Decision Memo for Implantable Cardioverter Defibrillators (CAG-00157R4)

A. The Centers for Medicare & Medicaid Services (CMS) has determined that the evidence is sufficient to
conclude that the use of implantable cardioverter defibrillators (ICDs, also referred to as defibrillators) is
reasonable and necessary for the treatment of illness or injury or to improve the functioning of a
malformed body member under section 1862(a)(1)(A) of the Social Security Act.

CMS is finalizing relatively minimal changes to the ICD NCD from the 2005 reconsideration. We
summarize the changes below and fully explain the changes in the Analysis section of the NCD decision
memo.

○ Patient Criteria
  ▪ Add cardiac magnetic resonance imaging (MRI) to the list of diagnostic imaging studies that
can evaluate left ventricular ejection fraction (LVEF);
  ▪ Require patients who have severe non-ischemic dilated cardiomyopathy but no personal
history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular
fibrillation to have been on optimal medical therapy (OMT) for at least 3 months;
  ▪ Require a patient shared decision making (SDM) interaction prior to ICD implantation for
certain patients.

○ Additional Patient Criteria
  ▪ Remove the Class IV heart failure requirement for cardiac resynchronization therapy (CRT).

○ Exceptions to Waiting Periods
  ▪ Add an exception for patients meeting CMS coverage requirements for cardiac pacemakers,
and who meet the criteria for an ICD;
  ▪ Add an exception for patients with an existing ICD and qualifying replacement.

○ Registry Requirement
  ▪ End the data collection requirement.

We are finalizing changes to the 20.4 NCD that reflect the 2005 reconsideration as described below:

B. Covered Indications

  1. Patients with a personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to
ventricular fibrillation. Patients must have demonstrated:
- An episode of sustained ventricular tachyarrhythmia, either spontaneous or induced by an electrophysiology (EP) study, not associated with an acute myocardial infarction and not due to a transient or reversible cause; or
- An episode of cardiac arrest due to ventricular fibrillation, not due to a transient or reversible cause.

2. Patients with a prior myocardial infarction and a measured left ventricular ejection fraction (LVEF) ≤ 0.30. Patients must not have:
   - New York Heart Association (NYHA) classification IV heart failure;
   - Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
   - Had a myocardial infarction within the past 40 days; or
   - Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B2, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

3. Patients who have severe ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, and have New York Heart Association (NYHA) Class II or III heart failure, left ventricular ejection fraction (LVEF) ≤ 35%. Additionally, patients must not have:
   - Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
   - Had a myocardial infarction within the past 40 days; or
   - Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B3, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

4. Patients who have severe non-ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, and have New York Heart Association (NYHA) Class II or III heart failure, left ventricular ejection fraction (LVEF) ≤ 35%, been on optimal medical therapy (OMT) for at least 3 months. Additionally, patients must not have:
- Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
- Had a myocardial infarction within the past 40 days; or
- Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B4, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

5. Patients with documented familial, or genetic disorders with a high risk of life-threatening tachyarrhythmias (sustained ventricular tachycardia or ventricular fibrillation), to include, but not limited to, long QT syndrome or hypertrophic cardiomyopathy.

For these patients identified in B5, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

6. Patients with an existing ICD may receive an ICD replacement if it is required due to the end of battery life, elective replacement indicator (ERI) or device/lead malfunction.

For each of these groups listed above, the following additional criteria must also be met:

1. Patients must be clinically stable (e.g., not in shock, from any etiology);
2. Left ventricular ejection fraction (LVEF) must be measured by echocardiography, radionuclide (nuclear medicine) imaging, cardiac magnetic resonance imaging (MRI), or catheter angiography;
3. Patients must not have:
   - Significant, irreversible brain damage; or
   - Any disease, other than cardiac disease (e.g., cancer, renal failure, liver failure) associated with a likelihood of survival less than 1 year; or
   - Supraventricular tachycardia such as atrial fibrillation with a poorly controlled ventricular rate.
C. Exceptions to waiting periods for patients that have had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months, or had a myocardial infarction within the past 40 days:

Cardiac Pacemakers: Patients who meet all CMS coverage requirements for cardiac pacemakers and who meet the criteria in this national coverage determination for an ICD may receive the combined device in one procedure at the time the pacemaker is clinically indicated;

Replacement of ICDs: Patients with an existing ICD may receive a ICD replacement if it is required due to the end of battery life, elective replacement indicator (ERI) or device/lead malfunction.

D. Other Indications:

For patients who are candidates for heart transplantation on the United Network for Organ Sharing (UNOS) transplant list awaiting a donor heart, coverage of ICDs, as with cardiac resynchronization therapy, as a bridge to transplant to prolong survival until a donor becomes available is determined by the local Medicare Administrative Contractors (MACs).

All other indications for ICDs not currently covered in accordance with this decision may be covered under Category B IDE trials (42 CFR 405.201).

See Appendix B for the NCD manual language.

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**Decision Memo**

**TO:** Administrative File: CAG-00157R4

**FROM:** Tamara Syrek Jensen, JD
Director, Coverage and Analysis Group

Joseph Chin, MD, MS
Deputy Director, Coverage and Analysis Group

Lori Ashby, MA
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David Dolan, MBA, MA
Lead Analyst

Printed on 2/15/2018. Page 4 of 77
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- **Patient Criteria**
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- **Additional Patient Criteria**
  - Remove the Class IV heart failure requirement for cardiac resynchronization therapy (CRT).

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  - End the data collection requirement.

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1. Patients must be clinically stable (e.g., not in shock, from any etiology);
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All other indications for ICDs not currently covered in accordance with this decision may be covered under Category B IDE trials (42 CFR 405.201).

See Appendix B for the NCD manual language.

II. Background

Throughout this document we use numerous acronyms, some of which are not defined as they are presented in direct quotations. Please find below a list of these acronyms and corresponding full terminology:

AAD – Antiarrhythmic Drug
ACC – American College of Cardiology
ACE – Angiotensin-Converting Enzyme
AF – Atrial Fibrillation
ARB – Angiotensin II Receptor Blocker
CABG – Coronary Artery Bypass Graft
CAD – Coronary Artery Disease
CED – Coverage with Evidence Development
CI – Confidence Interval
CMS – Centers for Medicare & Medicaid Services
CRT – Cardiac Resynchronization Therapy
CRT-D – Cardiac Resynchronization Therapy, Defibrillator
CRT-P – Cardiac Resynchronization Therapy, Pacemaker
ECG – Electrocardiogram
FDA – Food and Drug Administration
HF – Heart Failure
HR – Hazard Ratio
HRS – Heart Rhythm Society
ICD – Implantable Cardioverter Defibrillator
ITT – Intention to Treat
LBBB – Left Bundle Branch Block
LV – Left Ventricle / Left Ventricular
LVEF – Left Ventricular Ejection Fraction
LVSD – Left Ventricular Systolic Dysfunction
MI – Myocardial Infarction
NCA – National Coverage Analysis
CD – National Coverage Determination
NYHA – New York Heart Association
OMT – Optimal Medical Therapy
CMS initiated this national coverage determination (NCD) to consider coverage under the Medicare Program for implantable cardioverter defibrillators (ICDs; often referred to as “defibrillators”).

The scope of this review is limited to ICDs. While we reference cardiac resynchronization therapy defibrillator (CRT-D) devices in this document since these devices have defibrillator functions, CRT devices are outside the scope of this decision. Coverage determination under section 1862(a)(1)(A) for CRT devices are currently made by local Medicare contractors and are not currently subject to an NCD.

### Sudden Cardiac Death

Sudden cardiac death (SCD) is defined as the sudden and unexpected death from cardiac causes within an hour of the onset of symptoms (Bonow, Mann, Zipes, & Libby, 2012). Heart disease is the leading cause of death in the US and accounts for approximately 24% of all deaths, and SCD in turn is estimated to be the final cause of approximately 50% of all cardiac deaths (Centers for Disease Control and Prevention [CDC], 2017; Hayashi, Shimizu, & Albert, 2015). Stecker et al. (2014) summarize the “rule of 50’s” for SCD: it accounts for up to 50% of all cardiac deaths; 50% of the SCD’s are the first cardiac event; and SCD accounts for up to 50% of potentially productive years of life lost due to premature death or disabilities.

Sudden cardiac arrest is defined as the “sudden cessation of effective cardiac mechanical activity resulting in unresponsiveness, without normal breathing or signs of circulation” (Russo et al., 2013). If not resuscitated, patients with cardiac arrest will progress to sudden cardiac death. Sudden cardiac arrest and death in turn are most often associated with life-threatening ventricular tachyarrhythmias (rapid heart rates arising from the ventricles): sustained ventricular tachycardia (VT) or ventricular fibrillation (VF). Sustained VT may degenerate into VF and then asystole (in which all cardiac electrical activity stops, and the heart stops beating).

Sustained VT is defined as VT lasting ≥30 seconds or terminated by medical intervention (either cardioversion or pacing) before that time (Russo et al., 2013). While tachycardia is defined technically as a heart rate of ≥100 beats per minute, many experts believe that in hemodynamically unstable patients, “if the heart rate is <150 beats per minute, it is unlikely that the symptoms of instability are caused primarily by the tachycardia unless there is impaired ventricular function” (i.e., a depressed left ventricular ejection fraction). Thus, “a heart rate
>150 beats per minute is usually an inappropriate response to physiological stress (e.g., fever, dehydration) or other underlying conditions” and in unstable patients, this higher heart rate is likely a major contributor or the primary cause of the symptoms of instability (American Heart Association, 2016a; American Heart Association, 2016b).

## Dilated Cardiomyopathy

In cardiomyopathy, the heart muscle becomes enlarged, thick, or rigid. In the vast majority of cases, the underlying cardiomyopathy that increases SCD risk is due to ischemia (coronary artery disease causing blockages of the arteries which reduces blood flow and oxygen to the heart muscle, which in turn both decreases the heart’s ability to pump blood and oxygen to the vital organs (e.g., brain, kidneys), and increases the risk of life-threatening cardiac arrhythmias); we refer to this as ischemic cardiomyopathy.

The underlying cardiomyopathy may also be due to non-ischemic causes, such as infection, inflammation, familial or genetic conditions or idiopathic causes; we generally refer to this as non-ischemic cardiomyopathy, although in this document we break out familial or genetic etiologies into a separate SCD primary prevention risk category. The two major categories of cardiomyopathy – ischemic and non-ischemic – are not mutually exclusive; some patients may have elements of both.

## Heart Failure

Heart failure can affect the right side of the heart only, or it can affect both sides of the heart. Most cases involve both sides of the heart and in turn is a clinical diagnosis that is associated with a depressed left ventricular ejection fraction (LVEF) as measured by diagnostic imaging studies (echocardiography, radionuclide scanning, cardiac magnetic resonance imaging (MRI), or catheter angiography). Most trials on prevention of sudden cardiac death have used a LVEF ≤35% in their inclusion criteria. The New York Heart Association (NYHA) Functional Classification is the most commonly used classification system in both clinical practice and trial inclusion criteria (see Appendix D). It classifies the severity of heart failure in terms of symptoms (such as shortness of breath) and limitations in physical activity (e.g., Class I patients have no symptoms or limitations with ordinary activity; Class IV patients have symptoms at rest, and severe limitations in physical activity).

## Implantable Cardioverter Defibrillators

An ICD is a battery-driven electronic device placed in the chest, and connected to the heart by leads. The device monitors the heart’s electrical activity, detects the onset of life-threatening tachyarrhythmias, and tries to terminate these first by smaller electrical stimuli, and then with shock therapy. The “shock” is a defibrillation – a defibrillation.
delivery of an adequate amount of electrical current with the goal of temporarily stopping all cardiac electrical
signals (essentially producing a temporary state of asystole), which then allows the heart to “reset” itself back to
a normal sinus rhythm.

These shocks can be effective in terminating tachyarrhythmias but can be quite painful and disruptive to the
patient. Shocks are said to be “appropriate” if they are in response to true, life-threatening tachyarrhythmias;
“inappropriate” shocks are those triggered by arrhythmias that are not potentially life-threatening. Inappropriate
shocks can also be triggered by malfunction of the device/leads. Inappropriate shocks have been reported in 6% -20% of patients, and in 5.9% of patients in the 2016 DANISH trial (Køber et al., 2016; Priori et al., 2015;
Providência et al., 2015; van der Heijden et al., 2015). At the same time, defibrillators never actually “fire” (no
therapeutic shocks are delivered) in the majority of patients who receive them for primary prevention (Merchant,
Quest, Leon, & El-Chami, 2016; Priori et al., 2015).

ICDs can only treat arrest and life-threatening ventricular tachyarrhythmias, and thus their ability to reduce all-
cause mortality (i.e., improve patient survivability) depends on a relatively high frequency of arrhythmic deaths
compared to death by other causes, for any given population being treated with ICDs.

Adverse Events

Patients with ICDs [whether alone or combined with a cardiac resynchronization therapy device (CRT)] require
constant physician follow up typically for the rest of their lives, to include regular device interrogations every 3-6
months. Adverse events can include, in addition to inappropriate shocks, procedure-related pneumothorax
(collection of air in the chest cavity which causes the lung to collapse) and cardiac tamponade (pressure on the
heart that occurs due to the accumulation of fluid in the pericardial space), and device-related infections – all of
which are potentially life-threatening. Device infections have been reported previously in 1-7% of patients, and
the more recent DANISH trial observed rates of 4.7% in the CRT-D group and 5.1% in the ICD group (Køber et
al, 2016; Mulpuru, Pretorius, & Birgersdotter-Green, 2013; van der Heijden, 2015). In addition to being life-
threatening, such adverse events require further treatment, hospitalizations, imaging and procedures. When
battery life expires (typically within 5-10 years) a procedural replacement of the device itself is necessary.
Patients with defibrillators often live with anxiety and depressive symptoms due to fear of either a shock or
device failure (Freedenberg, Thomas, & Friedmann, 2011). As with any medical intervention, the question of
whether the demonstrated benefits outweigh the harms for particular patients persists.

III. History of Medicare Coverage

CMS issued an NCD in 1986 providing limited coverage of implantable defibrillators. The policy has expanded
coverage was expanded to patients with a previous myocardial infarction, low ejection fraction and a wide QRS
interval. The policy was also expanded to include coverage to patients enrolled in an Investigational Device
Exemption Category B device trial. A follow up decision memorandum to clarify this specific aspect of the policy
was published March 12, 2004.
On March 30, 2004, CMS accepted a request from Medtronic Inc. to expand coverage for ICDs. Medtronic Inc. made this request based on the results of the Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT) and specifically requested that Medicare expand coverage to the trial population. On December 16, 2004, CMS announced its concern on the absence of publication of the SCD-HeFT data and the potential for closing the NCD. Publication did not occur prior to the December 28, 2004 deadline so on that day CMS posted a final DM that continued current coverage. CMS opened a reconsideration of that decision on December 29, 2004 in anticipation of the SCD-HeFT publication, and a decision based on SCD-HeFT data was finalized on January 27, 2005.

