trial and the Frequent Optimization Study Using the QuickOpt Method (FREEDOM) trials failed to
 341 provide evidence supporting the benefit of CRT optimization and did not demonstrate superiority of the
 342 respective algorithms over nominal or empiric programming.⁶⁹⁻⁷¹ There are limited data supporting the
 343 use of LV-only stimulation in a small subset of patients who fail to respond to biventricular stimulation.⁷²
 344 Adaptive CRT (aCRT) is an algorithm that periodically measures intrinsic conduction and dynamically
 345 adjusts CRT pacing parameters. The algorithm withholds right ventricular pacing when intrinsic electrical
 346 conduction to the right ventricle is normal and provides adjustment of CRT pacing parameters based on
 347 electrical conduction. A prospective, multicenter, randomized, double-blind clinical trial demonstrated
 348 the safety and efficacy of the aCRT algorithm.⁷³ This algorithm can increase the longevity of the
 349 implantable device and replace a manual device optimization process with an automatic ambulatory
 350 algorithm, although echo optimization might still be needed, at least in nonresponders. The Clinical
 351 Evaluation on Advanced Resynchronization (CLEAR) study assessed the effects of CRT with automatically
 352 optimized atrioventricular and interventricular delays, based on a Peak Endocardial Acceleration (PEA)
 353 signal system. PEA-based optimization of CRT in patients with HF significantly increased the proportion
 354 of patients who improved with therapy during follow-up, mainly through an improved New York Heart
 355 Association (NYHA) class.⁷⁴

Bradycardia Mode and Rate Programming Recommendations	Class of Recommendation	Level of Evidence
In ICD patients who also have sinus node disease and guideline-supported indications for a bradycardia pacemaker, it is beneficial to provide dual-chamber pacing to reduce the risk of AF and stroke, to avoid pacemaker syndrome, and to improve quality of life.	I	B-R
In single- or dual-chamber ICD patients without guideline-supported	I	B-R

indications for bradycardia pacing, adjusting the pacing parameters is recommended so that ventricular stimulation is minimized to improve survival and reduce HF hospitalization.		
In ICD patients who have sinus rhythm, no or only mild LV dysfunction, and atrioventricular block where ventricular pacing is expected, it is reasonable to provide dual-chamber pacing in preference to single-chamber ventricular pacing to avoid pacemaker syndrome and to improve quality of life.	Ila	B-R
In ICD patients who have sinus rhythm, mild to moderate LV dysfunction, and atrioventricular block where ventricular pacing is expected, it is reasonable to provide CRT in preference to dual-chamber ventricular pacing to improve the combination of HF hospitalization, LV enlargement, and death.	Ila	B-R
In ICD patients who have chronotropic incompetence, it can be beneficial to program the ICD to provide sensor-augmented rate response, especially if the patient is young and physically active.	Ila	B-NR
In dual-chamber ICD patients with native PR intervals of 230 ms or less, it can be beneficial to program the mode, automatic mode change, and rate response so the patient's native atrioventricular conduction minimizes ventricular pacing.	Ila	B-R
In biventricular pacing ICD patients, it can be beneficial to adjust the therapy to produce the highest achievable percentage of ventricular pacing, preferably above 98%, to improve survival and reduce HF hospitalization.	Ila	B-NR
In biventricular pacing ICD patients, it can be reasonable to activate the algorithms providing automatic adjustment of atrioventricular delay and/or LV-RV offset to obtain a high percentage of synchronized pacing and reduce the incidence of clinical events.	IIb	B-R

356

357

358 **Tachycardia Detection Programming**

359 Following significant technological changes in ICDs in recent years, the concept of optimal ICD
360 programming has changed dramatically. From the dawn of this therapy in the early 1980s to the first
361 decade of the 21st Century, the rapid detection and treatment of VT and VF have been stressed. The
362 argument for rapid detection of VT and VF derived from a number of factors. Initial skepticism regarding
363 the feasibility of sudden death prevention with ICDs, the fact that early ICD patients had all survived one
364 or more cardiac arrests, concern for undersensing and underdetection (of VF in particular),
365 demonstration of an increasing defibrillation threshold with prolonged VF duration, and the increased
366 energy requirement of monophasic defibrillation all created a culture of programming for rapid
367 tachycardia detection and the shortest possible time to initial therapy.⁷⁵⁻⁷⁷ The initial generations of ICDs
368 did not record and save electrograms, leading to a reduced appreciation for the frequency and impact of
369 inappropriate shocks. With the advent and then dominance of primary prevention indications, avoidable
370 shocks assumed a relatively larger proportion of total therapy.⁷⁸⁻⁸³ Gradually, publications have
371 increased awareness of the frequency and the diverse range of adverse outcomes associated with
372 avoidable ICD therapy, and have demonstrated that avoidable ICD shocks can be reduced by evidence-
373 based programming of the detection rate, detection duration, anti-tachycardia pacing (ATP), algorithms
374 that discriminate supraventricular tachycardia (SVT) from VT, and specific programming to minimize the
375 sensing of noise.⁸¹⁻⁹²

376 **Duration Criteria for the Detection of Ventricular Arrhythmia**

377 Until recently, default device programming used short-duration “detection” criteria that varied by
378 manufacturer and a tachycardia rate of approximately 2.8 to 5 seconds before either anti-tachycardia
379 pacing or charging (including detection time plus duration or number of intervals).^{82,93} With increased
380 awareness of the potential harm from inappropriate shocks and the realization from stored pacemaker
381 electrograms that even long episodes of VT can self-terminate, a strategy of prolonged detection
382 settings has been explored. This strategy allows episodes to self-terminate without requiring device
383 intervention and reduces inappropriate therapy for non-malignant arrhythmias. The benefit of
384 programming a prolonged detection duration (30 or 40 beats) was first reported in the Prevention
385 Parameters Evaluation (PREPARE) study on exclusively primary prevention subjects (n=700), and
386 compared outcomes to a historical ICD cohort programmed at “conventional detection delays” with
387 about half programmed to 12 of 16 intervals within the programmed detection zone and half to 18 of 24
388 intervals.⁹⁴ The programming in PREPARE demonstrated a significant reduction in inappropriate shocks
389 for supraventricular arrhythmia and in avoidable shocks for VT. In addition, a composite endpoint was

390 reduced as well: the morbidity index, which consists of shocks, syncope, and untreated sustained VT.
391 Within the limitations of a non-randomized study, it was concluded that extending detection times
392 reduces shocks without increasing serious adverse sequelae.

393 In 2009, the Role of Long-Detection Window Programming in Patients with Left Ventricular Dysfunction,
394 Non-Ischemic Etiology in Primary Prevention Treated with a Biventricular ICD (RELEVANT) study
395 confirmed and expanded the results of the PREPARE trial in a cohort of 324 primary prevention CRT-D
396 patients with non-ischemic cardiomyopathy.⁹⁵ The subjects were treated with simplified VT
397 management, which implies much longer detection for VF episodes (30 of 40) compared with the
398 control group (12 of 16) and a monitor-only window for VT. As in PREPARE, the RELEVANT study group
399 experienced a significantly reduced burden of ICD interventions (81% reduction) without increasing the
400 incidence of syncope. Fewer inappropriate shocks and HF hospitalizations were reported in the
401 RELEVANT study group compared with the control group.

402 The Multicenter Automatic Defibrillator Implantation Trial: Reduce Inappropriate Therapy (MADIT-RIT),
403 a 3-arm study, compared a conventional programming strategy (a 1-second delay for VF [equivalent to
404 approximately 12 intervals including detection plus delay] and a 2.5-second delay for VT detection
405 [equivalent to approximately 16 intervals including detection plus delay]) (Arm A) to both a high-rate
406 cutoff with a VF zone starting at 200 bpm (Arm B) (discussed elsewhere) and to a delayed therapy
407 strategy with a 60-second delay for rates between 170 and 199 bpm, a 12-second delay at 200 to 249
408 bpm, and a 2.5-second delay at 250 bpm (Arm C).⁹⁶ The MADIT-RIT population was exclusively primary
409 prevention and included approximately an equal proportion of non-ischemic and ischemic
410 cardiomyopathy patients. All the patients were implanted with either a dual-chamber ICD or a CRT-D
411 programmed to deliver ATP before charging. After a mean 1.4-year follow-up, the prolonged detection
412 group (Arm C) was associated with a reduction in treated VT/VF leading to a 76% reduction in the
413 primary endpoint of the first inappropriate therapy ($P < .001$), as well as a significant reduction in the
414 first appropriate therapy, appropriate ATP, and inappropriate ATP, but not in appropriate or
415 inappropriate shock.

416 The Avoid Delivering Therapies for Non-Sustained Arrhythmias in ICD Patients III (ADVANCE III) trial
417 reported that a long detection was associated with a highly significant reduction of overall therapies
418 (appropriate and inappropriate ATP and/or shocks), inappropriate shocks, and all-cause
419 hospitalizations.⁹⁷ Importantly, like PREPARE, RELEVANT, and MADIT-RIT, the extended detection
420 duration used in the ADVANCE III trial (30 of 40) did not negatively impact the rate of syncopal events.

421 There was no significant difference in mortality between the optimal and the conventional programming
422 groups. Compared with the MADIT-RIT trial, the ADVANCE III control group had a longer detection
423 duration (primarily in the VF zone), and enrolled a larger cohort of subjects covering all ICD types (single,
424 dual, and CRT with ATP delivered during charging) for both primary and secondary prevention
425 indications. Finally, the Programming Implantable Cardioverter-Defibrillators in Patients With Primary
426 Prevention Indication (PROVIDE) trial randomized 1670 patients to conventional programming (12-beat
427 detection in each of 2 zones) or experimental programming (2 VT and 1 VF zone requiring 25-, 18-, and
428 12-beat detection, respectively).⁹⁸ PROVIDE observed a significant 36% reduction in the 2-year all-cause
429 shock rate, and an improved survival (hazard ratio [HR]: 0.7; 95% confidence interval [CI]: 0.50–0.98; $P =$
430 .036).

431 Whereas PREPARE, RELEVANT, MADIT-RIT and PROVIDE only enrolled primary prevention patients, a
432 subset of the ADVANCE III study evaluated the efficacy and safety of a long-detection approach in
433 secondary prevention patients who have a known higher burden of arrhythmic episodes. In this
434 particular subset of 25% of the enrolled patients, ADVANCE III reported that a long detection duration
435 reduced the overall therapies delivered, primarily due to a significant 36% reduction in appropriate
436 shocks.⁹⁹ Syncopal episodes related to arrhythmic events and deaths were similar between the 2 groups.

437 Following shortly on the heels of these trials, 2 meta-analyses including the above studies were
438 published in 2014. Tan et al presented the data from the RELEVANT, PREPARE, MADIT-RIT, ADVANCE III,
439 PROVIDE, and EMPIRIC trials.^{100,101} A 30% reduction in the risk of death was found in the therapy
440 reduction group when including all 6 studies; however, similar results were observed when separately
441 considering the 4 randomized trials and the 2 observational studies. Data on the appropriateness of
442 shocks were available only for RELEVANT, MADIT-RIT, ADVANCE III and PROVIDE, and a 50% reduction in
443 inappropriate shock was observed without an increased risk of syncope and appropriate shock.

444 A meta-analysis evaluated the impact of a prolonged arrhythmia detection duration on outcome¹⁰²—
445 thus excluding the EMPIRIC trial (which used 18 of 24 intervals for VF detection), the PREPARE trial
446 (which used a historical control group), and the high-rate therapy arm of the MADIT RIT. Analyzing the
447 cohort of patients enrolled in RELEVANT, Arm C of MADIT-RIT, ADVANCE III, and PROVIDE, the meta-
448 analysis reported a reduction of overall burden of therapies, driven by the greater than 50% reduction in
449 appropriate and inappropriate ATP and the 50% reduction in inappropriate shocks. A reduction in all-
450 cause mortality was observed without an increase in the risk of syncope.

451 All the reports above clearly stress the necessity to consider a long detection window setting as a
 452 "default" strategy for ICD programming. Moreover, they underline the importance of choosing to
 453 reprogram the ICD rather than use the manufacturers' out-of-the-box settings. A summary of the large
 454 comparative datasets of tachycardia detection is presented in Table 2.

455 **Table 2.** Tachycardia Detection Evidence

Study	Participants (N)	Short Detection Controls	Prolonged Detection Intervention	Findings
PREPARE	1391 Non-Randomized Primary Prevention	12 of 16 (58%) 18 of 24 (42%)	30 of 40	Reduction in inappropriate shocks (SVT), avoidable shocks (VT), and "morbidity index"
RELEVANT	324 Non-Randomized Primary Prevention	12 of 16	30 of 40	Reduction in inappropriate shocks (SVT), avoidable shocks (VT), and HF hospitalizations
MADIT-RIT	1500 Randomized Primary Prevention	2.5 s (170–199 bpm) 1 s (≥ 200 bpm)	60 s (170–199 bpm) 12 s (200–249 bpm) 2.5 s (≥ 250 bpm)	Reduction in first inappropriate therapy, first appropriate therapy, appropriate ATP, and inappropriate ATP; Improved Survival
ADVANCE-III	1902 Randomized Primary & Secondary Prevention	18 of 24	30 of 40	Reduction in overall therapies, inappropriate shocks, and all-cause hospitalizations
PROVIDE	1670 Randomized Primary	12 beats	25 beats (180–214 bpm) 18 beats	Reduction in all-cause shock rate; Improved survival

	Prevention		(214–250 bpm) 12 beats (>250 bpm)	
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457 **Limitations of Data on the Duration of Tachycardia Required for Detection**

458 Although the findings on the effect of tachycardia detection duration are based on roughly 7000
459 patients, there are limitations. Data on secondary prevention patients are limited to 25% of the 1902
460 patients enrolled in the ADVANCE III trial (n=477). Although this proportion is a fair representation of
461 the real-world population receiving an ICD, more data are needed to fully understand the impact of a
462 long-detection strategy in this subgroup of patients. MADIT-RIT and RELEVANT did not include single-
463 chamber ICDs and MADIT-RIT excluded patients with permanent atrial fibrillation. The PROVIDE and
464 MADIT-RIT trials were designed to assess the time to first therapy and not the overall rate of therapies.
465 MADIT-RIT, ADVANCE III, RELEVANT, and PROVIDE used devices from 3 different manufacturers with
466 detection strategies leading to different detection times, intervals, and definitions. Some manufacturers
467 of ICDs are not represented at all in these trials. Programming in the trial control groups was highly
468 heterogeneous, with time until ATP or charging for VF as varied as about 11–12 intervals (approximately
469 3.4 seconds at 200 bpm) in MADIT-RIT and PROVIDE and 18 intervals (approximately 5.4 seconds) in
470 ADVANCE III. An approximate translation of the impact of the number of intervals to detection and
471 tachycardia cycle length are listed in Table 3. A further limitation is the relatively short duration and lack
472 of inclusion of the patients with the most severe illness receiving an ICD. This limitation minimizes the
473 exposure to relatively rare events that might occur in non-clinical trial, “real world” patients. Lastly, as
474 ICD batteries deplete, the charge time lengthens. The effect of such a delay to shock therapy in addition
475 to prolonged detection times has not been studied.

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479

480 **Table 3.** Approximating the time taken to detect 30 intervals using fixed 8 of 10 interval detection plus
 481 adding a time delay, for a range of heart rates.

Arrhythmia characteristic		Interval-based detection	8 of 10 interval detection then delay	
Beats per minute	CL (ms)	Time to detect 30 intervals (s)	Time to detect 8 intervals (s)	Subsequent delay to approximate a 30-interval detection time
180	333	10.0	2.7	7.0
200	300	9.0	2.4	6.5
220	273	8.2	2.2	6.0
240	250	7.5	2.0	5.5
260	231	6.9	1.8	5.0
280	214	6.4	1.7	4.5
300	200	6.0	1.6	4.5

482

483 **Rate criteria for the detection of ventricular arrhythmia**

484 Ventricular tachyarrhythmia detection by implantable devices is primarily based on heart rate. Heart
 485 rates can be extremely rapid during ventricular tachyarrhythmias, and it is less likely that such rates are
 486 achieved during supraventricular tachyarrhythmias—thus making rate a powerful component of
 487 arrhythmia discrimination. However, VT can also present slower rates in the range of those of
 488 supraventricular tachyarrhythmias or even of sinus tachycardia. Therefore, any rate cut-off will always
 489 imply a trade-off between maximizing sensitivity for ventricular tachyarrhythmia detection at the
 490 expense of inappropriate detection of fast supraventricular tachyarrhythmias and maximizing specificity
 491 at the expense of some slow VTs going undetected.¹⁰³

492 Because ICD therapy was initially employed in secondary prevention patients, the cutoff rate was usually
 493 tailored to a rate slightly below that of the observed VT. With the development of ICD use in primary
 494 prevention, the detection rate came into question because there is no history of sustained tachycardia
 495 in these patients. The recognition of a significant rate of inappropriate therapies in primary prevention
 496 studies, and their potentially deleterious consequences, prompted the development of studies that
 497 tested whether programming faster rate criteria reduced avoidable ICD therapies, particularly shocks. In

498 many of these studies, however, testing also involved programming parameters other than rate, and
499 those have been discussed as described below.

500 In the MADIT-RIT trial of primary prevention patients, conventional therapy (rate cutoff 170 bpm,
501 n=514) was compared with a “high-rate group” in which rate cutoff was 200 bpm (n=500).⁹⁶ The primary
502 endpoint of first occurrence of inappropriate therapy was observed in 20% of the conventional group
503 and in 4% of the high-rate group ($P < .001$) over a mean follow-up of 1.4 years. ICD shocks occurred in 4%
504 and 2% of patients in the conventional and high-rate groups, respectively. The proportion of patients
505 with appropriate therapies was also significantly different (22% vs 9% in the conventional and high-rate
506 groups, respectively). It is important to note that all-cause mortality in the conventional group (6.6%)
507 was approximately double that of the high-rate group (3.2%, $P = .01$).

508 In a single-center observational study, 365 primary prevention patients were prospectively studied, with
509 programming including a single shock-only zone over 220 bpm.¹⁰⁴ During a mean follow-up of 42
510 months, 11% of the patients (7% in the first 2 years) experienced appropriate shocks, and only 6.6%
511 experienced inappropriate shocks. It was notable that in the monitoring zone over 170 bpm, self-limiting
512 VT episodes were detected in 12% of the patients, but they were symptomatic in only 1.9%. The
513 mortality rate was 17%, with one case of unexplained sudden death.

514 A recent primary prevention study revealed that there was considerable overlap between the
515 ventricular rates of supraventricular and ventricular arrhythmias, and the majority of inappropriate
516 shocks occurred at rates between 181 and 213 bpm.⁹⁸ These data also support the notion that for
517 primary prevention patients it is safe to increase the rate cutoff up to 200 bpm to reduce these
518 potentially avoidable therapies, a practice that was also supported by the results of the MADIT-RIT trial.

519 In secondary prevention patients, no trial has randomized the detection rate and compared outcomes.
520 However, the ADVANCE III Secondary Prevention sub-study confirmed the safety of not programming
521 therapy for rates < 188 bpm; syncope was rare at 2 to 3 episodes per 100 patient-years.¹⁰⁵ Previously
522 published recommendations suggest a VT zone starting at 10 to 20 bpm slower than the observed
523 tachycardia rate, usually including a 2- or 3-zone arrhythmia detection scheme (as discussed
524 elsewhere).¹⁰⁶ Clinicians should allow a larger rate differential when starting a patient on an anti-
525 arrhythmic drug that might slow the clinical tachycardia rate (e.g., amiodarone).

526 **Single- or Multi-Zone Detection**

527 Modern ICDs allow the rate to be classified into single or multiple zones. This classification permits
528 different criteria to be applied for detection (e.g., number of intervals) and for tiered therapy (e.g.,
529 different adaptive cycle lengths for slower vs faster VTs and more sequences of ATP for slower and
530 presumably hemodynamically more stable VTs). Additionally, because some manufacturers tie SVT
531 discrimination algorithms to specific VT zones, programming more than one tachycardia zone allows for
532 greater specificity in discriminating VT from SVT (see online Appendix B). Although there are trials in
533 which arms differ in whether a single zone or multiple zones are used, this is typically performed to
534 allow programming of various detection, discrimination, or therapies for comparison. Thus the number
535 of zones was not the randomization variable being directly compared. Therefore, the concept of single-
536 versus multi-zone programming as a head-to-head comparison is not well tested. The MADIT-RIT study
537 randomized primary prevention ICD patients into 1 of 3 arms with single-, dual-, or triple-zone
538 programming (the single-zone arm also had a monitoring zone). Although the trial's aim was to compare
539 conventional therapy with high-rate and delayed therapy, the outcome for the single-zone arm (high-
540 rate) was comparable to the triple-zone (delayed) arm and superior to the dual-zone (conventional) arm,
541 with regard to inappropriate shock.⁹⁶ This study is consistent with multiple studies in ICD programming
542 in which the use of multiple-zone programming has allowed for flexibility in programming strategies
543 with regard to detection, discrimination, and therapy. Additionally, there are observational data from
544 the ALTITUDE Real World Evaluation of Dual-Zone ICD and CRT-D Programming Compared to Single-
545 Zone Programming (ALTITUDE REDUCES) study that show that dual-zone programming is associated
546 with fewer shocks than single-zone programming, at least for rates <200 bpm.⁶⁴ Therefore, the authors
547 conclude that using more than 1 detection zone can be useful for modern ICD programming. It should
548 be noted that ATP before or during charging was used in the majority of studies described in both the
549 tachycardia detection and therapy sections and thus is recommended for longer detection.