Since there is an existing NCD for ICDs, this review is a reconsideration of the current policy. The current policy is codified in 20.4 of the Medicare National Coverage Determinations manual. Section 20.4 of the NCD Manual has been included at Appendix C.

A. Current Request

CMS opened this NCA to reconsider coverage indications for ICDs.

B. Benefit Category

Medicare is a defined benefit program. For an item or service to be covered by the Medicare program, it must fall within one of the statutorily defined benefit categories outlined in the Social Security Act.

Implantable cardioverter defibrillators qualify as:

- Prosthetic devices.

IV. Timeline of Recent Activities

<table>
<thead>
<tr>
<th>Date</th>
<th>Action</th>
</tr>
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| May 30, 2017 | CMS posts a tracking sheet announcing the opening of the NCA.  
The initial 30-day public comment period begins. |
<table>
<thead>
<tr>
<th>Date</th>
<th>Action</th>
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<tbody>
<tr>
<td>June 29, 2017</td>
<td>First public comment period ends. CMS receives 36 comments.</td>
</tr>
<tr>
<td>November 20, 2017</td>
<td>Proposed Decision Memorandum posted. 30-day public comment period begins.</td>
</tr>
<tr>
<td>December 20, 2017</td>
<td>30-day public comment period ends. CMS receives 24 comments.</td>
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**V. Food and Drug Administration (FDA) Status**

The FDA approved the first implantable defibrillator in 1985 while the first implantable cardioverter defibrillators were approved in 1988 and 1989. The FDA approves each device individually and has granted premarket approvals (PMA) for implantable defibrillators for the indications of providing antitachycardia pacing and ventricular defibrillation for automated treatment of life-threatening ventricular arrhythmias.

On September 28, 2012, the FDA approved the first subcutaneous implantable defibrillator, the EMBLEM S-ICD system by Boston Scientific. This device is indicated to provide defibrillation therapy for the treatment of life-threatening ventricular tachyarrhythmias in patients who do not have symptomatic bradycardia, incessant ventricular tachycardia, or spontaneous, frequently recurring ventricular tachycardia that is reliably terminated with antitachycardia pacing.


**VI. General Methodological Principles**

When making national coverage determinations, CMS generally evaluates relevant clinical evidence to determine whether or not the evidence is of sufficient quality to support a finding that an item or service falling within a benefit category is reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body member. The critical appraisal of the evidence enables us to determine to what degree we are confident that: 1) the specific assessment questions can be answered conclusively; and 2) the intervention will improve health outcomes for beneficiaries. An improved health outcome is one of several considerations in determining whether an item or service is reasonable and necessary.
A detailed account of the methodological principles of study design that the Agency utilizes to assess the relevant literature on a therapeutic or diagnostic item or service for specific conditions can be found in Appendix A.

Public comments sometimes cite published clinical evidence and give CMS useful information. Public comments that give information on unpublished evidence such as the results of individual practitioners or patients are less rigorous and therefore less useful for making a coverage determination. Public comments that contain personal health information will be redacted or will not be made available on the CMS website CMS responds in detail to the public comments on a proposed national coverage determination when issuing the final national coverage determination.

VII. Evidence

A. Introduction

CMS last reconsidered the ICD NCD (see Appendix C for § 20.4 of the NCD) in January of 2005. CMS has opened this national coverage analysis (NCA) to reconsider coverage indications for ICDs based on more recent scientific evidence and will discuss in this section relevant evidence established after that date.

For this reconsideration, we reviewed the published medical literature since 2005 to 2017 to determine reasonable and necessary indications for ICDs and whether the data collection questions have been answered.

B. Discussion of Evidence

1. Evidence Question(s)

Our review and analysis of the evidence on the clinical utility of ICDs in symptomatic patients who have not experienced a prior episode of cardiac arrest or sustained ventricular tachyarrhythmias (sustained ventricular tachycardia or ventricular fibrillation), and thus whether ICDs are reasonable and necessary to treat certain Medicare patients, is guided by the following questions:

• Is there evidence to conclude that ICDs decrease mortality for patients with ischemic dilated cardiomyopathy (IDCM) and reduced LVEF?
Is there evidence to conclude that ICDs decrease mortality for patients with non-ischemic dilated cardiomyopathy (NIDCM) and reduced LVEF?

2. External Technology Assessments

While CMS did not request an external technology assessment (TA) as part of this reconsideration, a TA had been requested prior to the opening of this NCA (2013) that assessed the effectiveness of defibrillators.


The authors, funded by the US Agency for Healthcare Research and Quality, performed a methodical technology assessment to include a systematic literature review and meta-analysis to assess the clinical effectiveness of ICD use for primary prevention of sudden cardiac death. The assessment focused on three key questions, examining:

- ICD versus no ICD, ICD with antitachycardia pacing versus ICD alone, or ICD with CRT versus ICD alone, and differences among subgroups;
- Early and late adverse events and inappropriate shocks after ICD implantation, and differences among subgroups; and
- Eligibility criteria and evaluation methods for patients included in comparative studies and the risk of SCD.

The assessment did not examine the possible added benefit of ICD in patients who are to receive CRT (i.e., CRT-D v. CRT-P).


Uhlig et al. Assessment on Implantable Defibrillators and the Evidence for Primary Prevention of Sudden Cardiac Death. 2013. Table 4, Page 27.
Uhlig and colleagues concluded that “there is a high strength of evidence that ICD therapy for primary prevention of SCD, versus no ICD therapy, shows benefit with regard to all-cause mortality and SCD in patients with reduced LVEF and ischemic or non-ischemic cardiomyopathy beyond the immediate post-MI or coronary revascularization periods. Studies failed to show statistically significant differences for all-cause mortality across subgroups. There is insufficient evidence for all-cause mortality for patients who receive CRT-Ds versus ICD alone for primary prevention. There is high strength of evidence that in-hospital adverse events are infrequent (1-3%) and moderate strength of evidence that up to one-fifth of patients receive inappropriate shocks from the ICDs.”

3. Internal Technology Assessment

Literature Search Methods

CMS searched Pubmed MEDLINE, Embase, ClinicalTrials.gov, and the Cochrane Central Registry of Controlled Trials, from 2004 through May 2017 (ensuring overlap of the literature reviewed in the 2005 NCD). Search keywords included combinations of: implantable cardioverter defibrillator, ICD, defibrillator, sudden cardiac death, ventricular arrhythmias, ischemic cardiomyopathy, non-ischemic cardiomyopathy, randomized, and cardiac resynchronization therapy. This evidence review primarily focuses on randomized controlled trials that assess the clinical utility of defibrillators compared to optimal medical therapy, and relevant formal Technology Assessments and professional society guidelines. In our analysis we reference other trials, observational studies and analyses, as well as relevant published correspondence and editorials, as these can assist both in interpreting these trials and identifying evidence gaps. Abstracts only and publications in languages other than English were excluded. Trials on CRT only were also excluded.

We reviewed all original research that is detailed in the ACC listing of published manuscripts from the ACC NCDR ICD Registry. We additionally performed a Pubmed search using the terms ICD or defibrillator and NCDR (National Cardiovascular Data Registry) focusing on all (including observational) studies, to assess the extent to which the published literature has addressed the ten “initial hypotheses” for this registry as stated in our 2005 NCD.

Technology Assessments

The authors, funded by the UK National Institute for Health Research, performed a methodical technical assessment to include a systematic literature review and meta-analysis to assess:

- The clinical effectiveness of ICDs for people at increased risk of sudden cardiac death as a result of ventricular arrhythmias despite receiving optimal pharmacological therapy;
- CRT with or without a defibrillator (CRT-D v. CRT-P) in addition to optimal pharmacological therapy for people with heart failure as a result of left ventricular systolic dysfunction (LVSD) and cardiac dyssynchrony; and
- CRT-D in addition to optimal pharmacological therapy for people with both conditions above.

Studies


As to the clinical effectiveness of ICDs, Colquitt and colleagues found that “ICDs reduced all-cause mortality in people at increased risk of SCD, defined in trials as those with previous ventricular arrhythmias/cardiac arrest, myocardial infarction (MI) > 3 weeks previously, non-ischemic cardiomyopathy (depending on data included) or ischemic/non-ischemic heart failure (HF) and left ventricular ejection fraction ≤ 35%. There was no benefit in people scheduled for coronary artery bypass graft. A reduction in SCD but not all-cause mortality was found in people with recent MI. CRT-P and CRT-D reduced mortality and HF hospitalizations, and improved other outcomes, in people with HF as a result of LVSD and cardiac dyssynchrony when compared with optimal pharmacological therapy (OPT). In people with both conditions, CRT-D reduced the risk of all-cause mortality and HF hospitalization, and improved other outcomes, compared with ICDs.”

The authors also concluded that “an RCT comparing CRT-D and CRT-P in people with HF as a result of LVSD and cardiac dyssynchrony is required, for both those with and those without an ICD indication. A RCT is also needed into the benefits of ICD in non-ischemic cardiomyopathy in the absence of dyssynchrony.”

Meta-Analyses


The purpose of this meta-analysis was to “investigate the association of primary prevention ICDs with all-cause
mortality in patients with nonischemic cardiomyopathy.” The authors pooled data from four trials: DEFINITE (2004), CAT (2002), SCD-HeFT (2005) and DANISH (2016); please see the Evidence Table (Figure 1) for summaries of the trials. A fifth trial (and one of four that examined non-ischemic cardiomyopathy patients only), AMIOVIRT (2003), was excluded because the trial compared ICD to an antiarrhythmic medication, amiodarone, rather than to standard optimal medical therapy. The authors included the non-ischemic patient subgroup from SCD-HeFT (the sole trial that included both ischemic and non-ischemic cardiomyopathy patients), and the DANISH subgroup of patients who did not receive a CRT (as CRT patients were excluded generally from the analysis).

Pooling these selected patients from these four trials, and using fixed- and random-effects models, the authors showed that use of an ICD significantly reduced all-cause mortality (HR 0.75; CI 0.61-0.93; P=.008, P=0.87 for heterogeneity). The authors concluded that these findings supported the 2012 AHA/ACCF/HRS guidelines recommending the use of ICDs in certain patients with non-ischemic cardiomyopathy and depressed LVEF.


This meta-analysis also assessed the association of ICDs for primary prevention with all-cause mortality in patients with non-ischemic cardiomyopathy. The authors pooled data from non-ischemic patients from six trials: CAT (2002), AMIOVIRT (2003), DEFINITE (2004), SCD-HeFT (2016), COMPANION (2004), and DANISH (2016). The authors did three separate analyses: combining all of the trials; combining ICD patients only, thus excluding CRT patients, for the five relevant trials; and combining CRT patients only, from the two relevant trials (DANISH and COMPANION). (See the Evidence Table, Figure 1, for summaries of these trials.)

For the first analysis, pooling all six trials, the authors showed that use of an ICD significantly reduced all-cause mortality risk by 23% (HR 0.77; CI 0.64–0.91). The second pooled analysis assessing ICD plus optimal medical therapy versus optimal medical therapy alone showed that use of an ICD significantly reduced all-cause mortality risk by 24% (HR 0.76; CI, 0.62–0.94). The third pooled analysis assessing ICD plus CRT (CRT-D) plus optimal medical therapy versus CRT plus optimal medical therapy alone, showed no significant difference between the two groups (HR 0.70; CI 0.39–1.26); thus no survival benefit when ICD is used in patients in addition to CRT. (P values were not reported.)

The authors concluded that “this incremental reduction of all-cause mortality with ICD is substantial and provides support to the existing [ACC/AHA/HRS] guidelines until we acquire additional data. . . Furthermore, adequately powered randomized studies are needed before recommending any change in existing guidelines . . .”

Randomized Controlled Trials
This section summarizes the randomized controlled trials since the 2005 NCD that assess the clinical utility of defibrillators in the primary prevention of sudden cardiac death. The evidence table (Figure 1) summarizes all major relevant randomized controlled trials; the trials since the 2005 NCD are in italics.

Figure 1: Major Randomized Controlled Trials Assessing Defibrillators for Primary Prevention of Sudden Cardiac Death

<table>
<thead>
<tr>
<th>Year</th>
<th>Trial / Author(s)</th>
<th>Population</th>
<th>Intervention/Comparator</th>
<th>Outcome</th>
<th>Time (mos)</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>1996</td>
<td>MADIT / Moss et al.</td>
<td>n = 196 NYHA 1-2 EF ≤35% Post MI 3 wks NSVT EP induced VT</td>
<td>ICD v AA drug (mostly amiodarone)</td>
<td>ACM</td>
<td>27m</td>
<td>HR 0.46 (95% CI, 0.26-0.82)</td>
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<tr>
<td>1997</td>
<td>CABG Patch (post CABG) / Bigger</td>
<td>n = 900 NYHA 1-3 EF ≤35% Sched for CABG Abnl SAECG</td>
<td>ICD v. OMT (OMT = drugs plus revascular.)</td>
<td>ACM</td>
<td>32m</td>
<td>HR 1.07 (95% CI, 0.81-1.42)</td>
</tr>
<tr>
<td>2002</td>
<td>MADIT II / Moss et al.</td>
<td>n = 1,232 NYHA 1-3 EF ≤30% Post-MI ≥1m, ≥3m if CABG</td>
<td>ICD v OPT</td>
<td>ACM</td>
<td>20m</td>
<td>HR 0.69 (95% CI, 0.51-0.93)</td>
</tr>
<tr>
<td>2004</td>
<td>DINAMIT (early post-MI) / Hohnloser et al.</td>
<td>n = 674 NYHA 1-3 EF ≤35% Post-MI (6-40 days, mean 18) Abnl HR</td>
<td>ICD v OPT</td>
<td>ACM</td>
<td>33m</td>
<td>HR 1.08 (95% CI, 0.76-1.55)</td>
</tr>
<tr>
<td>Year</td>
<td>Trial / Author(s)</td>
<td>Population</td>
<td>Intervention/Comparator</td>
<td>Outcome</td>
<td>Time (mos)</td>
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</table>
| 2009 | IRIS (early post-MI) / Steinbeck et al. | n = 898 Post-MI (5-31 days) AND one of:  
  a. EF ≤40%, HR ≥90bpm  
  b. rapid NSVT  
  c. both a and b | ICD v OPT | ACM | 37m | HR 1.04 (95% CI, 0.81-1.35) |