550 **Discrimination Between Supraventricular and Ventricular Arrhythmia**

551 The SVT-VT discrimination process classifies a sequence of sensed electrograms (EGMs) that satisfies
552 rate and duration criteria as either SVT (therapy withheld) or VT/VF (therapy given). *Discriminators* are
553 individual algorithm components that provide a partial rhythm classification or a definitive classification
554 for a subset of rhythms. *Discrimination algorithms* combine individual component discriminators to
555 produce a final rhythm classification. Discrimination algorithms vary among manufacturers and between
556 individual ICD models (see online Appendix B). The final rhythm classification can differ depending on the
557 technical details of how each individual discriminator is calculated, the nominal or programmed threshold

558 for each discriminator, the order in which discriminator components are applied, and the logical
559 connections between them (e.g., “and” vs “or”). In some ICDs, rhythms classified as VT/VF undergo a
560 subsequent sensing-verification step to confirm that EGMs represent true cardiac activation.

561 SVT-VT Discriminator Components

562 Individual discriminators can be considered in relation to the EGMs analyzed as ventricular-only or both
563 atrial and ventricular, by the rhythm that they identify (e.g., AF, sinus tachycardia, VT), or by the type of
564 EGM information analyzed (intervals vs morphology). Note that ventricular rate alone is a mandatory
565 discriminator, as discussed in the section above. We summarize the most commonly used
566 discriminators. More comprehensive discussions are available in the literature.¹⁰⁷⁻¹¹¹

567 *Rejection of Sinus Tachycardia by Onset*

568 Several interval-based discriminators focus on differences in the onset of sinus tachycardia (gradual and
569 parallel acceleration of atrial and conducted ventricular intervals) compared with VT (typically abrupt,
570 with at least transient atrioventricular dissociation). *Sudden (abrupt) onset* was one of the first single-
571 chamber, interval-based discriminators. It withholds therapy if acceleration across the sinus-VT rate
572 boundary is gradual. Because onset discriminators classify the rhythm only once, and thus cannot
573 correct misclassifications, they are now used infrequently and only with an override feature and/or
574 other discriminators.¹¹²⁻¹¹⁵ *Chamber of onset* is a related, interval-based, dual-chamber discriminator
575 that classifies a 1:1 tachycardia as SVT if the atrial rhythm accelerates at the device-defined onset. A
576 related, “Sinus Tachycardia®” discriminator classifies a tachycardia as VT if either the RR or the PR
577 intervals deviate sufficiently from the range of the immediately preceding sinus intervals.¹¹⁶

578 *Rejection of AF by Ventricular Interval Regularity*

579 Ventricular interval regularity (*interval stability*) is an explicit single-chamber, interval-based
580 discriminator that classifies the rhythm as AF if the ventricular intervals are sufficiently irregular.
581 Because interval variability in conducted AF decreases at faster rates, stability becomes unreliable in
582 discriminating VT from conducted AF at ventricular rates greater than 170 bpm.^{112,115} Interval stability
583 can also fail if drugs (e.g., amiodarone) cause monomorphic VT to become irregular or induce
584 polymorphic VT to slow into the SVT-VT discrimination zone.^{114,117}

585 *Diagnosis of VT by Dual-Chamber Components: Atrial vs Ventricular Rate and Atrioventricular*
586 *Association*

587 In contrast to the single-chamber discrimination algorithms above that diagnose SVT when their criteria
588 are fulfilled, 2 separate, interval-based, dual-chamber discrimination algorithms diagnose VT. First, **atrial**
589 **rate vs ventricular rate** diagnoses VT if the ventricular rate exceeds the atrial rate.¹¹⁸
590 Second, **atrioventricular dissociation** identifies isorhythmic VT during sinus tachycardia. Inversely, the
591 atrioventricular **association** discriminator diagnoses SVT in the presence of N:1 (e.g., 2:1, 4:1)
592 atrioventricular association consistent with atrial flutter at a fixed conduction ratio.

593 *The Ventricular Electrogram Morphology Discriminator*

594 This versatile, single-chamber discriminator is the only algorithm component that does not rely on inter-
595 EGM intervals. It classifies tachycardias as SVT if the morphology (shape) of the ventricular EGM is
596 sufficiently similar to the morphology during a conducted baseline rhythm. It can potentially
597 discriminate any SVT from VT, including SVTs that challenge other discriminators, such as abrupt-onset
598 1:1 SVTs and irregular VT during AF. Contemporary ICDs (including subcutaneous ICD [S-ICD]) analyze
599 EGMs from the shock electrodes, which record a larger field of view than EGMs from pace-sense
600 electrodes.¹¹⁹ They operate using a common series of steps and are susceptible to common failure
601 modes.^{111,120-123} The first common step is acquisition of a baseline rhythm template by mathematically
602 extracting EGM features and storing them. Both the acquisition of the initial template and the
603 subsequent template updating is automated in most ICDs. Nevertheless, physicians should confirm that
604 the conducted baseline beats match the template both at implant and during follow-up. For CRT
605 patients, the template must be manually collected. If the wavelet signal during template acquisition
606 appears clipped, adjustments specific to the manufacturer might be necessary.

607 SVT-VT Discrimination Algorithms

608 Discrimination algorithms combine component discriminators to provide a final rhythm classification of
609 VT/VF or SVT. The morphology discriminator frequently forms the primary component of single-chamber
610 algorithms with stability playing a secondary role and sudden onset used sparingly. By contrast, the
611 cornerstone of most dual-chamber algorithms is explicit or implicit comparison of atrial vs ventricular
612 rates. Because the ventricular rate is greater than the atrial rate in more than 80% of VTs, algorithms that
613 compare atrial and ventricular rates as their first step apply additional SVT discriminators to fewer than
614 20% of VTs, reducing the risk that they will misclassify VT as SVT.^{124,125} Most dual-chamber algorithms

615 further restrict single-chamber discriminators to tachycardias for which they offer the greatest benefit;
616 thus, stability is applied only if AF is confirmed by direct calculation of the atrial rate or the atrial rate is
617 greater than the ventricular rate. Similarly, sudden onset, chamber of onset, or 1:1 atrioventricular
618 association are applied only if the atrial rate equals the ventricular rate. The use of discriminators in
619 redetection varies among manufacturers and has not been systematically studied.

620 Assessing Clinical Benefits and Risks

621 *What evidence supports a benefit?*

622 1- The annual rate of inappropriate shocks has fallen dramatically from 37%–50% for SVT alone in
623 early studies to 1%–5% for all causes in modern clinical trials.^{97,118,126-128} This decrease is likely
624 due to differences in both clinical populations and the programming of multiple ICD parameters,
625 including longer detection time and higher rate cutoffs. Thus it is difficult to isolate the
626 differential effect of SVT-VT discrimination algorithms using clinical data. These studies have
627 programmed discrimination algorithms to ON, however, so it seems reasonable to use them.

628 2- Although clinical trials that reported dramatic reductions in shocks for SVT programmed
629 discrimination algorithms consistently, they have been programmed inconsistently in clinical
630 practice; and the rate of inappropriate shocks for SVT has been higher in observational studies
631 of remote-monitoring ICD databases. In the ALTITUDE REDUCES study on 15,991 patients in the
632 Latitude® database, SVT was the most common cause of shocks when the detection rate was
633 ≤ 180 bpm.¹²⁹ For detection rates ≤ 170 bpm, the rate of inappropriate shocks at 1 year was
634 significantly lower with dual-zone programming, which permits SVT-VT discrimination, than
635 single-zone programming, which does not (9.6% vs 4.3%). Similarly, Fischer analyzed shocks in
636 106,513 patients in the CareLink® database; programming SVT-VT discrimination ON was
637 associated with a 17% reduction in all-cause shocks.¹³⁰

638 3- Sophisticated simulations indicate that SVT-VT discrimination algorithms have substantial
639 benefit. For example, the SCD-HeFT study on primary-prevention patients did not use
640 discriminators. A validated Monte Carlo simulation predicted that use of single- or dual-chamber
641 SVT-VT discriminators alone would have reduced inappropriate shocks for SVT by 75.5% and
642 78.8%, respectively.¹³¹

643 *Which patients are most likely to benefit, and which are least likely to benefit?*

644 Despite limited direct evidence, it seems clear that patients will benefit most if the rates of their VTs and
645 SVTs overlap. This includes patients with slower monomorphic VT, those at risk for AF with rapid
646 ventricular rates, or those capable of exercising to sinus rates in the VT zone.^{103,132} In secondary-
647 prevention patients with slower VT, older discrimination algorithms reduced shocks for SVT compared
648 with rate-only detection. The benefit is less for primary-prevention patients, secondary-prevention
649 patients at risk only for VF, and those who cannot sustain rapid atrioventricular conduction. Patients
650 with permanent complete atrioventricular block do not benefit.

651 *What are the risks?*

652 The risk of the misclassification of either VT or VF as SVT by the discrimination algorithms can either
653 prevent VT detection or delay the time to therapy (underdetection), as documented in clinically-
654 significant situations.^{112,113,115,125} When modern algorithms are programmed to recommended
655 parameters, clinically significant underdetection is rare. Large clinical trials on multiple shock-reduction
656 strategies (including SVT-VT discrimination) report no or minimal and statistically-insignificant increases
657 in syncope.^{95,97,126,133} Most reports do not include the causes of syncope, and thus do not permit
658 identification of whether discrimination algorithms contributed to any of the syncopal episodes by
659 prolonging detection. However, in the PREPARE study, no syncopal episode was caused by untreated
660 tachycardia.¹³³ In general, discriminators that re-evaluate the rhythm classification during ongoing
661 tachycardia reduce the risk of underdetection compared with those that withhold therapy if the rhythm
662 is misclassified by the initial evaluation (e.g., onset, chamber of origin algorithms).

663 Additional Considerations

664 *SVT Limit*

665 SVT-VT discrimination applies from the VT detection rate to the SVT limit rate, which is programmable
666 independently of the VT/VF therapy zones with some manufacturers (preferable), but which might be
667 linked to one of the zone boundaries in others. The minimum cycle length for SVT-VT discrimination
668 should be set to prevent clinically significant delays in the detection of hemodynamically unstable VT.
669 PREPARE, EMPIRIC, and MADIT-RIT all support the safety of empirical programming at 200 bpm.^{96,101,134}
670 In MADIT-II, approximately 50% of SVT episodes were faster than 170 bpm, and a few were as fast as
671 250 bpm.⁸² In INTRINSIC RV, SVT comprised 19% of episodes, with rates between 200 and 250 bpm.¹³⁵
672 More limited and preliminary data from PainFree SST support programming up to 222–230 bpm.^{116,136}
673 We suggest the SVT limit not exceed 230 bpm in adults without a patient-specific indication, based on

674 the low incidence of SVTs in this rate range among ICD patients and the potential—however small—for
675 misclassifying hemodynamically unstable VT.

676 *Duration-Based “Safety-Net” Features to Override Discriminators*

677 These features deliver VT/VF therapy if a tachycardia satisfies the ventricular rate criterion for a
678 sufficient duration, even if the discrimination algorithm indicates SVT. The premise is that the ventricular
679 rate during transient sinus tachycardia or AF will decrease to below the VT rate boundary before the
680 override duration is exceeded. In one study, an override duration of 3 minutes delivered inappropriate
681 therapy to 10% of SVTs.¹¹² Because SVT is much more common than VT, programming an override
682 duration of less than 5–10 minutes results primarily or solely in inappropriate SVT therapy.¹²² Although
683 more data would be useful, in the absence of a documented benefit, we recommend programming this
684 feature OFF or long (minutes) without a patient-specific or device-specific indication.

685 *Dual-Chamber vs Single-Chamber Algorithms*

686 Clinical trials and simulated testing of induced arrhythmias that compared single- vs dual-chamber
687 discriminators have reported inconsistent results.^{10,33,137-139} Two meta-analyses found no superiority of
688 dual-chamber ICDs in terms of mortality or inappropriate therapies.^{11,140} Any benefit of dual-chamber
689 discrimination is likely restricted to specific patient groups.^{103,138} For example, the Dual Chamber and
690 Atrial Tachyarrhythmias Adverse Events (DATAS) trial of predominantly secondary prevention patients
691 with slower VTs reported modest benefit from dual-chamber discrimination, while the recent Reduction
692 and Prevention of Tachyarrhythmias and Shocks Using Reduced Ventricular Pacing with Atrial Algorithms
693 (RAPTURE) trial of primary-prevention patients programmed to a fast detection rate (>182 bpm) and
694 long detection duration (30/40 intervals) did not.^{103,138,139} Inappropriate therapy for SVT occurred in only
695 2% of the patients in each group. Recent data from PainFree SST notes very low rates of inappropriate
696 shocks (3.7% for single chamber; 2.8% for dual and triple chamber after 2 years). The choice of device
697 was not randomized, suggesting that when physicians chose a dual- or triple-chamber device (perhaps
698 due to known atrial arrhythmia or bradycardia), inappropriate shock rates were minimized.¹³⁶ The
699 Optimal Anti-Tachycardia Therapy in Implantable Cardioverter-Defibrillator Patients Without Pacing
700 Indications (OPTION) trial randomized 462 patients to single- or dual-chamber programming and noted
701 inappropriate shock rates of 10.3% for single chamber versus 4.3% for dual chamber after 27 months (P
702 = .015). Atrial lead-related complications were 1.3%, therapy was delivered from 170 bpm (VT) and 200
703 bpm (VF), and no difference in ventricular pacing percentage was noted.¹⁴¹ Dual-chamber algorithms

704 probably reduce the risk of underdetection compared with single-chamber algorithms because more
705 than 80% of VTs with a ventricular rate greater than the atrial rate undergo no further analysis.^{103,124,125}
706 However, the rate of clinically significant underdetection with modern programming is so low that this
707 difference is rarely of clinical significance. In most patients, improved SVT-VT discrimination should not
708 be considered an indication for a dual- vs single-chamber ICD. Even if a dual-chamber ICD is implanted,
709 dual-chamber discrimination should be programmed only if the atrial lead becomes chronic or if atrial
710 sensing is unreliable. Accurate sensing of atrial electrograms is essential for dual-chamber SVT-VT
711 discrimination. Atrial lead dislodgments, oversensing of far-field R waves, or undersensing due to low
712 amplitude atrial signals can cause misclassification of VT/SVT. On implant, it is important to position the
713 atrial lead to minimize far-field R waves.

714 *Ventricular Oversensing*

715 Excluding recalled leads, ventricular oversensing accounts for less than 10% of inappropriate shocks, but
716 it often results in repetitive shocks and severe symptoms.^{82,142-144} Recently introduced features reduce
717 inappropriate therapies from oversensing of physiological T waves and non-physiological signals related
718 to pace-sense lead failures as discussed below.

719 **Programming to Reduce T-Wave Oversensing**

720 The problem of T-wave oversensing relates to the basic requirement that ICDs reliably sense VF, which is
721 characterized by RR intervals shorter than the normal QT interval and some EGMs with low amplitudes
722 and slow rates. Approaches to minimizing T-wave oversensing include reprogramming ventricular
723 sensitivity, altering sensing bandwidth, and changing the sensing bipole.^{109,123,145} One manufacturer
724 provides an algorithm that withholds therapy after rate and duration criteria for VT/VF are fulfilled if a
725 specific pattern of T-wave oversensing is identified.¹⁴⁶ T-wave oversensing rates vary based on device
726 design; using an appropriate high band-pass filter results in very low rates of T-wave oversensing.¹⁴²
727 Because T-wave oversensing is unpredictable, features that minimize T-wave oversensing should be
728 enabled proactively at implant, providing they do not cause undersensing in VF.¹⁴⁶

729 **Lead-Related Oversensing**

730 Oversensed signals caused by pace-sense lead failure have specific interval patterns and EGM
731 characteristics.^{145,147,148} Present algorithms identify three features: (1) intervals can be too short to
732 represent successive ventricular activations; (2) such short intervals are often transient and can be

733 repetitive; (3) in true bipolar leads, oversensed signals are absent on the shock EGM. Algorithms can
734 provide warning alerts, withhold shocks after spurious detection of VT/VF, or both. All 3 criteria can
735 provide alerts, but only the third is applied to withhold shocks. The present algorithms were developed
736 to identify impending lead failures on recalled leads, notably the Sprint Fidelis. These algorithms might
737 not be appropriate for detecting failures in other leads.¹⁴⁴ There is a high false-positive rate when using
738 these algorithms, and caregivers must carefully review the device data that caused the alert to ensure
739 the lead experienced a true failure.¹⁴⁵

740 Alerts that combine both oversensing and abrupt changes in impedance trends provide earlier warning
741 of lead failure than a fixed impedance threshold.^{144,145,149} Such alerts can be delivered via wireless
742 remote monitoring and/or by notifying the patient via vibration or an audible tone. Caregivers must
743 respond rapidly to alerts to minimize inappropriate shocks.^{144,149} Wireless remote monitoring has been
744 reported to reduce response time.¹⁵⁰ The principal disadvantage of lead alerts is false-positive triggers.
745 The principal risk of shock-withholding algorithms is a failure to shock VF, which is extremely rare.¹⁵¹ In
746 addition to algorithmic approaches, oversensing due to failure of the cable leading to the ring electrode
747 can be prevented by changing the programming of the sensing configuration from true bipolar to
748 integrated bipolar. This approach is appropriate prophylactically or as temporary programming after a
749 ring electrode cable failure; it is not a permanent solution, however, because increased rates of high-
750 voltage cable fractures have been documented after sensing cable fractures.¹⁵²

751 **The Subcutaneous Defibrillator (S-ICD)**

752 The novel S-ICD follows many of the same principles as intravascular ICDs but is considered here
753 separately for duration criteria, rate criteria, and discrimination algorithms. Candidates for the S-ICD
754 must initially be screened with a modified tri-channel surface electrocardiogram that mimics the sensing
755 vectors of the S-ICD system. This test is designed to assess the R-wave to T-wave ratio for appropriate
756 signal characteristics and relationships. If the screening is not satisfactory for at least 1 of the 3 vectors
757 supine and standing, an S-ICD should not be implanted. On implant, the S-ICD automatically analyzes
758 and selects the optimal sensing vector.

759 Detection of VT or VF by the S-ICD is programmable using a single or dual zone. In the single-zone
760 configuration, shocks are delivered for detected heart rates above the programmed rate threshold: the
761 “shock zone.”¹³⁴ In the dual-zone configuration, arrhythmia discrimination algorithms are active from
762 the lower rate: the “conditional shock zone.” In this latter zone, a unique discrimination algorithm is

763 used to classify rhythms as either shockable or non-shockable. If they are classified as supraventricular
764 arrhythmias or non-arrhythmic oversensing, therapy is withheld.

765 With dual-zone programming, the shock zone uses rate as the sole method for rhythm analysis. In
766 contrast, the conditional shock zone uses a stepwise discrimination algorithm to distinguish shockable
767 from non-shockable rhythms. The conditional shock zone has a morphology analysis process based on a
768 normal rhythm trans-thoracic QRS:T wave template. The template uses up to 41 fiducial points to
769 reconstruct morphology for the template as well as the programmed targeted heart rate zones. The
770 comparison of the template to the high-rate rhythm electrocardiogram for discrimination constitutes
771 the static waveform analysis. A good template match designates a sensed beat as supraventricular,
772 thereby preventing a shock. A poor match to the static QRS:T morphology template moves the algorithm
773 to a dynamic waveform analysis that compares single-beat morphologies in groups of 4 beats for
774 uniformity. A consistent dynamic waveform match adjusts the sensing to evaluate QRS width. If a
775 tachycardia has a prolonged QRS width compared with the template width (>20 ms) and is of sufficient
776 duration, it will lead to a shock.

777 The system uses an initial 18 of 24 duration criteria (non-programmable) prior to initiating capacitor
778 charging; however, this duration is automatically extended following nonsustained ventricular
779 tachyarrhythmia events. A confirmation algorithm is also used at the end of capacitor charging to ensure
780 persistence of the ventricular arrhythmia prior to shock delivery. Shocks for spontaneous (noninduced)
781 episodes are delivered at a nonprogrammable 80 J regardless of the therapy zone of origination.