**Both Ischemic and Non-ischemic CM**

| Year | Trial / Author(s) | Population | Intervention/Comparator | Outcome | Time (mos) | All pts:  
  - POS: CRT-P (v OPT), HR 0.81 (95% CI, 0.69-0.96)  
  - POS: CRT-D (v OPT), HR 0.80 (95% CI, 0.68-0.95)  
  2nd (ACM alone):  
  - NEG: CRT-P, HR 0.76 (95% CI, 0.58-1.01)  
  - POS CRT-D, HR 0.64 (95% CI, 0.48-0.86) |
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<tr>
<td>2004</td>
<td>COMPANION / Bristow et al.</td>
<td>n = 1,520 (ICM = 838 NICM = 682) NYHA 3-4 EF ≤35% QRS ≥120 msec</td>
<td>1:2:2 OPT : CRT-P : CRT-D</td>
<td>ACM or Hosp (all)</td>
<td>12m : 16m : 16m</td>
<td></td>
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<tr>
<td>2005</td>
<td>SCD-HeFT / Bardy et al.</td>
<td>n = 2,521 (ICM = 52% NICM = 48%) NYHA 2-3 EF ≤35% OPT for 3m prior to R</td>
<td>1:1:1 ICD : amiodarone : placebo (baseline OPT for all)</td>
<td>ACM</td>
<td>46m</td>
<td>POS: ICD v placebo, HR 0.73 (95% CI, 0.52-1.02)</td>
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</table>

*Non-ischemic CM*
<table>
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<tr>
<th>Year</th>
<th>Trial / Author(s)</th>
<th>Population</th>
<th>Intervention/Comparator</th>
<th>Outcome</th>
<th>Time (mos)</th>
<th>Result</th>
</tr>
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<tbody>
<tr>
<td>2002</td>
<td>CAT / Bansch et al.</td>
<td>n = 104 NYHA 2-3 EF ≤30% New onset CM (≤9m)</td>
<td>ICD v OPT</td>
<td>ACM</td>
<td>66m</td>
<td>HR 0.81 (95% CI, 0.38-1.71)</td>
</tr>
<tr>
<td>2003</td>
<td>AMIOVIRT / Strickberger et al.</td>
<td>n = 103 NYHA 1-3 EF ≤35% Asymptomatic NSVT</td>
<td>ICD v. amiodarone [?baseline OPT]</td>
<td>ACM</td>
<td>36m</td>
<td>HR 0.87 (95% CI, 0.31-2.42)</td>
</tr>
<tr>
<td>2004</td>
<td>DEFINITE / Kadish et al.</td>
<td>n = 458 NYHA 1-3 EF ≤35% NSVT</td>
<td>ICD v OPT</td>
<td>ACM</td>
<td>29m</td>
<td>HR 0.65 (95% CI, 0.40-1.06)</td>
</tr>
<tr>
<td>2016</td>
<td>DANISH / Køber et al.</td>
<td>n = 1116 NYHA 2-3 (or 4 if CRT planned) EF ≤35% NT-proBNP &gt;200 pg/mL</td>
<td>ICD or CRT-D + OPT v OMT alone (OPT + CRT if needed)</td>
<td>ACM</td>
<td>68m</td>
<td>CRT: HR 0.91 (95% CI: 0.64-1.29) No CRT: HR 0.83 (95% CI: 0.58-1.19)</td>
</tr>
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</table>


Note that the Multicenter Unsustained Tachycardia Trial (MUSTT), which was initiated in 1989 and reported in 1999, is not included in the analysis since it did not randomize to drug therapy or ICD (Buxton et al., 1999). The 2014 Technology Assessment by Colquitt et al. explicitly excluded this trial (for further critique of that trial, see Colquitt et al., 2014).

The aim of this trial (IRIS) was to evaluate whether patients who met guideline criteria for an ICD had improved survival when the ICD was implanted early, within 40 days after a myocardial infarction (MI), compared to patients who received optimal medical therapy alone in the early phase. Existing guidelines recommended use of an ICD for such patients, but only > 40 days after MI, as the 2004 DINAMIT trial showed no ICD survival benefit with implantation within 6-40 days after MI.

This was a multicenter, open-label, investigator initiated, randomized controlled trial conducted at 80 centers in Germany and other European countries. A total of 898 patients (86% of whom were still in the hospital) were randomized to ICD (n=445) or to individualized, optimal medical therapy alone (n=453) at a mean (±SD) of 13±7 days after infarction. Randomization was stratified to ensure a balanced number of patients with ST elevated and non-ST elevated myocardial infarction between the ICD and control groups.

The primary outcome was time to all-cause mortality post-acute-MI and major secondary outcomes included SCD, non-SCD, and noncardiac deaths. Analysis was intention-to-treat and used time-to-event statistical methods.

Trial inclusion criteria focused on clinically stable patients with either depressed left ventricular ejection fraction (< 40%), documented nonsustained ventricular tachycardia, or both. Exclusion criteria focused on patients with life-threatening ventricular arrhythmias requiring treatment, insufficient documentation of acute MI, or an indication for coronary revascularization before study enrollment.

Patient demographics included median age 63±11 years in the ICD group and 62±11 years in the control group, and 78% male in the ICD group and 76% male in the control group. Other baseline patient characteristics were balanced, with the exception that diabetes and left bundle-branch block were slightly more frequent in the ICD group (P=0.03 and P=0.05, respectively). Greater than 90% of patients were discharged on appropriate post-MI medications. Patients were followed for a median of 37 months.

Trial results demonstrated that prophylactic, early implantation of an ICD (within 40 days after acute MI) does not reduce all-cause mortality (hazard ratio [HR], 1.04; 95% confidence interval [CI], 0.81 to 1.35; P=0.78). There were fewer SCDs in the ICD group than in the control group (27 vs. 60; HR, 0.55; 95% CI, 0.31 to 1.00; P=0.049), but the number of non-SCDs was higher (68 vs. 39; HR, 1.92; 95% CI, 1.29 to 2.84; P=0.001). Thus, the risk of SCD was reduced with ICD therapy, but was offset by an increase in the risk of non-SCD. Prespecified subgroup analysis suggested that the results did not depend on which of the enrollment criteria patients met.

The authors concluded that “prophylactic ICD therapy did not reduce overall mortality among patients with acute myocardial infarction and clinical features that placed them at increased risk.” And thus, “with different risk markers, generating a larger population at risk, and a longer follow-up period” their trial “confirmed the results of DINAMIT.”
The aim of this trial (DANISH) was to evaluate whether ICDs improve survival for patients with nonischemic cardiomyopathy and systolic heart failure, compared to usual care consisting of optimal medication therapy (OMT) and cardiac resynchronization therapy (CRT) for those with an indication for CRT. The investigators stated that clinical uncertainty remained as the largest trial focused on nonischemic patients alone showed no ICD benefit, while a second trial did show an ICD benefit but nonischemics were only one of two subgroups.

This was a multicenter, open-label, investigator-initiated, randomized controlled trial conducted at all five centers where ICDs were implanted in Denmark. A total of 1,116 patients were randomized to ICD or standard of care therapy into two stratum – with 471 in the no-CRT stratum (234 ICD and 237 OMT) and 645 in the CRT stratum (322 CRT-D, a CRT / ICD combination device, and 323 CRT alone). All trial patients received baseline OMT, and 58% received CRT. Randomization was stratified by center and by patients scheduled to receive CRT, to ensure balance between control and intervention arms. The primary outcome was time to all-cause mortality. Secondary outcomes were cardiac death, SCD, resuscitated cardiac arrest or sustained VT, and change from baseline in quality of life. Analysis was intention-to-treat and used time-to-event statistical methods. Prespecified subgroup analyses included age and CRT status.

The trial included symptomatic patients (NYHA class II or III, or NYHA class IV if CRT was planned) with nonischemic systolic heart failure (left ventricular ejection fraction ≤35%, not due to coronary heart disease) and an increased level (>200 pg/ml) of N-terminal pro–brain natriuretic peptide (NT-proBNP). Major exclusion criteria were permanent atrial fibrillation with a resting heart rate higher than 100 beats per minute or renal failure that was being treated with dialysis.

Patient demographics included median age 64±8 years in the ICD group and 63±7 years in the control group, and 73% male in the ICD group and 72% male in the control group. Other baseline patient characteristics were balanced, with the exception that in the no-CRT stratum, the ICD group had a significantly higher baseline NT-proBNP (1,277 pg/ml versus 862 pg/mL). Among all patients with wide left bundle-branch block (QRS duration ≥150 msec), 93% received CRT. As baseline optimal medication therapy, almost all patients received beta-blockers and inhibitors of the renin-angiotensin system and 60% received mineralocorticoid-receptor antagonists. Patients were followed for a median of 68 months.

Trial results demonstrated that ICDs did not improve survival for patients with symptomatic, nonischemic systolic heart failure compared to usual clinical care (HR 0.87; 95% CI 0.68 to 1.12; P=0.28). Neither was there a benefit for the secondary outcome of all CV deaths (HR 0.77, 95% CI 0.57 to 1.05, P=0.10). However, the risk of SCD specifically (also a secondary outcome) was halved with an ICD (HR 0.5; 95% CI 0.31 to 0.82; P=0.005). The results were independent of whether a patient received a CRT device, as there was no significant interaction in the prespecified subgroup analysis for CRT. The prespecified subgroup analysis for age in turn suggested that younger patients (<59 years) may have an overall survival benefit with an ICD (HR 0.51, CI 0.29–0.92, p=0.02). While the overall trial p-value for NT-proBNP was p=0.06, the stratum-specific p-value for patients with NT-proBNP <1177 pg/ml was p=0.02 (HR, 0.59, 95% CI 0.38–0.91), suggesting an ICD benefit for this subgroup. No other prespecified subgroup analysis demonstrated a treatment-by-subgroup interaction.
The authors concluded that “prophylactic ICD implantation in patients with symptomatic systolic heart failure not caused by coronary artery disease was not associated with a significantly lower long-term rate of death from any cause than was usual clinical care.”

Observational Studies Using the NCDR

The eight studies reviewed below all used the NCDR prominently in their analyses. These studies were not specifically designed to target a particular “initial hypothesis” for the registry as identified by CMS in 2005; nor were protocols for these studies submitted to or approved by CMS. However, we believe the research questions for each of these studies below are related to one or more of these ten hypotheses; and collectively, these eight studies are related to all ten hypotheses. We are aware that there are numerous other published studies that may be related to one or more of these hypotheses, or aspects of them.

We add the ten “initial hypotheses” for the registry again below for readers’ convenience.

Initial hypotheses that were to be addressed by the NCDR database included the following:

1. The clinical characteristics of the patients receiving ICDs are similar to those of patients involved in the primary prevention randomized clinical trials.
2. The indications for ICD implantation in patients are similar to those in the primary prevention randomized clinical trials.
3. The in-hospital procedure related complications for patients are similar to those in the primary prevention randomized clinical trials.
4. Certified providers competent in ICD implantation are implanting ICD devices in patients.
5. Patients who receive an ICD represent patients for which current clinical guidelines and the evidence base recommend implantation.
6. The clinical characteristics and indications for ICD implantation do not differ significantly among facilities.
7. The clinical characteristics and indications for ICD implantation do not differ significantly among providers.
8. The in-hospital procedure related complications for ICD implantation do not differ significantly among facilities.
9. The in-hospital procedure related complications for ICD implantation do not differ significantly among providers.
10. The in-hospital procedure related complications for ICD implantation do not differ significantly among device manufacturer, types, and/or programming.

Al-Khatib SM1, Hellkamp A, Bardy GH. et al. Survival of patients receiving a primary prevention implantable cardioverter-defibrillator in clinical practice vs clinical trials JAMA. 2013 Jan 2;309(1):55-62
The aim of this study was to determine whether trial-eligible patients receiving a primary prevention ICD have an all-cause mortality rate that differs from the two largest primary prevention clinical trials MADIT-II and SCD-HeFT.

This was a retrospective analysis comparing patients from the NCDR ICD Registry who met MADIT-II or SCD-HeFT study criteria to the respective patients from those clinical trials. The registry was queried for all patients implanted from January 1, 2006, through December 31, 2007 who had a history of a myocardial infarction and an LVEF 30% or less. Those meeting the MADIT-II criteria (2,464 propensity score-matched patients) or the SCD-HeFT criteria (3,352 propensity score-matched patients) were included. In turn, all patients enrolled in MADIT-II (n=1,232) and patients randomized to receive placebo or ICD therapy in SCD-HeFT (n=1,676) were included.

Comparability of clinical characteristics between the registry and trial populations was assessed for the entire study population. Survival in the registry matched patients was compared to those who received primary prevention ICDs in MADIT-II (n = 742) and SCD-HeFT (n = 829).

As for baseline demographics, registry patients were substantially older and had a higher burden of comorbidities compared to clinical trial patients. Both registry and clinical trial patients were predominantly male and white (and these characteristics were incorporated into the match criteria). The median follow-up in MADIT-II and the matched registry patients was 19.5 months and 35.8 months, respectively. The median follow-up in SCD-HeFT and the matched registry patients was 46.1 months and 35.0 months, respectively.

Study results demonstrated no statistically significant difference in survival between MADIT-II-like registry patients and MADIT-II ICD patients (2-year mortality rates: 13.9% and 15.6%, respectively). The survival curves for the SCD-HeFT-like registry patients was coincidental with the SCD-HeFT ICD arm (3-year mortality rates: 17.3% and 17.4%, respectively). No survival differences were identified after restricting results to patients ≥ 65 years of age and upon covariate adjustment.

The authors concluded that "there was no significant difference in survival between clinical trial patients randomized to receive an ICD and a similar group of clinical registry patients who received a primary prevention ICD. Our findings support the continued use of primary prevention ICDs in similar patients seen in clinical practice."

The aim of this study was to assess the extent to which the clinical characteristics and long-term (~3 years) outcomes of registry patients with left ventricular systolic dysfunction undergoing primary prevention ICD implantation differ from those enrolled in the randomized controlled trials.

This was a retrospective analysis of patients, January 2006 to December 2009, with a new ICD (no prior ICD) for primary prevention and LVEF <50% implanted at one of 7 health plans of the Health Maintenance Organization Research Network (Henry Ford Health System, Kaiser Permanente [Colorado, Northern California, Northwest Southern California], Marshfield Clinic, and Meyers/Fallon Community Health Plan/U. Mass). The study also includes data on adjudicated arrhythmia episodes resulting in device therapies including shocks and ATP through manual medical record abstraction at the study sites with central clinical review and adjudication of source documentation. During the enrollment period, 3254 patients underwent primary prevention ICD placement with a final cohort comprised 2621 patients with LVSD after exclusions. The characteristics of the study cohort was compared to those of the MADIT-II (n=742) and SCD-HeFT (n=849) trials.

The purpose of this study is to assess the extent to which the clinical characteristics and long-term (~3 years) outcomes of registry patients with left ventricular systolic dysfunction undergoing primary prevention ICD implantation differ from those enrolled in the randomized, controlled trials.