782 When programmed to include a conditional shock zone, the S-ICD VT detection algorithm has been
783 demonstrated to be more effective than transvenous ICD systems programmed at nominal settings to
784 prevent the detection of induced supraventricular arrhythmias.¹⁵³ Furthermore, in the clinical evaluation
785 of the conditional shock zone, the S-ICD system was strongly associated with a reduction in
786 inappropriate shocks from supraventricular arrhythmias and did not result in prolongation of detection
787 times or increased syncope.¹⁵⁴

788 **Integrating Tachycardia Detection Data Into Programming Recommendations**

789 When taking data from specific single-manufacturer studies and producing generic guidelines applicable
790 across all ICDs, some compromises and potential pitfalls have been encountered. Nevertheless, it is our
791 intention to convey the general principles of good quality evidence (e.g., extending detection time) to
792 apply to ICD programming in general. Thus, attempts have been made to translate interval-based

793 detection to time-based detection and to provide a range of reasonable heart rate cutoffs that are
 794 inclusive of those proven in good-quality trials. We encourage programming ICDs to manufacturer-
 795 specific therapies of proven benefit; however, when evidence is lacking, the guidelines provide a
 796 framework for programming within the evidence base. See online Appendix B for manufacturer-specific
 797 examples of optimal ICD programming.

798

Tachycardia Detection Programming Recommendations	Class of Recommendation	Level of Evidence
<p>For primary prevention ICD patients, tachyarrhythmia detection duration criteria should be programmed to require the tachycardia to continue for at least 6-12 seconds* or for 30 intervals before completing detection, to reduce total therapies.</p> <p><i>*Tachyarrhythmia detection duration is directly related to the tachyarrhythmia rate. Direct evidence to support a delay >2.5 seconds for rates over 250 bpm is not available, but can be inferred from evidence that 30 detection intervals are safe at that rate.</i></p>	I	A
<p>For primary prevention ICD patients, the slowest tachycardia therapy zone limit should be programmed between 185 and 200bpm*, to reduce total therapies.</p> <p><i>*Higher minimum rates for detection might be appropriate for young patients or for those in whom SVT-VT discriminators cannot reliably distinguish SVT from VT, provided there is no clinical VT below this rate.</i></p>	I	A
<p>For secondary prevention ICD patients, tachyarrhythmia detection duration criteria should be programmed to require the tachycardia to continue for at least 6-12 seconds* or for 30 intervals before completing detection, to reduce total therapies.</p> <p><i>*Tachyarrhythmia detection duration is directly related to the tachyarrhythmia rate. Direct evidence to support a delay >2.5</i></p>	I	B-R

<i>seconds for rates over 250 bpm is not available, but can be inferred from evidence that 30 detection intervals are safe at that rate.</i>		
<p>Discrimination algorithms to distinguish SVT from VT should be programmed to include rhythms with rates faster than 200 bpm and potentially up to 230 bpm (unless contraindicated*) to reduce inappropriate therapies.</p> <p><i>*Discrimination algorithms and/or their individual components are contraindicated in patients with complete heart block or if the algorithm/component is known to be unreliable in an individual patient. Dual-chamber discriminators that misclassify VT as SVT if the atrial lead dislodges are discouraged in the perioperative period. Dual-chamber discriminators are contraindicated in patients with known atrial lead dislodgment, atrial undersensing or oversensing of far field R waves, and in those with permanent AF.</i></p>	I	B-R
It is recommended to activate lead-failure alerts to detect potential lead problems.	I	B-NR
<p>For secondary prevention ICD patients where the clinical VT rate is known, it is reasonable to program the slowest tachycardia therapy zone at least 10 bpm below the documented tachycardia rate but not faster than 200bpm*, to reduce total therapies.</p> <p><i>*Higher minimum rates for detection might be appropriate for young patients or for those in whom SVT-VT discriminators cannot reliably distinguish SVT from VT, provided there is no clinical VT below this rate.</i></p>	IIa	C-EO
It can be useful to program more than one tachycardia detection zone to allow effective use of tiered therapy and/or SVT-VT discriminators and allow for a shorter delay in time-based detection programming for faster arrhythmias.	IIa	B-R
When a morphology discriminator is activated, it is reasonable to re-acquire the morphology template when the morphology match	IIa	C-LD

is unsatisfactory, to improve the accuracy of the morphology discriminator.		
It is reasonable to choose single chamber ICD therapy in preference to dual chamber ICD therapy if the sole reason for the atrial lead is SVT discrimination, unless a known SVT exists that may enter the VT treatment zone, to reduce both lead-related complications and the cost of ICD therapy.	IIa	B-NR
For the S-ICD, it is reasonable to program 2 tachycardia detection zones: 1 zone with tachycardia discrimination algorithms from a rate ≤ 200 bpm and a second zone without tachycardia discrimination algorithms from a rate ≥ 230 bpm, to reduce avoidable shocks.	IIa	B-NR
Programming a non-therapy zone for tachycardia monitoring might be considered to alert clinicians to untreated arrhythmias.	IIb	B-NR
It may be reasonable to disable the SVT discriminator time-out function, to reduce inappropriate therapies.	IIb	C-EO
It may be reasonable to activate lead "noise" algorithms that withhold shocks when detected VT/VF is not confirmed on a shock or other far-field channel to avoid therapies for non-physiological signals.	IIb	C-EO
It may be reasonable to activate T-wave oversensing algorithms, to reduce inappropriate therapies.	IIb	C-LD
It may be reasonable to program the sensing vector from bipolar to integrated-bipolar in true-bipolar leads at risk for failure of the cable to the ring electrode to reduce inappropriate therapies.* <i>*This is not intended as a long-term solution when a cable fracture has been identified.</i>	IIb	C-EO

799

800

801 **Tachycardia Therapy Programming**

802 Although therapies delivered by the ICD can abort sudden cardiac death, appropriate and inappropriate
803 ICD shocks have been associated with a considerable increase in the risk of mortality.^{82,83,155-158} In the
804 Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT), the risk of mortality was 5-fold higher in
805 patients who received appropriate ICD shocks and 2-fold higher in patients who received inappropriate
806 shocks.⁸³ Similarly, pooling data from 4 studies of 2135 ICD patients, shocked VT was associated with a
807 32% increase in the risk of mortality. In that analysis, shocked patients had poorer survival than patients
808 treated with ATP only.¹⁵⁵ ICD shocks are likely a marker of more advanced heart disease and subsequent
809 death, but defibrillation therapies have been associated with troponin release and increased LV
810 dysfunction with the potential of further mortality risk.

811 The incidence of appropriate and inappropriate ICD shocks depends on the patient's characteristics,
812 including the indication for the device, concomitant medical therapies including antiarrhythmic
813 medications, programming of the ICD, and the duration of follow-up. With regard to ICD programming,
814 faster VT/VF detection rates, longer detection durations, use of a single zone, use of SVT discriminators,
815 and delivery of ATP have been shown to reduce both appropriate and inappropriate shocks and to
816 improve quality of life.^{91,101,126,129,130,133,159,160} This programming might improve survival.¹²⁶ Indeed, several
817 studies have shown that ATP is effective at terminating slow and fast VT with exceedingly low rates of
818 adverse events like syncope.^{93,135,161-165} The initial bias of the ICD community was to reserve ATP therapy
819 for those patients in whom the therapy was demonstrated to be effective, usually during an
820 electrophysiologic study. However, the approach of physician-directed programming based on the
821 knowledge of induced arrhythmias was found to be significantly inferior to the routine strategic
822 (EMPIRIC) programming of ATP. It is not reflective of the arrhythmias experienced outside the EP lab for
823 primary and secondary prevention patients with ischemic and non-ischemic substrates.^{101,166} Although
824 the ideal number of ATP bursts has not been definitively determined, current data support the use of up
825 to 2 ATP attempts, given additional attempts yield very little additional efficacy.^{93,135,161-165,167,168} In one
826 study, up to 5 attempts were found to be safe.¹⁶⁸ The most effective ATP duration is likewise uncertain;
827 however, in the ATP Delivery for Painless ICD Therapy (ADVANCE-D) trial—a prospective randomized
828 clinical trial of 925 patients—8-pulse ATP was as effective and safe as 15-pulse ATP.¹⁶⁹ The PITAGORA
829 ICD clinical trial randomized 206 patients with an ICD to 2 ATP strategies: an 88% coupling interval burst
830 versus a 91% coupling interval ramp. The results of the trial showed that over a median follow-up of 36
831 months and compared with ramp pacing, burst pacing was more effective for terminating fast VT

832 episodes (between CL 240 and 320 ms).¹⁷⁰ In a prospective study of 602 patients, a strategy of tiered
833 ATP and low energy shock was efficacious and safe in patients with VT CL greater than 250 ms, with
834 extremely low syncope rates.¹⁷¹ However, a “real-world” retrospective study on 2000 patients with 5279
835 shock episodes from the LATITUDE remote monitoring system showed that the success rate of first
836 shock as first therapy was approximately 90%, but the success rate was lower after failed ATP.
837 Therefore, that study recommended programming a higher level of energy after ATP.¹⁷² Finally, a
838 substudy of the Effectiveness and Cost of ICD Follow-Up Schedule with Telecardiology (ECOST) study,
839 which randomly assigned 433 patients to remote monitoring (n=221; active group) versus ambulatory
840 follow-up (n=212; control group) showed that remote monitoring was highly effective in the long-term
841 prevention of inappropriate ICD shocks through early detection and prevention of atrial fibrillation with
842 a rapid ventricular rate, nonsustained VT, or diverted VT episodes.¹⁷³

843 **Benefits and Risks**

844 The goal of ICD therapy is to prolong life while causing as little morbidity as possible. Although survival is
845 quantifiably objective, morbidity is more subjective and includes both physical and emotional
846 components. Clearly, shocks are usually painful to the patient, whereas ATP is typically not
847 uncomfortable. However, there can be other morbidities related to both therapies, including mild to
848 extreme emotional distress, syncope, palpitations, and proarrhythmia yielding more therapies and
849 occasionally leading to death. Paradoxically, the need for life-saving therapies, including shocks and
850 potentially ATP, might also be associated with increased mortality; however, the causal relationships are
851 unclear. Also, the prevalence of tachycardia amenable to ATP or hemodynamic significance varies with
852 the mechanism of the risk (e.g., long QT vs ischemic cardiomyopathy). In addition, although the risk of
853 having a hemodynamically important or life-threatening arrhythmia can vary from patient group to
854 patient group, the largest proportion of patients in whom ICD therapy is applied has yet to have a
855 previously recorded arrhythmia, and we must therefore strategically choose on the basis of other
856 factors how we will treat the first event and subsequent events.

857 **Classification of Therapy**

858 The literature uses definitions of therapies that differ from each other and that impact their results and
859 conclusions. The occurrence rates of these events are dependent not only on their definition, but are
860 also highly dependent on the programming of the defibrillation system. Both shock and nonshock
861 therapies can be categorized as being appropriate, inappropriate, and avoidable. Whereas appropriate

862 and inappropriate therapies refer to therapies that were actually delivered, avoidable therapies are
863 theoretical events in the future. These potential future tachycardia therapies, delivered for either
864 appropriately or inappropriately detected events, can frequently be avoided by establishing
865 programming to either prevent the initiation of the arrhythmia or to allow the condition to pass without
866 therapy.

867 Appropriate

868 A response to a sustained ventricular arrhythmia (VT, VF) or hemodynamically poorly tolerated
869 arrhythmias (e.g., associated with syncope, rate over 200 bpm or hemodynamically compromising
870 supraventricular arrhythmias).

871 Inappropriate

872 A response to signals generated by something other than sustained ventricular arrhythmias or
873 hemodynamically poorly tolerated arrhythmias. Possible signals include supraventricular rhythms such
874 as sinus tachycardia, atrial fibrillation, atrial flutter, reentrant SVT, atrial tachycardia, or instances of
875 signal misinterpretation. Signal misinterpretation includes multiple counting of single events (e.g., atrial,
876 T-wave or R-wave), environmental signals such as electromagnetic interference, frequent PVCs and
877 nonsustained ventricular arrhythmias, extracardiac physiologic signals (e.g., diaphragmatic or pectoral
878 myopotentials), other implantable electronic devices (e.g., pacemakers, left ventricular assist devices,
879 nerve stimulators), inappropriate lead placement or dislodgement, conductor or insulation failures,
880 header connection instability, and pulse generator failure.

881 Avoidable

882 Programming of detection and therapy parameters and algorithms so that shock or ATP therapy is
883 withheld from arrhythmias that would be expected to be hemodynamically tolerated. Examples include
884 self-terminating ventricular arrhythmias, ATP-susceptible ventricular arrhythmias, and overdrive
885 suppression responsive rhythms. Many appropriate and most inappropriate therapies are also
886 potentially avoidable.

887 Phantom

888 These are not true therapies; however, there is the patient's perception that a therapy was delivered.
889 Interrogation of the ICD and/or coincident rhythm monitoring does not identify a tachycardia or
890 therapy.

891

892 Unintended Consequences of ICD Therapy and ICD Therapy Programming

893 In the SCD-HeFT and MADIT II trials, inappropriate shocks more than doubled the risk of death. Mortality
894 rates were substantially higher after shocks: 10% within days after the first shock, 25% within 1 year,
895 and 40% by 2 years. The leading cause of death was progressive HF. In an analysis of the MADIT-CRT
896 trial, the patients with appropriate shocks experienced increased mortality when compared with the
897 patients without ICD shocks, after accounting for mechanical remodeling effects; this was not the case
898 for patients who received appropriate ATP only.¹⁵⁶ ICD shocks have also been associated with
899 independent predictors of mortality in the large ALTITUDE registry of 3809 ICD recipients and in a meta-
900 analysis of ICD trials in which ATP was applied.^{155,157} Emotional morbidities associated with ICD shocks
901 are well recognized and include anxiety, depression, and post-traumatic stress disorders.¹⁷⁴⁻¹⁷⁶ Phantom
902 shocks can result from fear and/or anxiety and have a reported incidence of 5% in a European study of
903 ICD recipients over 35 months of follow-up.¹⁷⁷ If possible, and when safe, it is best to avoid both the
904 discomfort and psychological impact of shocks for ventricular arrhythmias, supraventricular arrhythmias,
905 noise events including lead failures, and for self-terminating arrhythmias, as is discussed in the section
906 on tachycardia detection. The 1500-patient MADIT RIT study demonstrated a mortality reduction by
907 changing both tachycardia detection criteria and tachycardia therapy (shocks and ATP). Therefore it is
908 difficult to assign the outcome result to ATP, shocks, or both when compared with older, more
909 conventional programming.¹²⁶ In addition, in a randomized study of remote follow-up of ICDs, home
910 monitoring showed an incidence of 52% fewer inappropriate shocks, 72% fewer hospitalizations due to
911 inappropriate shocks, 76% fewer capacitor charges, and a significant positive impact on battery
912 longevity.¹⁷⁸

913 *ATP:* Several large clinical trials have established the safety and efficacy of ATP as a first-line therapy to
914 treat even very fast VTs.^{93,95,101,133} The use of first-line ATP involving VT at rates between 188 and 250
915 bpm in the PainFREE Rx II trial resulted in a 71% relative shock reduction.⁹³ In the PREPARE study, a
916 primary prevention cohort of 700 patients was programmed with 30 of 40 detection intervals with ATP-
917 first for VT between 182–250 bpm with SVT discriminators active up to 200 bpm. The results

918 demonstrated a robust absolute risk reduction for shocks at 1 year from 17% to 9% without an increase
919 in arrhythmic syncope when compared with historical controls.¹³³ Similar findings were noted in the
920 RELEVANT study, which evaluated a cohort of patients with non-ischemic heart disease and cardiac
921 resynchronization defibrillators.⁹⁵ In the earlier EMPIRIC study, standardized VT detection and ATP
922 therapy parameters demonstrated a reduction in shocks when compared with physician-tailored
923 treatment in a randomized assessment of 900 primary prevention patients.¹⁰¹ The use of ATP during ICD
924 capacitor charging has been clinically validated as safe and effective.¹⁶³ It is important to recognize that
925 inappropriate therapies including inappropriate ATP, delivered primarily in the setting of
926 supraventricular arrhythmias, have been associated with increased mortality in the MADIT-RIT and
927 MADIT-CRT trials.^{156,179} However, the overall safety of ATP and its role as a contributor to improved
928 survival is well established, particularly in terms of preventing avoidable ICD shocks.

929 *Customized vs Strategic Programming:* Because primary prevention patients have no prior ventricular
 930 arrhythmias, programming individual devices on implant is largely empiric. There are more data for
 931 secondary prevention patients, but how the patient will behave in the future is still uncertain. The ability
 932 to individualize the antitachycardia programming for patients with both primary and secondary
 933 prevention indications was tested in the EMPIRIC trial and found to be an inferior approach to prevent
 934 these therapy events.¹⁰¹ The application of standardized programming and borrowing data from the
 935 PainFREE Rx II and PREPARE studies resulted in a comprehensive review of programming and its
 936 application across manufacturers.

937 *Secondary Prevention:* For the secondary prevention ICD patient, specific knowledge of the patient's
 938 arrhythmia history facilitates the creation of an effective antitachycardia programming strategy. Using
 939 what is known about the ventricular arrhythmia, including any electrocardiograms, available telemetry
 940 strips, and EMS recordings, provides insight into the arrhythmia mechanism. In cases of monomorphic
 941 VT, discerning the rate (cycle length) and the hemodynamic impact is useful in making choices,
 942 particularly for detection at a minimum; the device must be programmed with active VT detection zones
 943 sufficient to cover the clinical arrhythmia. Slower, monomorphic VT that is better tolerated
 944 hemodynamically favors a robust approach using ATP termination with at least 2–3 sequences and at
 945 least 8 pulses. The use of a second burst of ATP has also been shown to increase effectiveness from 64%
 946 to 83% in the fast VT range of 188 to 250 bpm.¹⁶⁷ Although a second burst has clear value, value beyond
 947 2 bursts is limited, except in rare situations.¹⁰¹ The use of ICDs in patients with implanted left ventricular
 948 assist devices allows prolongation of detection times and programming of multiple ATP attempts
 949 without significant risk to the patient, and it reduces the opportunity for shock therapies. Adjunct
 950 medications and ablation of VT (or SVT) might also be considered for cases in which slow VT occurs or if
 951 there is an overlap between the SVT and VT rates, leading to ICD therapies.

952

953

Tachycardia Therapy Programming Recommendations	Class of Recommendation	Level of Evidence
It is recommended in all patients with structural heart disease and ATP capable ICD therapy devices that ATP therapy be active for all ventricular tachyarrhythmia detection zones to	I	A

include arrhythmias up to 230 bpm, to reduce total shocks except when ATP is documented to be ineffective or proarrhythmic.		
It is recommended in all patients with structural heart disease and ATP-capable ICD therapy devices that ATP therapy be programmed to deliver at least 1 ATP attempt with a minimum of 8 stimuli and a cycle length of 84%–88% of the tachycardia cycle length for ventricular tachyarrhythmias to reduce total shocks, except when ATP is documented to be ineffective or proarrhythmic.	I	A
It is indicated to program burst ATP therapy in preference to ramp ATP therapy, to improve the termination rate of treated ventricular tachyarrhythmias.	I	B-R
It is reasonable to activate shock therapy to be available in all* ventricular tachyarrhythmia therapy zones, to improve the termination rate of ventricular tachyarrhythmias. <i>*Rarely, to limit patient discomfort and anxiety, hemodynamically stable slow VT can be treated without programming a backup shock.</i>	IIa	C-EO
It is reasonable to program the initial shock energy to the maximum available energy in the highest rate detection zone to improve the first shock termination of ventricular arrhythmias unless specific defibrillation testing demonstrates efficacy at lower energies.	IIa	C-LD

954

955

956

957 Intraprocedural Testing of Defibrillation Efficacy

958 The efficacy of the ICD for the primary and secondary prevention of SCD has been well established in
959 several landmark clinical trials.¹⁸⁰⁻¹⁸⁵ Most of these trials have required induction, detection, and
960 termination of VF at the time of implantation as a measure of defibrillation efficacy and as a surrogate of
961 the ICD's ability to prevent SCD. Testing defibrillation efficacy has been considered an integral part of
962 ICD implantation for many years, and it is performed to establish the appropriate connection of high-
963 voltage electrodes and to test the ability of the ICD to detect and terminate VF with a shock. However,
964 identifying system failures or high defibrillation thresholds is difficult, mainly due to the low prevalence,
965 which also depends upon the definition employed, about 5% combined. Significant improvements over
966 the past 2 decades have reduced energy requirements for defibrillation.¹⁸⁶⁻¹⁸⁹ Similarly, current
967 transvenous ICD technology is capable of delivering energies of 35–40 J, raising the question of the value
968 of routine defibrillation testing (DT). Physicians have therefore gravitated to implanting ICDs with
969 minimum or no DT with wide variability in practice, despite a paucity of rigorous data. DT is currently
970 being performed during ICD implant in only about half the procedures.¹⁹⁰⁻¹⁹⁵ Studies evaluating DT are
971 summarized in Table 4.

972 One of the most important reasons to avoid DT at the time of ICD implantation is that testing might
973 result in complications or even death. The risks of DT include (a) those related to VF itself, which can
974 lead to circulatory arrest and hypoperfusion, (b) risks related to the shocks delivered to terminate VT,
975 and (c) risks related to anesthetic drugs that are required for heavier sedation, which are used to
976 provide patient comfort during testing.