Study results demonstrated that the mean age of the registry cohort was higher than that of the populations from the RCTs, ranging from almost 7 years higher than the mean age of the SCD-HeFT population to nearly 3 years higher than the average age of the population of MADIT-II. LVEF in the cohort was 25% (± 6.9%) compared to the RCTs (MADIT-II 23 ± 5%; SCD-HeFT 23.5 ± 7%). One quarter of the cohort population was female, significantly higher than that of MADIT-II, but similar to that of SCD-HeFT. Almost one third of the population was non-White, compared with 23% in SCD-HeFT; information on race was not reported in MADIT-II. The registry cohort population had a substantially higher comorbidity burden, including higher diabetes mellitus, hypertension, and atrial fibrillation/flutter. Registry patients were more likely to receive β blockers and statins and less likely to receive digoxin. Data on ICD therapy and outcomes (mortality and hospitalizations) was still being collected at the time that this manuscript was published.

The authors concluded that "this community-based cohort of patients receiving ICDs differs substantially from the populations that were enrolled in the RCTs that form the basis of clinical practice guidelines. Specifically, these [cohort] patients are older, included more women, have a substantial burden of important co-existing illnesses, and are more likely to receive evidence-based medications for heart failure."


The aim of this study was to examine the relationship between physician annual ICD implantation volume and in-hospital complications.
This was a retrospective cohort study of patients who had an ICD implanted between April 2006 and March 2010. Between April 2006 and March 2010, 4011 physicians performed 356,515 initial ICD implantations at 1 of 1463 hospitals which qualified for this study. The primary outcome was any adverse event that occurred during the implantation or preceding hospital discharge. Major adverse events were cardiac arrest, cardiac perforation, valve injury, coronary venous dissection, hemothorax, pneumothorax, deep vein thrombosis, transient ischemic attack, stroke, myocardial infarction, pericardial tamponade, and arteriovenous fistula.

Physicians were categorized as electrophysiologists, nonelectrophysiologist cardiologists (“cardiologists”), thoracic surgeons (“surgeons”), physicians who met the training standards for ICD implantation promulgated by the Heart Rhythm Society or none of the above. Physician procedure volume was annualized by dividing the total number of ICD implantations a physician performed by the number of years the physician contributed data to the NCDR ICD Registry.

Physicians were ranked by their annualized ICD implantation volume and divided them into quartiles of increasing procedure volume for analysis characteristics (number of beds, teaching status, urban location). The authors examined the relationship between physician annual ICD implantation volume and in-hospital complications, using hierarchical logistic regression to adjust for patient characteristics, implanting physician certification, hospital characteristics, hospital annual procedure volume, and the clustering of patients within hospitals and by physician. Patient characteristics included demographics (overall for the study: mean age 67 years, 27% female, 19% nonwhite race, and 5% Hispanic ethnicity). The authors also repeated the analysis for ICD subtypes: single chamber, dual chamber, and biventricular.

Study results demonstrated that ICD implant related complications occurred in 3.1% (10,994 patients) with 0.39% (1,375 patients) dying from these complications. The rate of adverse events was lower among patients who received a single-chamber ICD (1.9%) than those implanted with a dual-chamber ICD (2.9%). The complication rate decreased with increasing physician procedure volume from 4.6% in the lowest quartile to 2.9% in the highest quartile, and the mortality rate decreased from 0.72% to 0.36%. The relationship between physician procedure volume and decreased complications remained significant after adjusting for patient, physician, and hospital characteristics (OR 1.55 for complications in lowest-volume quartile compared with highest; 95% confidence interval, 1.34-1.79).

The authors concluded that “physicians who implant more ICDs have lower rates of procedural complications and in-hospital mortality, independent of hospital procedure volume, physician specialty, and ICD type.” Thus, “concentrating ICD implantation in the hands of fewer physicians may improve the clinical outcomes of this increasingly common procedure.”

The aim of this study was to estimate the association of implanting physician certification with in-hospital procedural complication rates following ICD implantation.

This was a retrospective cohort study of ICD Registry patients implanted, between January 2006 and June 2007 with an ICD without an epicardial lead or prior ICD implant. The investigators grouped implant procedures by the certification status of the implanting physician into mutually exclusive categories: electrophysiologists, nonelectrophysiologist cardiologists, thoracic surgeons, and other specialists. A total of 111,293 ICD implant procedures met the study criteria with 70.9% (78,857) of the procedures being performed by electrophysiologists, 21.9% (24,399) by nonelectrophysiologist cardiologists, 1.7% (1,862) by thoracic surgeons, and 5.5% (6,175) by other specialists. Overall study patient demographics included mean age 68 years, 27% female, 82% white race, 5% Hispanic ethnicity. The association of physician certification and risk of in-hospital complications was assessed using hierarchical generalized logistic regression to account for clustering of patients within hospitals.

Study results demonstrated that the characteristics of patients undergoing ICD implantation differed by physician specialty in several respects, notably by age, race, and payer status. Thoracic surgeons had greater mix of Medicare beneficiaries (75%) compared to the other specialties which ranged from 67-70%. The average patient age for thoracic surgeon patients (70.1 years) was approximately two years older than the other specialties. Thoracic surgeons treated a smaller mix of African Americans but higher mix of Hispanic ethnicity when compared to the other specialties. The category of “other specialists” were more likely than the named physician specialist categories to treat minority patients.

There were only clinically modest differences in measures of cardiac status such as history of congestive heart failure, NYHA class, prior cardiac arrest, history of atrial fibrillation, history of ventricular fibrillation, diabetes, hypertension, chronic lung disease, cerebrovascular disease, use of revascularization procedures, LVEF, QRS duration, and blood urea nitrogen measurements across physician certification categories.

Compared with patients whose ICD was implanted by electrophysiologists, patients whose ICD was implanted by either nonelectrophysiologist cardiologists or thoracic surgeons were at increased risk of complications in both unadjusted (electrophysiologists, 3.5% [2743/78,857]; nonelectrophysiologist cardiologists, 4.0% [970/24,399]; thoracic surgeons, 5.8% [108/1862]); with adjusted analyses (relative risk [RR] for nonelectrophysiologist cardiologists, 1.11 [95% confidence interval {CI}, 1.01-1.21]; RR for thoracic surgeons, 1.44 [95% CI, 1.15-1.79]).

The authors concluded that “in this registry, nonelectrophysiologists implanted 29% of ICDs. Overall, implantations by a nonelectrophysiologist were associated with a higher risk of procedural complications and lower likelihood of receiving a CRT-D device when indicated compared with patients whose ICD was implanted by an electrophysiologist.”
The aim of this study was to assess patient characteristics associated with not meeting the inclusion criteria of the clinical trials that demonstrated the efficacy of primary prevention ICDs.

This was a retrospective cohort study. The authors used data from the ICD Registry to identify patients who received ICDs for primary prevention from January 2006 to December 2008. They determined if patients met the inclusion criteria to at least one of four ICD primary prevention trials: Multicenter Automatic Defibrillator Implantation Trial (MADIT), MADIT-II, Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT), and the Multicenter Unsustained Tachycardia Trial (MUSTT). Of the 333,993 patients identified from the ICD Registry, a total of 150,264 remained after exclusions for sites with incomplete data reporting, secondary prevention, prior syncope, prior tachycardia arrest, sustained VT, and previous ICD placement.

Study demographics included, for the cohort meeting trial-based criteria (n=128,821), a mean age of 67±13 years, 80% white race and 73% male. The cohort not meeting trial-based criteria (n=21,443) had a mean age of 64±14 years, was 82% white and 74% male.

Study results demonstrated that 150,264 qualifying registry patients, 85.7% (128,821) met inclusion criteria for at least one of the primary prevention trials, 77.9% met SCD-HeFT criteria, 39.0% met MADIT II criteria, 1.6% met MUSTT criteria, and 0.9% met MADIT I criteria. On average, patients with older age (> 65 years), prior percutaneous coronary intervention, and prior coronary artery bypass grafting were more likely to meet trial criteria. For non-CRT ICDs, the proportion of implantations that did not meet trial inclusion criteria increased as patient age decreased, from 18% in those > 65 years old to 36% in the age group < 35 years old. In multivariate analysis, the significant predictors for not meeting trial criteria included prior cardiac transplantation (odds ratio [OR] 2.1), pediatric electrophysiology operator (OR 2.0), and high-grade atrioventricular conduction disease (OR 1.4).

The authors concluded that among patients in the ICD Registry receiving first-time ICDs for primary prevention, the majority met trial-based criteria.
The aim of this study was to assess temporal trends in patient characteristics and outcomes among older patients undergoing primary prevention ICD therapy in US hospitals between 2006 and 2010.

The authors performed multivariable hierarchical logistic regression to evaluate the temporal trends. The cohort included 117,100 patients from the NCDR ICD Registry: Medicare fee-for-service beneficiaries aged ≥65 years and older with LVEF ≤35% who underwent primary prevention ICD implantation, including those receiving concomitant CRT between 2006 and 2010, and could be matched to Medicare claims. Outcomes were all-cause mortality, heart failure hospitalization at 180 days, and device-related complications.

Study results demonstrated that between 2006 and 2010 there were only modest changes in baseline patient clinical characteristics. In the same timeframe, fewer single lead devices and more cardiac resynchronization therapy devices were used over time. There were statistically significant improvements in all outcomes, including 6-month all-cause mortality (7.1% in 2006, 6.5% 2010; adjusted odds ratio 0.88; 95% CI 0.82-0.95), 6-month rehospitalization (36.3% in 2006, 33.7% in 2010; adjusted odds ratio 0.87; 95% CI 0.83-0.91), and device-related complications (5.8% in 2006, 4.8% in 2010; adjusted odds ratio 0.80; 95% CI 0.74-0.88).

As for temporal trends in patient demographics, "the mean age for the entire group was 75.1±6.2 years. Between 2006 and 2010, the proportion of patients aged ≤75 years increased from 47.4% to 48.5% and those aged ≥85 years increased from 6.5% to 7.6% (P<0.001). The proportion of women undergoing ICD implantation increased modestly (27.3% to 28.4%, P=0.001). The proportion of white non-Hispanic patients decreased from 86.0% to 84.5%, whereas the proportion of blacks (7.7% to 8.8%), and Hispanics (4.1% to 4.7%), increased (P<0.001)."

The authors concluded that, given relatively stable clinical characteristics of patients selected for ICD implantation over this time period, the simultaneous improvements in outcomes "suggest meaningful advances in the care for this patient population."


The aim of this study was to assess the relationship between hospital ICD implantation volume and procedural complications.

The authors performed multivariable hierarchical logistic regression to evaluate the relationship, with the hypothesis that hospitals that perform higher volumes of procedures will generally have better clinical outcomes.

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The study included 224,233 patients from 1,201 US hospitals participating in the NCDR who had an initial ICD implantation between January 2006 and December 2008 and met study exclusion criteria. The primary outcome was any adverse event that occurred in hospital (to include during the implantation procedure). The study accounted for a wide range of patient demographics, clinical history and risk factors, ICD types (single chamber, dual chamber, biventricular) and data, implanting physician certification, and hospital characteristics such as size (patient beds), geographic location (rural, suburban, urban), and type (private/community, government, university).

Study demographics for all patients included a mean age of 66.8 years, 81% white, and 73% male, with slight variations when hospitals were categorized in quartiles based on annual ICD implantation volumes.

Study results demonstrated that “the rate of adverse events declined progressively with increasing procedure volume (p trend < 0.0001). This relationship remained significant (p trend < 0.0001) after adjustment for patient clinical characteristics, operator characteristics, and hospital characteristics. The volume-outcome relationship was evident for all ICD subtypes, including single-chamber (p trend = 0.004), dual-chamber (p trend < 0.0001), and biventricular ICDs (p trend = 0.02).”

The authors concluded that “patients who have an ICD implanted at a high-volume hospital are less likely to have an adverse event associated with the procedure than patients who have an ICD implanted at a low-volume hospital.”


The aim of this study was to determine the number, characteristics, and in-hospital outcomes of patients who receive a non–evidence-based ICD and examine the distribution of these implants by site, physician specialty, and year of procedure.

The authors performed a retrospective cohort analysis of 111,707 patients from the NCDR who received an ICD implantation between January 2006 and June 2009. The primary outcome was any in-hospital outcome, including death, any post-procedure complication, cardiac tamponade, pneumothorax, infection, hematoma, and length of hospital stay. Patients were classified as receiving a non–evidence-based ICD implant if they met at least one of following criteria: (1) had an MI within 40 days before ICD implantation; (2) had CABG surgery within 3 months before ICD implantation; (3) had NYHA class IV symptoms; or (4) had newly diagnosed heart failure at the time of ICD implantation. Patients who did not receive evidence-based ICD implantation were compared to those patients who did.
Study results demonstrated that 22.5% or the 111,707 study patients received non–evidence-based ICD implantation. “Patients who received a non–evidence-based ICD compared with those who received an evidence-based ICD had asignificantly higher risk of in-hospital death (0.57% [95% CI 0.48%-0.66%] vs 0.18% [95% CI 0.15%-0.20%]; P < .001) and any post-procedure complication (3.23% [95% CI 3.01%-3.45%] vs 2.41% [95% CI 2.31%-2.51%]; P < .001).” There was substantial variation in non–evidence-based ICDs by site. The rate of non–evidence-based ICD implants was significantly lower for electrophysiologists than non-electrophysiologists. There was no clear decrease in the rate of non–evidence-based ICDs over time.

Study demographics of patients receiving any non–evidence-based ICD implant included median age of 67 years (IQR 57 – 75 years). The majority were white (78%) and male (75%). Most patients had heart failure (91.8%) and ischemic heart disease (77.2%).

The authors concluded that “a substantial number of ICDs were implanted in patients who were similar to those who either were excluded from major clinical trials of primary prevention ICDs or shown not to benefit from ICD therapy in other trials. Such patients not only have more comorbidities than patients receiving an evidence-based device, but they are at a higher risk of in-hospital death and any post-procedure complication.” They also found no clear decrease in the overall number of non–evidence-based ICD implants over time. Thus the authors conclude that more effort is needed to improve physician adherence to evidence-based practice.

4. Medicare Evidence Development & Coverage Advisory Committee (MEDCAC)

A MEDCAC meeting was not convened on this issue.

5. Evidence-Based Guidelines

American College of Cardiology, American Heart Association, and Heart Rhythm Society Guidelines

The Class of Recommendation (COR) indicates the strength of the recommendation and estimates the magnitude of benefit versus risk.

- **Class 1 (Strong):** Is recommended. Should be performed/administered.
- **Class IIa (Moderate):** Is reasonable. Can be useful/effective/beneficial.
- **Class IIb (Weak):** May/might be reasonable. Usefulness/effectiveness is unknown/unclear/uncertain or not well established.
- **Class III: No Benefit (Moderate):** Is not recommended. Is not indicated/useful/effective/beneficial.
- **Class III: Harm (Strong):** Potentially harmful/Causes harm. Should not be performed/administered/other.

The Level of Evidence (LOE) rates the quality of the evidence based on the type, quantity, and consistency of the data from clinical trials and other sources.

- **Level A**
  - High-quality evidence from more than 1 RCT
  - Meta-analyses of high quality RCTs
  - One or more RCTs corroborated by high-quality registry studies
- **Level B-R**
  - Moderate-quality evidence from 1 or more RCTs
  - Meta-analyses of moderate-quality RCTs
- **Level B-NR**
  - Moderate-quality evidence from 1 or more well-designed, well-executed nonrandomized studies, observational studies, or registry studies
  - Meta-analysis of such studies
- **Level C-LD**
  - Randomized or nonrandomized observational or registry studies with limitations of design or execution
  - Meta-analyses of such studies
  - Physiological or mechanistic studies in human subjects
- **Level C-EO**
  - Consensus of expert opinion based on clinical experience

The following recommendations were put forward:
In patients with ischemic heart disease, who either survive SCA due to VT/VF or experience hemodynamically unstable VT (LOE: B-R) (1-4) or stable VT (LOE: B-NR) (5) not due to reversible causes, an ICD is recommended if meaningful survival greater than 1 year is expected.

In patients with ischemic heart disease and unexplained syncope who have inducible sustained monomorphic VT on electrophysiological study, an ICD is recommended if meaningful survival of greater than 1 year is expected (7) (LOE: B-NR).

In patients with ischemic heart disease and with NYHA class II or III HF despite GDMT, an ICD is recommended if meaningful survival of greater than 1 year is expected. (1, 2) (LOE: A).

In patients with NICM, HF with NYHA class II–III symptoms and an LVEF of 35% or less, despite GDMT, an ICD is recommended, if meaningful survival of greater than 1 year is expected (1-5) (LOE: B-NR).

In patients with HCM who have survived an SCA due to VT or VF, or have spontaneous sustained VT causing syncope or hemodynamic compromise, an ICD is recommended if meaningful survival greater than 1 year is expected (1, 6, 9, 10) (LOE: B-NR).

In patients with cardiac sarcoidosis who have sustained VT or are survivors of SCA or have an LVEF of 35% or less, an ICD is recommended, if meaningful survival of greater than 1 year is expected (1-5) (LOE: B-NR).

In patients with neuromuscular disorders, primary and secondary prevention ICDs are recommended for the same indications as for patients with NICM if meaningful survival of greater than 1 year is expected (1, 2) (LOE: B-NR).

In patients with a cardiac channelopathy and SCA, an ICD is recommended if meaningful survival of greater than 1 year is expected (7-13) (LOE: B-NR).

In patients with catecholaminergic polymorphic ventricular tachycardia and recurrent sustained VT or syncope, while receiving adequate or maximally tolerated beta blocker, treatment intensification with either combination medication therapy (e.g., beta blocker, flecainide), left cardiac sympathetic denervation, and/or an ICD is recommended (2-6) (LOE: B-NR).

In patients with Brugada syndrome with spontaneous type 1 Brugada electrocardiographic pattern and cardiac arrest, sustained VA or a recent history of syncope presumed due to VA, an ICD is recommended if a meaningful survival of greater than 1 year is expected (4, 6) (LOE: B-NR).

In patients with early repolarization pattern on ECG and cardiac arrest or sustained VA, an ICD is recommended (3, 4) (LOE: B-NR).

In patients with short QT syndrome who have a cardiac arrest or sustained VA, an ICD is recommended if meaningful survival greater than 1 year is expected (3-5) (LOE: B-NR).

In patients resuscitated from SCA due to idiopathic polymorphic VT or VF, an ICD is recommended if meaningful survival greater than 1 year is expected (9-13) (LOE: B-NR).

In patients with adult congenital heart disease and hemodynamically unstable VT, an ICD is recommended after evaluation and appropriate treatment for residual lesions/ventricular dysfunction if meaningful survival of greater than 1 year is expected (13-17) (LOE: B-NR).

In cases of myocarditis, an ICD is recommended if meaningful survival of greater than 1 year is expected (13-17) (LOE: B-NR).

In patients who meet criteria for an ICD who have inadequate vascular access or are at high risk for infection, and in whom pacing for bradycardia or VT termination or as part of CRT is neither needed nor anticipated, a subcutaneous implantable cardioverter-defibrillator is recommended (1-5) (LOE: B-NR).

Patients considering implantation of a new ICD or replacement of an existing ICD for a low battery should be informed of their individual risk of SCD and nonsudden death from HF or noncardiac conditions and the effectiveness, safety, and potential complications of the ICD in light of their health goals, preferences and values (1-5) (LOE: B-NR).
In ESC guidelines, “Class I” is defined as “evidence and/or general agreement that a given treatment or procedure is beneficial, useful, effective” and means that the intervention “is recommended” or “is indicated.” In turn, “Level of evidence A” is “data derived from multiple randomized clinical trials or meta-analyses” while level B is “data derived from a single randomized clinical trial or large non-randomized studies.” For definitions of all classes and levels of evidence, see the 2015 ESC guideline.

Secondary prevention of sudden cardiac death
For “secondary prevention of sudden cardiac death and ventricular tachycardia,” the ESC states that ICD implantation “is recommended” (Class I, Level A) in patients “with documented VF or hemodynamically not tolerated VT in the absence of reversible causes or within 48 h after myocardial infarction who are receiving chronic optimal medical therapy and have a reasonable expectation of survival with a good functional status >1 year.”
ESC states that ICD implantation “should be considered” (Class IIa, Level C) in patients “with recurrent sustained VT (not within 48 h after myocardial infarction) who are receiving chronic optimal medical therapy, have a normal LVEF and have a reasonable expectation of survival with good functional status for >1 year.” [ESC 1618]

On “sustained VT,” ESC states: “implantation of an ICD in patients with sustained VT increases survival compared with anti-arrhythmic drug therapy. To date, no trial has been conducted comparing catheter ablation for sustained VT without ICD implantation and ICD placement only. In view of the scarcity of data and the rather high rate of recurrence following catheter ablation for sustained VT, ICD implantation should be considered in all patients with LV dysfunction (ejection fraction <45%) and sustained VT.”

Primary prevention of sudden cardiac death
For primary prevention of sudden cardiac death, ESC states that “ICD therapy is recommended to reduce SCD in patients with symptomatic HF (NYHA class II–III) and LVEF ≤35% after ≥3 months of optimal medical therapy who are expected to survive for at least 1 year with good functional status:
• Ischemic etiology (at least 6 weeks after myocardial infarction) – Class I, Level A
• Nonischemic etiology – Class I, Level B”

Note that these ESC Guidelines were updated in 2015 and thus do not consider the DANISH trial, reported in 2016.
For patients who are asymptomatic or have preserved left ventricular function, ESC states: “currently there are no RCTs demonstrating the value of an ICD in asymptomatic patients (NYHA class I) with systolic dysfunction
(LVEF ≤35–40%) or in patients with HF and preserved LVEF >40–45%, so ICDs are not recommended for primary prevention in these patients.”

Special cases
For patients with New York Heart Association Class IV listed for heart transplantation, ESC states that ICD implantation “should be considered” (Class IIa, Level C) “for primary and secondary prevention of SCD in patients who are listed for heart transplant.”
ESC also supports consideration of ICD implantation for a number of familial or genetic diseases.

ESC states that subcutaneous defibrillators “should be considered” (Class IIa, Level C) “as an alternative to transvenous defibrillators in patients with an indication for an ICD when pacing therapy for bradycardia support, cardiac resynchronization or antitachycardia pacing is not needed.” ESC states that subcutaneous defibrillators “may be considered” (Class IIb, Level C) “as a useful alternative to the transvenous ICD system when venous access is difficult, after the removal of a transvenous ICD for infections or in young patients with a long-term need for ICD therapy.”

ESC states that wearable cardioverter defibrillators “may be considered” (Class IIb, Level C) for adult patients with poor LV systolic function who are at risk of sudden arrhythmic death for a limited period, but are not candidates for an implantable defibrillator (e.g. bridge to transplant, bridge to transvenous implant, peripartum cardiomyopathy, active myocarditis and arrhythmias in the early post-myocardial infarction phase). In discussing “gaps in evidence,” ESC further states that “wearable defibrillators may be an interesting therapeutic option in selected patients but require larger randomized trials before clear indications can be fully defined.”

Psychosocial management after ICD implantation
ESC states that: “assessment of psychological status and treatment of distress are recommended in patients with recurrent inappropriate shocks;” and “discussion of quality-of-life issues is recommended before ICD implantation and during disease progression in all patients.” (Both Class I, Level C.)

6. Professional Society Recommendations / Consensus Statements / Other Expert Opinion

Expert Consensus Statement

The writing group included members who are considered leaders in their field and were selected from the Heart Rhythm Society (HRS), American College of Cardiology (ACC), American Heart Association (AHA), Heart Failure Society of America (HFSA), and the Society of Thoracic Surgeons (STS). The group evaluated data to provide clinicians with guidance on four situations where ICD therapy might be beneficial in selected populations of patients who are not represented in clinical trials. The four situations evaluated were: 1) use of an ICD in patients with an abnormal troponin that is not due to a MI, 2) use of an ICD within 40 days after a MI, 3) use of an ICD within the first 90 days after revascularization, and 4) use of an ICD in the first 9 months after initial diagnosis of nonischemic cardiomyopathy.

The writing group recommends implantation of an ICD in the following patient populations:

- Patients with abnormal cardiac biomarkers that are not thought to be due to an MI and who otherwise would be candidates for implantation on the basis of primary prevention or secondary prevention criteria.
- Patients who, within 40 days of an MI, require non-elective permanent pacing, who also would meet primary prevention criteria for implantation of an ICD, and recovery of left ventricular function is uncertain or not expected.
- Patients who, within 40 days of an MI, develop sustained (or hemodynamically significant) ventricular tachyarrhythmias > 48 hours after an MI and in the absence of ongoing ischemia.
- Patients within 40 days of an MI and who have an ICD that requires elective replacement due to battery depletion, after careful assessment of comorbidities and the current clinical situation.
- Patients within 90 days of revascularization who have previously qualified for the implantation of an ICD for secondary prevention of SCD (resuscitated from cardiac arrest due to ventricular tachyarrhythmia) and have abnormal left ventricular function.
- Patients within 90 days of revascularization who have previously qualified for the implantation of an ICD for secondary prevention of SCD (resuscitated from cardiac arrest due to ventricular tachyarrhythmia) that is unlikely related to myocardial ischemia/injury and have normal left ventricular function.
- Patients within 90 days of revascularization who require nonelective permanent pacing, who would also meet primary prevention criteria for implantation of an ICD, and in whom recovery of LV function is uncertain or not expected.
- Patients within 90 days of revascularization with structural heart disease and sustained (or hemodynamically significant) VT that was not clearly related to acute MI or ischemia.
- Patients within 90 days of revascularization with an ICD that requires replacement due to battery depletion, after careful assessment of comorbidities and the current clinical situation.
- Patients < 9 months from the initial diagnosis of nonischemic cardiomyopathy who require nonelective permanent pacing, who would meet primary prevention criteria for implantation of an ICD, and recovery of LV function is uncertain or not expected.
- Patients < 9 months from the initial diagnosis of nonischemic cardiomyopathy with sustained (or hemodynamically significant) ventricular tachyarrhythmia.

**Appropriate Use Criteria**

The American College of Cardiology Foundation, the Heart Rhythm Society, and other specialty/subspecialty societies conducted a review of common clinical scenarios where ICDs and CRT are considered. As stated by the writing committee, “The clinical scenarios covered in this document address secondary prevention, primary prevention, comorbidities, generator replacement at elective replacement indicator, dual-chamber ICD, and CRT.”

7. Public Comment

Public comments sometimes cite the published clinical evidence and give CMS useful information. Public comments that give information on unpublished evidence such as the results of individual practitioners or patients are less rigorous and therefore less useful for making a coverage determination.

CMS uses the initial public comments to inform its proposed decision. CMS responds in detail to the public comments on a proposed decision when issuing the final decision memorandum. All comments that were submitted without personal health information may be viewed in their entirety by using the following link https://www.cms.gov/medicare-coverage-database/details/nca-view-public-comments.aspx?NCAId=288.

Initial Comment Period: 5/30/2017 – 6/29/2017

During the initial 30-day public comment period, CMS received 36 comments. Of these 36 comments, one was omitted from publication on the CMS website due to excessive personal health information content, and one commenter posted twice. Most of the comments suggested changes to the language in the current NCD from 2005 for implantable defibrillators. Several of these comments asked for CMS to update the covered indications based on current professional guidelines, and numerous commenters supported ending the registry requirement for ICD implantation. A number of commenters also made reference to the DANISH study and asked that CMS not limit coverage for patients with non-ischemic cardiomyopathy based on the findings from one trial using subgroup analyses that were not sufficiently powered and including patients with substantially elevated NT-proBNP who may have been more likely to die of non-SCD causes.

The majority of comments were provided by physicians/cardiologists, electrophysiologists, and other healthcare professionals. There were three comments that represented five professional societies, including the American Heart Association (AHA), American Stroke Association (ASA), Heart Failure Society of America (HFSA), Heart Rhythm Society (HRS), and American College of Cardiology (ACC). Additional groups who offered comments were AdvaMed, Emory Healthcare, Path to Improved Risk Stratification, and Mercy Health. We also received three comments from manufacturers of ICDs, including Medtronic, Boston Scientific, and Abbott.
During the 30-day comment period following the release of the proposed decision memorandum, CMS received 24 comments, with two comments coming from the same individual. In general, the public comments were overwhelmingly supportive of our proposed decision and were particularly supportive of our proposal to add cardiac MRI to the list of diagnostic imaging studies that can evaluate LVEF, as well as ending the data collection requirement. Several commenters also supported our proposal to have exceptions to the waiting periods for patients meeting CMS coverage criteria for cardiac pacemakers and who meet the criteria for an ICD, as well as for patients with an existing ICD and qualifying replacement.

While most of the comments agreed with our proposal to require an SDM interaction prior to ICD implantation for certain patients, many did not agree with the proposal that the SDM interaction come from an independent physician or qualified non-physician practitioner. Some comments also questioned the proposal to require the use of an evidence-based decision tool on ICDs as part of the SDM interaction. A few comments also questioned the proposed requirement that patients who have severe ischemic and/or non-ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to VF to have been on optimal medical therapy for at least three months.

The majority of comments were provided by professional societies and medical groups. Two comments represented four professional societies, including the AHA, ASA, HRS, and ACC. Additional groups who offered comments were AdvaMed, the Coalition to Transform Advanced Care (C-TAC), Cardiology Advocacy Alliance (CAA), Medical Imaging & Technology Alliance (MITA), Center to Advance Palliative Care (CAPC), Society for Cardiovascular Magnetic Resonance (SCMR), HCA Healthcare, Intermountain Healthcare, Michigan Medicine, Mercy Health, and Allina Health. We also received three comments from manufacturers of ICDs, including Medtronic, Boston Scientific, and Abbott. The remaining comments were from physicians/cardiologists, other healthcare professionals, and a university professor.