977 Periprocedural Mortality

978 Although improved ICD technology has led to the need for fewer inductions of VF at the time of
979 implantation testing, procedure-related mortality has not been completely eliminated. Using modern
980 ICD technology with transvenous systems and biphasic waveforms, the perioperative mortality rate
981 within 30 days of implantation is reported to be 0.2% to 0.4%.^{191,196} Recent data from the NCDR Registry
982 demonstrated an in-hospital mortality of 0.03% following ICD implantation, with death occurring in the
983 lab in 0.02%.¹⁹⁶ A Canadian report from 21 implanting centers estimates that 3 of 19,067 (0.016%)
984 deaths are related to defibrillation testing.

985

986 DT-Related Complications

987 Complications occurring during ICD implantation procedures are infrequent and many can be directly or
988 indirectly related to DT. Adverse effects related to DT testing include myocardial injury, depression of
989 contractile function leading to worsening of HF, persistent hypotension, CNS injury, thromboembolic
990 events, or respiratory depression.

991 Transient central nervous system hypoperfusion and cerebral ischemic changes can be demonstrated
992 during intra-operative electroencephalographic (EEG) monitoring at the time of DT. However, EEG
993 recovery occurs within less than 30 seconds, with a slightly longer time to the return of middle cerebral
994 blood flow.¹⁹⁷⁻¹⁹⁹ However, the clinical relevance of this transient finding is unclear because DT does not
995 appear to cause cognitive dysfunction 24–48 hours following ICD implantation.^{200,201} Although an
996 increase in biochemical markers of myocardial injury can be observed during ICD implantation or after
997 spontaneous clinical shocks, true intraoperative myocardial infarction (MI) is rare, even when extensive
998 DT is performed.²⁰²⁻²⁰⁵ In 2 recent studies using transvenous ICDs and a more abbreviated testing
999 protocol, there was no significant increase in CK, CK-MB, myoglobin, and NT-proBNP before and after
1000 DT, whereas elevated levels of high-sensitive Troponin T were observed after DT.^{206,207} In the NCDR ICD
1001 Registry, the incidence of MI during ICD implantation was reported to be 0.02%.¹⁹⁶

1002 Defibrillator shocks and VF transiently depress contractile function, although fatal pulseless electrical
1003 activity is rare at the time of ICD implantation.^{202,206,208-210} Refractory VF has been reported to occur
1004 during DT, but this is also uncommon, particularly with contemporary devices. One study reported that
1005 all tested ICD shocks failed and at least 3 external rescue shocks were required in 0.5% of patients.²⁰³ A
1006 Canadian study reported that 27 of 19,067 (0.14%) implants required prolonged resuscitations during
1007 DT.²¹¹

1008 Thromboembolic complications can occur during DT in the presence of intracardiac thrombus or when
1009 there are less than 3 weeks of therapeutic and uninterrupted anticoagulation in the setting of atrial
1010 fibrillation. Stroke or TIA is reported to occur in 0.026%–0.05% of cases.^{204,211} Multiple strategies have
1011 been employed, but none were documented to reduce the incidence of thromboembolism, including
1012 the avoidance of DT. These include pre-procedure transesophageal echocardiography to exclude left-
1013 atrial appendage thrombus and deferring testing when a thrombus is identified, or using transthoracic
1014 echocardiography to detect left-ventricular thrombi.

1015 Anesthetic agents can contribute to complications related to a depressant effect on myocardial
1016 contractility or can lead to respiratory depression if oversedation occurs. Heavier sedation is typically
1017 used in patients undergoing DT. Although patients with underlying chronic obstructive pulmonary
1018 disease or sleep apnea might be at increased risk, oversedation and respiratory depression could occur
1019 in any patient. Randomized trial data can help to identify which adverse events are directly (or
1020 indirectly) related to DT. For example, stroke or TIA might be “directly” related to DT due to
1021 dislodgement of intra-cardiac thrombus during conversion of atrial fibrillation in the absence of
1022 therapeutic anticoagulation, and an episode of prolonged hypotension could result in reduced cerebral
1023 perfusion. Respiratory depression, respiratory failure requiring intubation, or hypotension might be
1024 direct results of DT or might be due to the drugs required to perform testing. Pulseless electrical activity
1025 or even death can occur with hemodynamic complications related to induction of VF or multiple
1026 external shocks. In contrast, DT can indirectly increase the risk for pneumothorax, perforation,
1027 tamponade, lead dislodgment, or infection as more leads are inserted, or the procedure might be
1028 prolonged due to the system modifications required to improve defibrillation efficacy; however, all
1029 these complications can also occur in the absence of DT. In addition, due to the rates and types of
1030 adverse events reported in the literature, it appears that overall complication rates are primarily driven
1031 by mechanical complications or infection, most of which are not related to DT.

1032 In a sub-study of the Resynchronization for Ambulatory Heart Failure Trial (RAFT), in which 145 patients
1033 were randomized to DT compared with no DT at the time of initial ICD implantation, the risk of
1034 perioperative complications was extremely low, regardless of DT performance.²¹² There was, however, a
1035 nonsignificant increase in the risk of death or HF hospitalization in the group that underwent DT.
1036 Likewise, no significant difference in implant-related complications was demonstrated in DT compared
1037 with the groups without DT in the Safety of Two Strategies of ICD Management at Implantation (SAFE-
1038 ICD) study, a prospective observational study of 2120 patients performed at 41 centers.²¹³ Similar
1039 findings were observed in the prospective randomized Test-No Test Implantable Cardioverter
1040 Defibrillator (TNT-ICD) pilot study on 66 patients, in which there was no difference in adverse events
1041 between patients who underwent testing compared with those who did not.²¹⁴

1042 The Shockless Implant Evaluation (SIMPLE) trial is the largest randomized study assessing the effect of
1043 DT on clinical outcomes.²¹⁵ This large-scale study randomized 2500 patients to DT or not at the time of
1044 ICD implantation; 1253 patients were randomly assigned to DT and 1247 were assigned to no-testing,
1045 and were followed for a mean of 3.1 years (SD 1.0). The primary outcome of arrhythmic death or failed

1046 appropriate shock was noninferior (90 [7% per year]) in the no-testing group compared with patients
1047 undergoing DT (104 [8% per year]; HR: 0.86; 95% CI 0.65–1.14; P non-inferiority $<.0001$). The first safety
1048 composite outcome occurred in 69 (5.6%) of 1236 patients with no testing and in 81 (6.5%) of 1242
1049 patients with DT, $P = .33$. The second, pre-specified safety composite outcome, which included only
1050 events most likely to be directly caused by testing, occurred in 3.2% of patients with no testing and in
1051 4.5% with DT, $P = .08$. Heart failure needing intravenous treatment with inotropes or diuretics was the
1052 most common adverse event (in 20 [2%] of 1236 patients in the no-testing group vs 28 [2%] of 1242
1053 patients in the testing group, $P = .25$). In summary, routine DT at the time of ICD implantation is
1054 generally well tolerated without a statistically significantly increased rate of complications, but it also
1055 does not improve shock efficacy or reduce arrhythmic death.

1056 Finally, the No Regular Defibrillation Testing In Cardioverter Defibrillator Implantation (NORDIC-ICD)
1057 trial, another prospective randomized parallel group multicenter non-inferiority trial conducted in 48
1058 centers in Europe, assessed the effects of DT at the time of ICD implantation on first shock efficacy.²¹⁶
1059 The primary endpoint was different from the SIMPLE trial and assessed the average first-shock efficacy
1060 for all true VT and VF episodes occurring in any patient during follow-up. NORDIC-ICD randomized 540
1061 patients to DT and 537 to no DT at the time of ICD implantation. During a median follow-up of 22.8
1062 months, the first shock efficacy was demonstrated to be noninferior in the patients undergoing ICD
1063 implantation without DT, with a difference in first shock efficacy of 3.0% in favor of the no-DT test group
1064 (95% CI -3.0%–9.0%; P non-inferiority $<.001$). Overall, 112 procedure-related serious adverse events were
1065 reported within 30 days of ICD implantation in 94 patients (17.6%) undergoing DT compared with 74
1066 patients (13.9%) not undergoing DT ($P = .095$). The authors concluded that defibrillation efficacy without
1067 DT was noninferior to ICD implantation with DT in left-sided ICD implants. Because no major benefit or
1068 harm associated with DT was detected, in patients with a left-sided pectoral implantation it is
1069 reasonable to omit routine VF induction and DT during ICD implantation, assuming stable ICD lead
1070 position and good sensing and capture function.²¹⁷⁻²²⁰ This approach is particularly applicable to patients
1071 with ischemic and idiopathic dilated cardiomyopathy, given these entities were well represented in the
1072 studied cohort. Patients well represented within the cohort included those with implantation in the left
1073 pectoral location, those indicated for primary and secondary prevention of sudden cardiac death, and
1074 patients with ischemic and non-ischemic cardiomyopathies. Fewer data are available regarding other
1075 cardiomyopathies, such as patients with hypertrophic obstructive cardiomyopathy, congenital
1076 channelopathies, patients undergoing generator replacement, and procedures in the right pectoral
1077 location. In these instances, and when there is any question of the adequacy of the lead position or

1078 function, DT is reasonable. It is worth emphasizing that a nontesting strategy requires an anatomically
1079 well-positioned defibrillation lead in the right ventricle with adequate sensing of intrinsic R-waves (>5–7
1080 mV), adequate pacing thresholds, and a thorough verification of proper lead connection.

1081 Other important considerations include the use of alternative right ventricular defibrillation lead sites
1082 such as the mid-septum. Pooled data from 2 randomized studies do not indicate a clinically relevant
1083 elevation of energy required for defibrillation with mid-septal sites. Positioning of the RV defibrillation
1084 lead in other positions such as the right ventricular outflow tract has not been systematically
1085 addressed.²²¹

1086 The SIMPLE trial data was consistent between subgroups, both from patients with single- or dual-coil
1087 ICD leads and with or without the use of amiodarone. More recently, the Multicenter Comparison of
1088 Shock Efficacy Using Single vs Dual-Coil Lead Systems and Anodal vs Cathodal Polarity Defibrillation in
1089 Patients Undergoing Transvenous Cardioverter-Defibrillator Implantation (MODALITY) study was
1090 reported.²²² This was a multicenter registry that prospectively followed 469 consecutive patients
1091 undergoing DT at the time of implant; 158 (34%) had dual-coil and 311 (66%) had single-coil lead
1092 systems configuration, 254 (54%) received anodal shock, and 215 (46%) received cathodal shock. In 35
1093 (7.4%) patients, the shock was unsuccessful. No significant differences in the outcome of DT using a
1094 single- versus dual-coil lead were observed, but the multivariate analysis showed an increased risk of
1095 shock failure using cathodal shock polarity (OR: 2.37; 95% CI 1.12–5.03). These and other registry data
1096 support the use of either single- or dual-coil leads, preferably programmed to deliver anodal
1097 shocks.^{211,213,223}

1098 Performing DT has not been determined to be harmful or inappropriate. One reason to perform DT in
1099 specific populations is that high defibrillation thresholds have been reported in 2.2% to 12% of subjects
1100 undergoing DT. The probabilistic nature of DT with the failure of a single shock 10 J below the maximum
1101 ICD output does not necessarily imply long-term ICD failure. Determinations of DT using multiple shock
1102 protocols have reported that a safety margin of only 5.2 ± 1.1 J has a 97.3% rate of successful VT/VF
1103 conversion;²²⁴ however, the inability to convert VF at maximum output occurs in approximately 1% of
1104 procedures during DT. The long-term outcomes of these patients have not been evaluated without
1105 modification of the lead system. Further supporters of DT suggest that routine testing is necessary to
1106 identify system integrity and sensing failures. R-wave amplitude ≤ 5 –7 mV at implant almost invariably
1107 reliably sense VF.^{190,221} Failure to sense and some inner insulation failures might only be detected by DT.
1108 This situation has not been systematically evaluated.