Prior “Personal History”

Comment: Several commenters asked what the term “prior personal history” was specifically referring to. Another commenter asked that we remove the term “personal” stating that this is not commonly used with the phrase “prior history.”

Response: The term “prior personal history” should have been specified throughout the proposed decision. We are changing the phrase “prior personal history” to “personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation” throughout the document. We have removed the term “prior” to eliminate redundancy in the phrase, recognizing that “history” can only be prior. We are keeping the term “personal” to specify that this refers to the health history of the individual, and not other types of history (e.g., family).
Comment: Several commenters disagreed with the proposed requirement that the SDM encounter must occur with an "independent" physician or qualified non-physician practitioner. Many believed this would delay the procedure and increase burden for patients and providers by creating an unnecessary step, and that the implanting physician is capable of providing the SDM interaction. Some commenters disagreed with a requirement for an SDM encounter prior to initial ICD implantation, stating informed consent would be sufficient for the patient.

Response: We believe that a SDM encounter prior to initial ICD implantation is a critical step in empowering patient choice in their treatment plan. While ICDs have remained a common treatment option for many years, the strength of evidence for an ICD benefit is different for different patient populations. As mentioned in the Analysis section of this decision memo, the joint 2017 guidelines by AHA/ACC/HRS state that "In patients with VA [Ventricular Arrhythmia] or at increased risk for SCD, clinicians should adopt a shared decision-making approach in which treatment decisions are based not only on the best available evidence but also on the patients’ health goals, preferences, and values,” topics that are not typically covered when obtaining informed consent. The SDM interaction requires the use of an evidence-based tool to ensure topics like the patients’ health goals and preferences are covered before ICD implantation. We want to ensure that the patient receives more information than the risks and benefits of the procedure. We also recognize that requiring a SDM encounter with an "independent" physician or non-physician professional could create unnecessary burden, so we have decided to remove the word "independent" from the SDM requirement. In order to provide flexibility for this requirement, we are indicating that the SDM interaction may occur at a separate visit.

Comment: There were a number of comments regarding the SDM tool that is part of the SDM encounter requirement. Many wanted clarity around what is considered an "evidence-based decision tool," while some believed that a decision tool was not necessary for the SDM encounter. Another commenter stated that existing tools have not yet been "validated." There were a couple of comments requesting that the decision tool on ICDs used for SDM include information about when and how the ICD might be deactivated in the future.

Response: CMS believes in the importance of using an evidence-based decision tool as part of the SDM encounter. While we are not specifying the type of SDM tool that is required for the SDM encounter, the Analysis section of this decision memo gives an example of an existing SDM tool for ICDs, which is an evidence-based decision aid for patients with heart failure who are at risk for sudden cardiac death and are considering an ICD. This decision tool was funded by the National Institutes on Aging and the Patient-Centered Outcomes Research Institute, and can be found at https://patientdecisionaid.org/wp-content/uploads/2017/01/ICD-Infographic-5.23.16.pdf. This tool is based on published clinical research and interviews with patients; it also includes discussion of the option for future ICD deactivation.

Comment: One commenter suggested that an SDM encounter should be required for all ICD indications.
Response: While we encourage SDM for all covered indications, we will only require a SDM interaction for certain patient populations to limit the added burden of this interaction. We also recognize that certain patient populations, such as patients with a personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, lack alternate treatment options and SDM would have less of an impact compared to patient populations with more treatment options available.

**Inappropriate Shocks**

Comment: One commenter questioned a statement in the Background section of this decision memo that ICDs never actually “fire” in the majority of patients who receive them for primary prevention, and suggested that the references cited did not provide primary data to support the statement.

Response: We believe this statement is supported by the evidence. Merchant (2016) cites relevant primary data in the section of his article with the subheading “Presence of appropriate ICD therapies during first battery life.”

**ICD Battery Life**

Comment: One comment referred to our reference of ICD battery life being 5-7 years, stating that ICDs are now lasting 7-10 years.

Response: We acknowledge that like other types of batteries, newer types of ICD batteries have longer lives than older types. We are changing our statement to reflect that ICD batteries may last 5-10 years depending on the type of battery and the extent of its use for the patient while implanted.

**Removal of NYHA Class IV Indication for CRT**

Comment: One comment requested clarification on what is meant when we said we are removing the Class IV heart failure requirement for CRT, and if this means that we are proposing to create a separate coverage decision for CRT.
Response: As we discuss in the Analysis section, the requirement of patients having NYHA Class IV heart failure for CRT was not included in the proposed decision. We believe CRT is a separate therapy and should be evaluated in a distinct determination. Because there is no national coverage criteria for CRT, we have removed this section from the NCD on ICDs. At this time, CMS has not proposed to open a separate NCD on CRT.

**MRI for Evaluating LVEF**

Comment: While we received a number of comments supporting the addition of cardiac MRI to the list of diagnostic imaging studies that can evaluate LVEF, one commenter suggested including cardiac MRI for the evaluation of ventricular scarring or infiltration as a finding supporting the indication for ICD implantation.

Response: There is strong evidence supporting the addition of cardiac MRI to our list of modalities that may be used to evaluate LVEF. Our focus here is on depressed LVEF as it is an important, trial-based risk factor for SCD in patients who have no history of documented cardiac arrest or life-threatening ventricular tachyarrhythmia. We recognize that MRI as well as other diagnostic tests may be useful in the clinical work up of such patients, but we are not mandating which tests should be used for which patients, or when in the course of their clinical evaluation.

Comment: One commenter requested that CMS consider excluding cardiac MRI for measuring LVEF from the requirements of the forthcoming Appropriate Use Criteria (AUC) for advanced imaging services since the NCD would include cardiac MRI as an approved method of measuring LVEF.

Response: CMS appreciates this comment; however, AUC requirements are outside the scope of this NCD.

**Optimal Medical Therapy (OMT)**

Comment: One commenter recommended the indications for severe ischemic and/or non-ischemic dilated cardiomyopathy, NYHA Class II or III heart failure patients with LVEF less than or equal to 35% in covered indication B3, be separated by removing the OMT requirement for ischemic dilated cardiomyopathy.
Based on our review of the clinical trial evidence (to include trials evaluated in the 2005 NCD but not re-evaluated in the current NCD), we believe that the evidence is equivocal on whether patients with ischemic cardiomyopathy and systolic heart failure would benefit from a new requirement for at least 3 months of OMT alone prior to ICD placement, if they otherwise met criteria for an ICD. We thus accept this recommendation to split the ischemic and nonischemic categories, and to apply the 3 month OMT requirement to the nonischemic heart failure population, but not to the ischemic population.

Comment: A couple commenters expressed concern about the requirement of being on OMT at least three months before implantation, with some commenters suggesting that it may not be possible for some patients to meet the requirement of having been on OMT for three months prior to implantation due to being unable to tolerate certain medications. One commenter suggested the wording “clinicians must have tried for at least three months to optimize medical therapy to the extent tolerated by the patient.” Another commenter suggested wording that would allow ICD implantation prior to completing three months of OMT if a patient’s condition worsens during the OMT period.

Response: We understand this concern and appreciate these comments. OMT must be tailored to the patient. If a patient cannot tolerate a given medication, then that medication is not optimal medical therapy for that patient. We also note that the evidence for ICDs is based on chronic stable patients and based on the best available evidence do not believe ICDs should be implanted prior to the completion of three months of OMT for patients with non-ischemic dilated cardiomyopathy whose condition continues to worsen during that time. Therefore, we are not making any changes to the wording of the OMT requirement.

Subcutaneous ICD

Comment: One commenter suggested that the NCD should specifically address the indications for the subcutaneous ICD.

Response: The coverage indications in this NCD apply to subcutaneous ICDs.

Covered Indications

Comment: There were a couple comments that suggested re-wording the phrase “uncontrolled supraventricular tachycardia such as from atrial fibrillation” that is listed as part of the exclusion criteria for this NCD, with one commenter suggesting the phrase “persistent supraventricular tachycardia such as atrial fibrillation with a poorly controlled ventricular rate,” and another suggesting that patients with atrial fibrillation could receive AV nodal ablation and ICD placement in one procedure.
Response: We are re-wording the phrase “uncontrolled supraventricular tachycardia such as from atrial fibrillation” to “supraventricular tachycardia such as atrial fibrillation with a poorly controlled ventricular rate,” as this clarifies that a patient would not be automatically excluded if they have atrial fibrillation. We will not be including the term “persistent” because this may be misinterpreted as referring to persistent atrial fibrillation. We did not review the merits of AV nodal ablation with or without simultaneous ICD placement as part of this NCD, and the commenter did not present supporting evidence or citations. However, we note that AV nodal ablation is a therapy aimed at preventing atrial fibrillation and controlling the heart rate. A problem with patients with atrial fibrillation and poorly controlled ventricular rates, as noted in trial publications and by commenters alike, is that this could lead to inappropriate defibrillator shocks.

Comment: One comment suggested removing “symptomatic” from the second bullet point under coverage indication B1, and defining “ongoing ischemia” and the timeframe for “acute myocardial infarction.” Another comment also suggested to change “MI” to “Type 1 MI” to provide clarity regarding whether demand ischemia, troponin leaks, etc. are subject to the myocardial infarction waiting period.

Response: We have re-worded coverage indication B1 to more closely resemble the wording from the 2005 NCD. This includes removing the terms “symptomatic” and “ongoing ischemia” from the decision language in order to provide clarity on covered indications for patients with a personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation. We leave definitions of common clinical terms such as acute myocardial infarction, myocardial infarction, and ischemia, to professional medical societies and clinical experts who define such terms in published guidelines and clinical trials. We note that universal definitions for some of these terms exist, and such terms may be updated faster than CMS national coverage determinations.

Comment: One commenter suggested that we include specific language for CRT indications in our exceptions to waiting periods for patients who meet CMS coverage requirements for cardiacpacemakers, and who meet criteria for coverage for an ICD.

Response: As previously mentioned in this document, CRT devices are outside the scope of this decision, and CRT devices are currently covered at local contractor discretion and not currently subject to an NCD.

Comment: There was a comment that suggested adding patients who previously met criteria in sections B2 or B3 before an MI or revascularization event occurred, and patients who are status post Left Ventricular Assist Device (LVAD) placement to our list of exceptions to the waiting periods required in sections B2 and B3.

Response: We appreciate the comment and recommendations. The waiting periods are based on evidence from well-known clinical trials as cited in the Evidence section of this decision. In the absence of new evidence, we are...
unable to add these patient populations to our list of exceptions to waiting periods for patients that have had a CABG, or PCI with angioplasty and/or stenting, within the past 3 months, or had a myocardial infarction within the past 40 days.

Comment: One commenter suggested that discontinuing the data collection requirement would eliminate coverage for ICDs for the primary prevention of sudden cardiac death.

Response: ICDs for the primary prevention of sudden cardiac death will be covered under the indication for “patients who have severe ischemic and/or non-ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation.”

Comment: One comment suggested that the term “significant, irreversible brain damage” could be too subjective and lead to appropriate candidates being denied an ICD, and asked us to instead consider the wording “significant, irreversible brain damage that renders the patient unable to provide informed consent or participate in shared decision making.”

Response: The phrase “significant, irreversible brain damage” listed as part of the exclusion criteria is consistent with wording in the 2005 NCD, and with trial exclusion criteria and medical society clinical guidelines. We believe clinicians fully understand that this exclusion, for example, applies to comatose patients in a vegetative state, but may not apply to all patients with persistent speech or motor dysfunction as a result of a prior stroke, but who otherwise meet criteria and are good candidates for ICD placement. We do not believe patients or clinicians would benefit by more granular definition.

Comment: One commenter stated that for patients who present with syncope in the setting of an ischemic cardiomyopathy, non-ischemic cardiomyopathy or congenital heart disease, VT or VF can be diagnosed using any one of numerous sources including history and physical exam and any number of diverse diagnostic tests.

Response: Patients who experience syncope must meet one of the criteria categories in this decision in order to receive an ICD. There is no evidence to support that syncopal patients who do not meet any of these criteria would nonetheless have improved survival by ICD placement. Please note that patients with unexplained syncope and inducible tachyarrhythmias are already covered in paragraph B(1) of our decision. We recognize the role of the physician in patient care and the roles of professional society guidelines in clinical practice and do not further specify valid tests for diagnosing spontaneous ventricular tachyarrhythmias.

Comment: A couple commenters suggested adding ICD coverage for New York Heart Association class IV heart failure patients awaiting heart transplantation.
Response: We are aware of the use of ICDs and cardiac resynchronization therapy (CRT) for patients who are candidates for heart transplantation on the United Network for Organ Sharing (UNOS) transplant list awaiting a donor heart, as bridge to transplant to prolong survival until a donor becomes available. Patients with end-stage heart failure awaiting heart transplantation have been treated with specific medications, ICDs and CRT devices including ones with defibrillator functions. While CRT is outside the scope of this NCD, patients who are candidates for heart transplantation awaiting a donor heart have specific individual considerations in optimizing treatment pending transplantation. There may be interactions with other NCDs as well (e.g., ventricular assist devices (NCD 20.9) and heart transplant (NCD 260.9)). For these reasons, coverage determinations under section 1862(a)(1)(A) for this specific limited group of patients will remain with the local Medicare Administrative Contractors (MACs).

Comment: One comment suggested language to clarify the criteria for cardiac pacemaker implantation during the primary prevention waiting period. The commenter suggested stating the specific time exceptions of “even if this occurs within 40 days post MI or if revascularization occurred within in the past 3 months.”

Response: CMS agrees that including the specific time exceptions in this section would provide clarity. Since this applies to both cardiac pacemakers and ICD replacement, the wording has been placed at the beginning of section C.

Future Research

Comment: Some commenters emphasized the importance of further research, and at least one encouraged “ongoing collaboration and coordination” among federal agencies and medical societies “to answer outstanding questions.”

Response: As stated in the Analysis section of this document, we agree that further research could be done on risk stratification and specific subpopulations. We encourage studies such as, but not limited to, definitive evaluation of an ICD benefit in Medicare patients who are elderly, or who meet criteria for CRT, or who have substantially improved LVEF when they have reached the point of generator exchange.

VIII. CMS Analysis

Introduction: National coverage determinations are determinations by the Secretary with respect to whether or not a particular item or service is covered nationally by Medicare (§1869(f)(1)(B) of the Act). In order to be covered by Medicare, an item or service must fall within one or more benefit categories contained within Part A or Part B, and must not be otherwise excluded from coverage. Moreover, with limited exceptions, the expenses
incurred for items or services must be reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of a malformed body member (§1862(a)(1)(A) of the Act).

When making national coverage determinations, we evaluate the evidence related to our analytic questions based on the quality, strength and totality of evidence presented in the reviewed literature. As part of this evaluation, it is important to consider whether the evidence is relevant to the Medicare beneficiary population. In determining the generalizability of the results of the body of evidence to the Medicare population, we consider, at minimum, the age, race and gender of the study participants.

Evidence Review Summary: Existing Medicare coverage policies and professional medical society guidelines alike are based on evidence from randomized controlled trials that support the use of ICDs in patients with heart failure and either ischemic or non-ischemic dilated cardiomyopathy. However, we note that many of these trials were conducted 15-20 years ago and since that time medical therapy has evolved substantially. Since our last decision, several meta-analyses, systematic evidence reviews and technology assessments have been published and provide evidence that ICDs are reasonable and necessary for certain patients aligning with our prior criteria. In general, meta-analysis when designed appropriately and performed rigorously can provide high strength evidence.

Questions: For this reconsideration, CMS focused on the following questions:

- Is there evidence to conclude that ICDs decrease mortality for patients with ischemic dilated cardiomyopathy (IDCM) and reduced LVEF?
- Is there evidence to conclude that ICDs decrease mortality for patients with non-ischemic cardiomyopathy (NIDCM) and reduced LVEF?

Is there evidence to conclude that ICDs decrease mortality for patients with ischemic dilated cardiomyopathy (IDCM) and reduced LVEF?

Yes. Uhlig and colleagues (2013) analyzed 14 trials including IRIS and concluded that “there is a high strength of evidence that ICD therapy for primary prevention of SCD, versus no ICD therapy, shows benefit with regard to all-cause mortality and SCD in patients with reduced left ventricular ejection fraction and ischemic or non-ischemic cardiomyopathy beyond the immediate post-MI or coronary revascularization periods.” Colquitt and colleagues (2014) analyzed 13 trials including IRIS and found that “ICDs reduced all-cause mortality in people at increased risk of SCD, defined in trials as those with previous ventricular arrhythmias/cardiac arrest, myocardial infarction (MI) > 3 weeks previously, non-ischemic cardiomyopathy (depending on data included) or ischemic/non-ischemic heart failure (HF) and left ventricular ejection fraction \( \leq 35\% \).”

Is there evidence to conclude that ICDs decrease mortality for patients with non-ischemic cardiomyopathy (NIDCM) and reduced LVEF?

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Yes. Overall, based on the preponderance of evidence including past randomized controlled trials, new trials, meta-analyses, systematic evidence reviews and technology assessments, we conclude that the evidence is sufficient to conclude that ICDs decrease mortality for symptomatic patients with severe ischemic and severe non-ischemic dilated cardiomyopathy and reduced LVEF.

Specifically for patients with heart failure and non-ischemic cardiomyopathy, Al-Khatib and colleagues (2017) analyzed four trials including DANISH and showed that “use of an ICD significantly reduced all-cause mortality (HR 0.75; CI 0.61-0.93; P=.008, P=0.87 for heterogeneity).” Golwala and colleagues (2017) analyzed six trials including DANISH and showed that “use of an ICD significantly reduced all-cause mortality by 23% (HR 0.77; CI 0.64–0.91).”

We believe this decision reflects the currently available evidence, including professional society guidelines. Further, by adding patient shared decision making we are empowering patients to be part of their health treatment decisions. In addition, by removing the data collection we are reducing the burden on providers.

The general criteria from the 2005 NCD which applies to all categories of patients who are candidates for an ICD are mostly unchanged or represent minor modifications based on recent published consensus guidelines by professional medical societies, evidence reviewed and/or public comments. We go into detail below regarding covered indications.

**Patients with a Personal History of Sustained Ventricular Tachyarrhythmia or Cardiac Arrest Due to Ventricular Fibrillation:** A literature search did not find any new trials since the 2005 NCD relevant to the clinical utility of ICDs for patients with a prior personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation. Therefore, no changes were made to this section of the decision.

**Patients with a Prior Myocardial Infarction and a measured LVEF ≤ 0.30:** There were several systematic reviews and meta-analyses assessing these patients with prior MI and a measured LVEF ≤ 0.30. These analyses confirmed the benefit of ICDs in these patients. Therefore, no changes in coverage were made for these patients.

**Patients Who Have Severe Ischemic Dilated Cardiomyopathy but No Personal History of Sustained Ventricular Tachyarrhythmia or Cardiac Arrest Due to Ventricular Fibrillation:** For patients who have ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, ICD coverage is minimally changed from the 2005 NCD.
For Medicare patients with ischemic cardiomyopathy, the baseline standard of care of MADIT II (2002) and SCD-HeFT (2005) may no longer reflect the contemporary standard of care (Bardy et al, 2005; Moss et al., 2002). However, other than IRIS (2009), which demonstrated that ICDs do not improve survivability if implanted within 31 days after myocardial infarction, confirming the results of DINAMIT (2004) and supporting our prior coverage criterion, no other major trials have been completed assessing the clinical utility of ICDs versus optimal medical therapy in patients with ischemic cardiomyopathy, since the 2005 reconsideration (Hohnloser et al, 2004; Steinbeck et al., 2009).

Patients Who Have Severe Non-Ischemic Dilated Cardiomyopathy but No Personal History of Sustained Ventricular Tachyarrhythmia or Cardiac Arrest Due to Ventricular Fibrillation: For patients who have non-ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation is minimally changed from our existing coverage policy.

Results from the DANISH study called into question the utility of ICDs for certain patients with heart failure and non-ischemic cardiomyopathy. While we agree that DANISH contributed to the evidence base for the treatment of heart failure patients at risk for SCD, there are several factors to consider: 1) the study population may be different with the inclusion of NT-proBNP and 58% CRT composition of DANISH in the ICD and control groups; 2) DANISH was not powered for subgroup analyses on age and CRT status; and 3) the absence of a parallel registry made it difficult to determine whether eligible patients who enrolled were similar to those who were not.

Two aspects of the baseline optimal medical therapy in the DANISH trial stand out. First, it is the only trial that includes CRT as a component of baseline optimal medical therapy. While an important consideration, cardiac resynchronization therapy is distinct and generally beyond the scope of this decision. Second, it reflects the marked advancement of pharmacological therapy that has occurred since the earlier ICD trials. As with other medical conditions, better medical treatments and care develops over time with evidence and experience. The determination of effect of these changes on trial results compared to prior trials is a topic of discussion which will not be answerable in this decision. Importantly, with additional treatment options, a detailed discussion of the potential benefits and harms with each patient would be a key step to choosing the best treatment for any particular individual.

The requirement of being on optimal medical therapy for at least three months is in the inclusion criteria of multiple trials, including the 2016 DANISH trial, and we believe it is reasonable for non-ischemic heart failure patients. We found no evidence to support making any other changes to the 2005 NCD on the use of ICDs for patients with non-ischemic dilated cardiomyopathy, or to make changes to the LVEF requirements from the 2005 reconsideration.

Other Clinical Circumstances: This decision provides the same coverage of ICDs for patients with no history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation and familial or genetic disorders. Patients with endstage heart failure awaiting heart transplantation have been treated with CRT devices including ones with defibrillator functions. Since CRT was not part of this analysis, determinations for this specific limited group of patients will remain with the local Medicare Administrative Contractors. The rationale, evidence, benefits and harms, professional society recommendations and consensus guidelines were all considered for these clinical circumstances.
The requirement of patients having NYHA Class IV heart failure for CRT was removed in this decision since we believe CRT is a separate therapy and should be evaluated in a distinct determination. Because there is no national coverage criteria for CRT, we have removed this section of the NCD.

**General Patient Criteria:** Based on the evidence reviewed and the 2017 society guidelines, we made no changes for eligible clinically stable patients requirement. For eligible clinically stable patients, we made no changes from the 2005 ICD reconsideration were made. We continue to require that these patients must not have significant irreversible brain damage or any disease other than cardiac disease associated with a likelihood of survival of less than one year. We added cardiac MRI to the list of diagnostic imaging studies that can evaluate left ventricular ejection fraction, based on new evidence in part cited in recent professional medical society guidelines and public comments (Al-Khatib et al., 2017; Russo et al., 2013).

**Exceptions to Waiting Periods for Patients that have had a CABG, or PCI with Angioplasty and/or Stenting, within the Past 3 Months, or Had a Myocardial Infarction within the Past 40 Days:** We made certain exceptions to the waiting period requirements for symptomatic patients who have ischemic and non-ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, acknowledging that the wording in the 2005 decision could inadvertently require a second procedure when only one procedure is needed. In patients with an existing ICD who then suffer an MI or undergo a coronary revascularization procedure, we would now allow replacement of the device, if clinically indicated, forgoing the previously mentioned waiting periods. This exception also applies to patients without an existing ICD who meet requirements for both a cardiac pacemaker and a defibrillator; thus, if the pacemaker is needed for pacing needs, the defibrillator may be implanted in the same procedure, and a patient would no longer be required to undergo a second procedure for the defibrillator after the expiration of the waiting periods. This is standard medical practice and recommended in the HRS/ACC/AHA expert consensus statement (Kusumoto et al. 2014). We note that specific coverage criteria for pacemakers goes beyond the scope of this NCD, which focuses on the clinical utility of ICDs.

We also made an exception to the required waiting periods for patients with an existing ICD that requires replacement due to the end of battery life, elective replacement indicator (ERI), or device malfunction, with documentation that the device is at ERI level, or that there is a device/lead malfunction. This is practical medical practice and recommended in the HRS/ACC/AHA expert consensus statement (Kusumoto et al. 2014).

**Patient Shared Decision Making:** Since there are some outstanding questions regarding the appropriate populations benefitting from ICDs, we are including a requirement for a patient shared decision making (SDM) interaction in our decision for certain patient populations. We do not believe an SDM interaction would be beneficial for patients with a personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation given the lack of alternative treatment options, nor do we believe SDM is necessary for patients with an existing ICD who qualify for a replacement ICD since a previous implantation had already occurred. SDM is especially important in treatments where there are complex considerations on benefits, harms, indications and existing effective treatments. Barry and Edgman-Levitan (2012) noted: “[t]he process by which the optimal decision may be reached for a patient at a fateful health crossroads is called shared decision making and involves, at minimum, a clinician and the patient, although other members of the health care team or friends and family members may be invited to participate. In shared decision making, both parties share information: the clinician offers options and describes their risks and benefits, and the patient expresses his or her preferences and values. Each participant is thus armed with a better understanding of the relevant factors and shares responsibility in the decision about how to proceed.” Ideally SDM integrates the use of evidence-based decision tools including treatment pictograms to characterize benefits and harms.
The importance of individual patient values and preferences in decision making applies to the use of ICDs. As endorsed in the joint 2017 guidelines by AHA/ACC/HRS, “In patients with VA [Ventricular Arrhythmia] or at increased risk for SCD, clinicians should adopt a shared decision-making approach in which treatment decisions are based not only on the best available evidence but also on the patients’ health goals, preferences, and values”.

An example of an existing SDM tool for ICDs is the joint effort between The Colorado Program for Patient Centered Decisions, with funding from the National Institutes on Aging (K23AG040696) and the Patient-Centered Outcomes Research Institute (P>1000116-01) to develop an evidence-based decision aid tool for patients with heart failure considering an ICD who are at risk for sudden cardiac death (primary prevention).

https://patientdecisionaid.org/wp-content/uploads/2017/01/ICD-Infographic-5.23.16.pdf. The decision aid tools and website were developed based on research study findings and interviews with patients.

In addition to the tool, they developed a website which leads patients step-by-step through some information on ICDs designed to increase patients’ knowledge of their medical condition, the risks and benefits of available treatments and to empower patients to become more involved in the decision-making process.

https://patientdecisionaid.org/icd/.

Registry Requirement: In 2005, CMS had some questions about the evidence (see Appendix C for § 20.4 of the NCD). Based on our concerns at the time, we required additional data to be collected via a registry (see https://www.cms.gov/medicare-coverage-database/details/ncd-details.aspx?NCDId=110&ncdver=3&NCAId=288&bc). We assessed the extent to which the published literature has addressed the following ten “initial hypotheses” for the registry data collection requirement in the 2005 NCD. Based on our analysis, the eight peer-reviewed publications were directly related to the 10 initial hypotheses. CMS believes that additional data collection is no longer needed. We have listed the 10 hypotheses below for the ease of the reader. In addition, we have summarized the publications and have explained why they answered the hypotheses.

1. The clinical characteristics of the patients receiving ICDs are similar to those of patients involved in the primary prevention randomized clinical trials.
2. The indications for ICD implantation in patients are similar to those in the primary prevention randomized clinical trials.
3. The in-hospital procedure related complications for patients are similar to those in the primary prevention randomized clinical trials.
4. Certified providers competent in ICD implantation are implanting ICD devices in patients.
5. Patients who receive an ICD represent patients for which current clinical guidelines and the evidence base recommend implantation.
6. The clinical characteristics and indications for ICD implantation do not differ significantly among facilities.
7. The clinical characteristics and indications for ICD implantation do not differ significantly among providers.
8. The in-hospital procedure related complications for ICD implantation do not differ significantly among facilities.
9. The in-hospital procedure related complications for ICD implantation do not differ significantly among providers.
10. The in-hospital procedure related complications for ICD implantation do not differ significantly among device manufacturer, types, and/or programming.
In sum, we believe the research questions for each of the studies specifically listed in the Evidence Section addresses one or more of the ten CMS “initial hypotheses;” and collectively, these eight studies are related to all ten hypotheses. Again, we are aware that there are numerous other published studies that may be related to one or more of these hypotheses, or aspects of them. These real world studies have provided support for implementation of this technology outside the controlled trial environment. The initial data collection requirement through the NCDR has served to generate and improve the evidence base for the use of ICDs in certain Medicare beneficiaries. We acknowledge the substantial contribution of the NCDR, the input of a number of professional societies and contribution of manufacturers in this positive collaborative effort over the past decade. We believe it has served its purpose and are ending the registry data collection requirement. However, we encourage the continuation and improvement of a voluntary registry for purposes of quality improvement, safety, and appropriate use verification.

Considerations for Further Research: Based on our analysis, we recognize that further research could be done on risk stratification and specific subpopulations. We also recognize the separate but related CRT technology. In discussion of research gaps in a 2015 AHRQ-funded Technology Assessment, Rickard et al conclude: “The effectiveness of CRT-D versus CRT-P in patients with an LVEF ≤35% has not been adequately addressed.” In the 2014 NICE Health Technology Assessment, Colquitt et al conclude: “A RCT comparing CRT-D [CRT with a defibrillator] and CRT-P [a CRT pacemaker alone] in people with heart failure as a result of left ventricular systolic dysfunction and cardiac dyssynchrony is required, for both those with and those without an ICD indication.