1109

1110 Contraindications to Defibrillation Threshold Testing

1111 A great paucity of systematic data limits the assessment of the literature regarding contraindications to
1112 DT. Most implanters tend to avoid DT in patients perceived to be at high risk. Information derived from
1113 an NCDR-ICD registry identified advanced age, impaired LVEF, NYHA Class IV HF, atrial fibrillation/flutter,
1114 need to withhold warfarin, and several other factors as high-risk situations. Unfortunately, the strength
1115 of these associations was weak, given the odds ratios were under 2.²²³ Other registries have identified
1116 patients with broader QRS durations, advanced NYHA class, and cardiac resynchronization therapy as
1117 reason for not performing DT.²¹³ There are no convincing data to identify high-risk patients, and clinical
1118 judgment has likely kept the highest risk patients, particularly those who were hemodynamically
1119 unstable, from being tested in the current literature.

1120 S-ICD

1121 Patients receiving a non-transvenous ICD system should routinely undergo DT, given there are no
1122 current data regarding the safety and efficacy of not performing DT with this lead configuration and
1123 device.

1124 Conclusion

1125 In providing focused recommendations for ICD programming and defibrillation testing of patients
1126 implanted with a device we have intentionally left many questions unanswered. There are hundreds of
1127 choices for which there is inadequate data to provide evidence or consensus-based
1128 recommendations. This document is a long overdue effort to provide analysis and guidance to the
1129 clinician as to how to make strategic programming choices in the implementation of ICD therapy. The
1130 four continental electrophysiology societies limited the discussion and recommendations to four areas
1131 for which there was sufficient consensus and data. In the review process, clearly articulated opinions
1132 pointed out that additional recommendations are desirable. However, there is an information gap of
1133 insufficient data filled with opinions and logical arguments. Generalizations and inferences were made
1134 from the existing data; e.g. taking data from pacemaker trials and applying to this to ICD bradycardia
1135 programming, logical arguments bridging the differences between primary and secondary prevention
1136 patients for tachycardia detection and therapy, and the use of non-inferiority data to make decisions
1137 about defibrillation testing. This document is a beginning; necessary because there is now sufficient

1138 data to support recommendations that improve the safety, morbidity and mortality of patients with
 1139 ICDs.

1140

1141 **Table 4.** Defibrillation Testing

Study (n)	Patients (DT/No DT)	Results and remarks
CREDIT ¹⁹³ (361) Prospective Multicenter Registry	64% / 36%	<p>More frequent DT for new implants vs generator replacements (71% vs 32%, $P = .0001$), DT for primary and secondary prevention indications (64% vs 63%, $P = NS$).</p> <p>Reasons for no DT were as follows: unnecessary (44%); persistent atrial fibrillation (37%); no anesthetist (20%); and patient or physician preference (6%).</p> <p>DT was not performed in a third of ICD implants, usually due to a perceived lack of need or relative contraindication.</p> <p>Non-consecutive patients, single manufacturer.</p>
Ontario DT Registry ²²⁵ (2173) Prospective Multicenter Registry	PP: 65% / 45% SP: 67% / 43% GR: 24% / 76%	<p>Multivariate predictors for DT included new ICD implant (OR: 13.9; $P < .0001$), DCM (OR: 1.8; $P < .0001$), amiodarone (OR: 1.5; $P = .004$), and LVEF >20% (OR: 1.3; $P = .05$). History of AF (OR: 0.58; $P = .0001$) or OAC use (OR: 0.75; $P = .03$) was associated with a lower likelihood of having DT. Complications, including death, were similar: DT 8.7% vs no DT 8.3% ($P = .7$).</p> <p>All consecutive implants at 10 centers in Ontario</p>
NCDR ²²³ (64,277) Prospective Multicenter Registry	71% / 29%	<p>No DT; older, higher incidence of HF, lower LVEF, atrial arrhythmias, and a primary prevention indication; hospital adverse events; DT 2.56% vs. 3.58% no DT ($P < .001$). Death or any no DT complication (OR [95%CI] 1.46 [1.33–1.61]; $P < .001$), DT is not performed on many (29%) patients in clinical practice.</p>

		Generator replacement excluded.
Israel DFT Registry ²²⁶ (3596) Prospective Multicenter Registry	17% / 83%	<p>Variables associated with ICD testing: implantation for secondary prevention (relative risk [RR] 1.87), prior ventricular arrhythmias (RR 1.81), use of AADs (RR 1.59), and sinus rhythm (RR 2.05). No significant differences in the incidence of mortality, malignant ventricular arrhythmias, or inappropriate ICD discharges were observed between patients who underwent DT compared with those who were not tested.</p> <p>All consecutive implants during 1 year at 22 centers: HOCM: 6.2% DT, 6.3% no DT; ARVC: 0.6% DT, 0.5% no DT; congenital heart disease: 0.8% DT, 2.1% no DT; Long QT: 1.2% DT, 0.26% no DT; Brugada syndrome: 0.3% DT, 0.44% no DT; family history cardiac death: 5.3% DT, 4.7% no DT.</p>
SAFE-ICD ²¹⁰ 2120 Prospective Observational Study	836 DT 1284 no DT	<p>Followed up for 24 months. Primary endpoint was composite of severe implant complications, sudden cardiac death, or resuscitation at 2 years.</p> <p>Primary endpoint: Of 34 patients, 12 intraoperative complications (8 in DT; 4 in no DT) and 22 during follow-up (10 in DT; 12 in no DT). Estimated yearly incidence: DT 1.15% (0.73 to 1.83) and no DT 0.68% (0.42 to 1.12); no difference.</p> <p>In 41 Italian centers. The only exclusion criterion was refusal to provide consent. Other ICD indications: 15% DT, 12% no DT.</p>

<p>Healey JS, et al²¹²</p> <p>(145)</p> <p>Randomized Multicenter Subgroup Study</p>	<p>75 DT</p> <p>70 no DT</p>	<p>All patients in DT arm achieved a successful DT (≤ 25 J); 96% without requiring any system modification. No patient experienced perioperative stroke, myocardial infarction, HF, intubation, or unplanned ICU stay. The composite of HF hospitalization or all-cause mortality occurred in 10% of no DT vs. 19% of the DT arm (HR: 0.53; 95% CI 0.21–1.31; $P = .14$).</p> <p>Conclusions: Perioperative complications, failed appropriate shocks, and arrhythmic death are uncommon regardless of DT. There was a nonsignificant increase in the risk of death or HF hospitalization with DT.</p> <p>Excluded: intracardiac thrombus, persistent or permanent AF without appropriate anticoagulation, right-sided implant, or felt ineligible for DT.</p>
<p>SIMPLE²¹⁵</p> <p>2500</p> <p>Randomized Multicenter Trial</p>	<p>1253 DT</p> <p>1247 no DT</p>	<p>Primary outcome: arrhythmic death or failed appropriate shock occurred in fewer patients (90 [7% per year]) in no DT vs DT (104 [8% per year]; HR: 0.86; 95% CI 0.65–1.14; P non-inferiority $<.0001$). The first safety composite outcome occurred in 69 (5.6%) of 1236 patients with no DT and in 81 (6.5%) of 1242 patients with DT, $P = .33$. The second safety composite outcome, including only events most likely to be directly caused by DT, occurred in 3.2% of patients without DT vs 4.5% with DT, $P = .08$.</p> <p>Routine DT at the time of ICD implantation is generally well tolerated, but does not improve shock efficacy or reduce arrhythmic death.</p> <p>Single manufacturer, excluded patients on active transplantation list, ICD expected to be right-sided implant. HOCM: 4.2% DT, 3.4% no DT; long QT, Brugada syndrome, or catecholaminergic polymorphic VT: 2.3%</p>

		DT, 1.9% no DT.
NORDIC ICD ²¹⁶ 1077 Randomized Multicenter Trial	540 DT 537 no DT	<p>ICD shocks were programmed to 40 J in all patients.</p> <p>Primary end point: First shock efficacy for all true VT and fibrillation episodes during 22.8 months of follow-up. Noninferior with or without DT. First shock efficacy 3.0% in favor of no DT. A total of 112 procedure-related serious adverse events occurred within 30 days in 94 DT patients (17.6%) and 89 events in 74 no-DT patients (13.9%).</p> <p>Excluded were the following: survived an episode of VF due to acute ischemia or potentially reversible causes, listed for heart transplant, life expectancy less than the study duration due to malignant conditions, terminal renal insufficiency, any conditions precluding DT (e.g., left atrial or ventricular thrombus), pre-existing or previous ICD or CRT-D, or if the device was intended to be implanted on the right side.</p>

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Intraprocedural Testing of Defibrillation efficacy Recommendations	Class of Recommendation	Level of Evidence
Defibrillation efficacy testing is recommended in patients undergoing a subcutaneous ICD implantation.	I	C-LD
It is reasonable to omit defibrillation efficacy testing in patients undergoing initial left-pectoral transvenous ICD implantation procedures where appropriate sensing, pacing, and impedance values are obtained with fluoroscopically well-positioned RV leads.	Ila	B-R
Defibrillation efficacy testing is reasonable in patients undergoing a right-pectoral transvenous ICD implantation or	Ila	B-NR

ICD pulse generator changes.		
Defibrillation efficacy testing at the time of implantation of a transvenous ICD should not be performed on patients with a documented nonchronic cardiac thrombus, atrial fibrillation or atrial flutter without adequate systemic anticoagulation, critical aortic stenosis, unstable CAD, recent stroke or TIA, hemodynamic instability, or other known morbidities associated with poor outcomes.	III (Harm)	C-LD

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Number Value: 0 = \$0; 1 = < \$10,000; 2 = > \$10,000 to < \$25,000; 3 = > \$25,000 to < \$50,000; 4 = > \$50,000 to < \$100,000; 5 = > \$100,000

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