While the 2016 DANISH trial has made a new contribution to this evidence base, Kober et al., concluded it is unclear whether patients who are eligible for CRT should routinely receive an ICD, and a randomized study (CRT-D to CRT-P) would be clinically important. Accordingly, we believe randomized controlled trials comparing CRT-D to CRT-P on the backdrop of contemporary, optimal pharmacological therapy in specific patient populations with dilated cardiomyopathy would be informative. Such trials would provide evidence and supporting documentation for a separate consideration of CRT.

We appreciate the importance of further research on a myriad of potential risk stratification models that combine clinical and demographic data with data from various studies (such as laboratory tests, interventional electrophysiological (EP) studies, cardiac magnetic resonance imaging (MRI), to name just a few), whether alone or in some combination or sequence, and for particular populations. As noted by Uhlig et al. in their 2013 technology assessment: “Prevention is the primary strategy to lower death from SCD. However, SCD is a particular management challenge because the majority of cases occur in individuals without a prior diagnosis of cardiac disease or other clear risk factors for SCD. The most common underlying cardiovascular diagnosis among people with SCD is coronary artery disease (CAD). Yet, in about half of the cases of SCD, SCD itself is the initial manifestation of CAD. The clinical strategy to prevent death from SCD involves identification of risk factors for ventricular tachyarrhythmias and SCD, to target individuals for medical and interventional treatments.”

While we encourage such research on risk stratification to continue, we acknowledge that other agencies are better equipped and have clearer authority to take the lead in vetting and supporting such a large and varied research portfolio, some of which is in earlier discovery and testing phases. However, it is important to note that all other indications for ICDs not currently covered in accordance with this decision may continue to be covered under Category B IDE trials (42 CFR 405.201). The end result of such research could improve health outcomes of Medicare beneficiaries.
Health Disparities

Current research has not addressed the disparities of ICD implantation in African Americans when compared to Caucasians. Despite being at higher risk for SCD, Hernandez et al. (2007) found that African Americans who were potentially eligible for an ICD were 30% less likely to receive an ICD compared to Caucasians, independent of other characteristics. Hernandez et al. (2007) also observed this disparity in women who were potentially eligible for an ICD, finding that they were 40% less likely than men to undergo ICD therapy. African Americans and women are both traditionally underrepresented in research and these findings illustrate the need for future research to address these disparities.

Summary

This NCA has focused on the use of ICDs in symptomatic patients who have severe ischemic and non-ischemic dilated cardiomyopathy with no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation. As we have discussed, we have not substantively changed the current coverage policy for ICDs for patients with a personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation. We have added exceptions to waiting periods for patients that have had a CABG, or PCI with angioplasty and/or stenting, within the past 3 months, or had a myocardial infarction within the past 40 days for patients who are pacemaker dependent and otherwise qualify for an ICD implant and for beneficiaries with an existing ICD which requires replacement, eliminating the possibility of unnecessary invasive procedures and allows for timely replacement of a potentially lifesaving system which requires replacement.

Additionally, we have required a shared decision making encounter to engage patients, to discuss their treatment options and the potential benefits and harms associated with ICDs, and to incorporate their views and beliefs in choosing the most appropriate treatment. Based on the evidence, we will reduce provider burden and documentation through discontinuation of the registry data collection requirement and the need to document ischemic cardiomyopathy etiology. While we agree that some research questions remain, the data and information gathered from the NCDR has substantially contributed to the evidence base and supports the ending of the registry data collection requirements from the 2005 NCD. We acknowledge that additional research questions remain that cannot be addressed by registries and we provide details about the study designs that could potentially address these questions. It is our understanding that such studies are currently being planned.

IX. Conclusion

A. The Centers for Medicare & Medicaid Services (CMS) has determined that the evidence is sufficient to conclude that the use of implantable cardioverter defibrillators (ICDs, also referred to as defibrillators) is reasonable and necessary for the treatment of illness or injury or to improve the functioning of a malformed body member under section 1862(a)(1)(A) of the Social Security Act.

CMS is finalizing relatively minimal changes to the ICD NCD from the 2005 reconsideration. We summarize the changes below and fully explain the changes in the Analysis section of the NCD decision memo.
Patient Criteria
- Add cardiac magnetic resonance imaging (MRI) to the list of diagnostic imaging studies that can evaluate left ventricular ejection fraction (LVEF);
- Require patients who have severe non-ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation to have been on optimal medical therapy (OMT) for at least 3 months;
- Require a patient shared decision making (SDM) interaction prior to ICD implantation for certain patients.

Additional Patient Criteria
- Remove the Class IV heart failure requirement for cardiac resynchronization therapy (CRT).

Exceptions to Waiting Periods
- Add an exception for patients meeting CMS coverage requirements for cardiac pacemakers, and who meet the criteria for an ICD;
- Add an exception for patients with an existing ICD and qualifying replacement.

Registry Requirement
- End the data collection requirement.

We are finalizing changes to the 20.4 NCD that reflect the 2005 reconsideration as described below:

B. Covered Indications

1. Patients with a personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation. Patients must have demonstrated:
   - An episode of sustained ventricular tachyarrhythmia, either spontaneous or induced by an electrophysiology (EP) study, not associated with an acute myocardial infarction and not due to a transient or reversible cause; or
   - An episode of cardiac arrest due to ventricular fibrillation, not due to a transient or reversible cause.

2. Patients with a prior myocardial infarction and a measured left ventricular ejection fraction (LVEF) ≤ 0.30. Patients must not have:
   - New York Heart Association (NYHA) classification IV heart failure;
   - Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
   - Had a myocardial infarction within the past 40 days; or
   - Clinical symptoms and findings that would make them a candidate for coronary revascularization.

   For these patients identified in B2, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

3. Patients who have severe ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, and have New York Heart Association (NYHA) Class II or III heart failure, left ventricular ejection fraction (LVEF) ≤ 35%. Additionally, patients must not have:
- Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
- Had a myocardial infarction within the past 40 days; or
- Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B3, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

4. Patients who have severe non-ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, and have New York Heart Association (NYHA) Class II or III heart failure, left ventricular ejection fraction (LVEF) \( \leq 35\% \), been on optimal medical therapy (OMT) for at least 3 months. Additionally, patients must not have:
   - Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
   - Had a myocardial infarction within the past 40 days; or
   - Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B4, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

5. Patients with documented familial, or genetic disorders with a high risk of life-threatening tachyarrhythmias (sustained ventricular tachycardia or ventricular fibrillation), to include, but not limited to, long QT syndrome or hypertrophic cardiomyopathy.

For these patients identified in B5, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

6. Patients with an existing ICD may receive an ICD replacement if it is required due to the end of battery life, elective replacement indicator (ERI) or device/lead malfunction.
1. Patients must be clinically stable (e.g., not in shock, from any etiology);
2. Left ventricular ejection fraction (LVEF) must be measured by echocardiography, radionuclide (nuclear medicine) imaging, cardiac magnetic resonance imaging (MRI), or catheter angiography;
3. Patients must not have:
   ▪ Significant, irreversible brain damage; or
   ▪ Any disease, other than cardiac disease (e.g., cancer, renal failure, liver failure) associated with a likelihood of survival less than 1 year; or
   ▪ Supraventricular tachycardia such as atrial fibrillation with a poorly controlled ventricular rate.

C. Exceptions to waiting periods for patients that have had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months, or had a myocardial infarction within the past 40 days:

   Cardiac Pacemakers: Patients who meet all CMS coverage requirements for cardiac pacemakers and who meet the criteria in this national coverage determination for an ICD may receive the combined device in one procedure at the time the pacemaker is clinically indicated;

   Replacement of ICDs: Patients with an existing ICD may receive a ICD replacement if it is required due to the end of battery life, elective replacement indicator (ERI) or device/lead malfunction.

D. Other Indications:

   For patients who are candidates for heart transplantation on the United Network for Organ Sharing (UNOS) transplant list awaiting a donor heart, coverage of ICDs, as with cardiac resynchronization therapy, as a bridge to transplant to prolong survival until a donor becomes available is determined by the local Medicare Administrative Contractors (MACs).

   All other indications for ICDs not currently covered in accordance with this decision may be covered under Category B IDE trials (42 CFR 405.201).

See Appendix B for the NCD manual language.

APPENDIX A
General Methodological Principles of Study Design
(Section VI of the Decision Memorandum)

When making national coverage determinations, CMS evaluates relevant clinical evidence to determine whether or not the evidence is of sufficient quality to support a finding that an item or service is reasonable and necessary. The overall objective for the critical appraisal of the evidence is to determine to what degree we are confident that: 1) the specific assessment questions can be answered conclusively; and 2) the intervention will
We divide the assessment of clinical evidence into three stages: 1) the quality of the individual studies; 2) the generalizability of findings from individual studies to the Medicare population; and 3) overarching conclusions that can be drawn from the body of the evidence on the direction and magnitude of the intervention’s potential risks and benefits.

The methodological principles described below represent a broad discussion of the issues we consider when reviewing clinical evidence. However, it should be noted that each coverage determination has its unique methodological aspects.

Assessing Individual Studies

Methodologists have developed criteria to determine weaknesses and strengths of clinical research. Strength of evidence generally refers to: 1) the scientific validity underlying study findings regarding causal relationships between health care interventions and health outcomes; and 2) the reduction of bias. In general, some of the methodological attributes associated with stronger evidence include those listed below:

- Use of randomization (allocation of patients to either intervention or control group) in order to minimize bias.
- Use of contemporaneous control groups (rather than historical controls) in order to ensure comparability between the intervention and control groups.
- Prospective (rather than retrospective) studies to ensure a more thorough and systematical assessment of factors related to outcomes.
- Larger sample sizes in studies to demonstrate both statistically significant as well as clinically significant outcomes that can be extrapolated to the Medicare population. Sample size should be large enough to make chance an unlikely explanation for what was found.
- Masking (blinding) to ensure patients and investigators do not know to that group patients were assigned (intervention or control). This is important especially in subjective outcomes, such as pain or quality of life, where enthusiasm and psychological factors may lead to an improved perceived outcome by either the patient or assessor.

Regardless of whether the design of a study is a randomized controlled trial, a non-randomized controlled trial, a cohort study or a case-control study, the primary criterion for methodological strength or quality is to the extent that differences between intervention and control groups can be attributed to the intervention studied. This is known as internal validity. Various types of bias can undermine internal validity. These include:

- Different characteristics between patients participating and those theoretically eligible for study but not participating (selection bias).
- Co-interventions or provision of care apart from the intervention under evaluation (performance bias).
- Differential assessment of outcome (detection bias).
Occurrence and reporting of patients who do not complete the study (attrition bias).

In principle, rankings of research design have been based on the ability of each study design category to minimize these biases. A randomized controlled trial minimizes systematic bias (in theory) by selecting a sample of participants from a particular population and allocating them randomly to the intervention and control groups. Thus, in general, randomized controlled studies have been typically assigned the greatest strength, followed by non-randomized clinical trials and controlled observational studies. The design, conduct and analysis of trials are important factors as well. For example, a well-designed and conducted observational study with a large sample size may provide stronger evidence than a poorly designed and conducted randomized controlled trial with a small sample size. The following is a representative list of study designs (some of that have alternative names) ranked from most to least methodologically rigorous in their potential ability to minimize systematic bias:

- Randomized controlled trials
- Non-randomized controlled trials
- Prospective cohort studies
- Retrospective case control studies
- Cross-sectional studies
- Surveillance studies (e.g., using registries or surveys)
- Consecutive case series
- Single case reports

When there are merely associations but not causal relationships between a study’s variables and outcomes, it is important not to draw causal inferences. Confounding refers to independent variables that systematically vary with the causal variable. This distorts measurement of the outcome of interest because its effect size is mixed with the effects of other extraneous factors. For observational, and in some cases randomized controlled trials, the method in that confounding factors are handled (either through stratification or appropriate statistical modeling) are of particular concern. For example, in order to interpret and generalize conclusions to our population of Medicare patients, it may be necessary for studies to match or stratify their intervention and control groups by patient age or co-morbidities.

Methodological strength is, therefore, a multidimensional concept that relates to the design, implementation and analysis of a clinical study. In addition, thorough documentation of the conduct of the research, particularly study selection criteria, rate of attrition and process for data collection, is essential for CMS to adequately assess and consider the evidence.

Generalizability of Clinical Evidence to the Medicare Population

The applicability of the results of a study to other populations, settings, treatment regimens and outcomes assessed is known as external validity. Even well-designed and well-conducted trials may not supply the evidence needed if the results of a study are not applicable to the Medicare population. Evidence that provides accurate information about a population or setting not well represented in the Medicare program would be considered but would suffer from limited generalizability.
The extent to which the results of a trial are applicable to other circumstances is often a matter of judgment that depends on specific study characteristics, primarily the patient population studied (age, sex, severity of disease and presence of co-morbidities) and the care setting (primary to tertiary level of care, as well as the experience and specialization of the care provider). Additional relevant variables are treatment regimens (dosage, timing and route of administration), co-interventions or concomitant therapies, and type of outcome and length of follow-up.

The level of care and the experience of the providers in the study are other crucial elements in assessing a study’s external validity. Trial participants in an academic medical center may receive more or different attention than is typically available in non-tertiary settings. For example, an investigator’s lengthy and detailed explanations of the potential benefits of the intervention and/or the use of new equipment provided to the academic center by the study sponsor may raise doubts about the applicability of study findings to community practice.

Given the evidence available in the research literature, some degree of generalization about an intervention’s potential benefits and harms is invariably required in making coverage determinations for the Medicare population. Conditions that assist us in making reasonable generalizations are biologic plausibility, similarities between the populations studied and Medicare patients (age, sex, ethnicity and clinical presentation) and similarities of the intervention studied to those that would be routinely available in community practice.

A study’s selected outcomes are an important consideration in generalizing available clinical evidence to Medicare coverage determinations. One of the goals of our determination process is to assess health outcomes. These outcomes include resultant risks and benefits such as increased or decreased morbidity and mortality. In order to make this determination, it is often necessary to evaluate whether the strength of the evidence is adequate to draw conclusions about the direction and magnitude of each individual outcome relevant to the intervention under study. In addition, it is important that an intervention’s benefits are clinically significant and durable, rather than marginal or short-lived. Generally, an intervention is not reasonable and necessary if its risks outweigh its benefits.

If key health outcomes have not been studied or the direction of clinical effect is inconclusive, we may also evaluate the strength and adequacy of indirect evidence linking intermediate or surrogate outcomes to our outcomes of interest.

**Assessing the Relative Magnitude of Risks and Benefits**

Generally, an intervention is not reasonable and necessary if its risks outweigh its benefits. Health outcomes are one of several considerations in determining whether an item or service is reasonable and necessary. CMS places greater emphasis on health outcomes actually experienced by patients, such as quality of life, functional status,
duration of disability, morbidity and mortality, and less emphasis on outcomes that patients do not directly experience, such as intermediate outcomes, surrogate outcomes, and laboratory or radiographic responses. The direction, magnitude, and consistency of the risks and benefits across studies are also important considerations. Based on the analysis of the strength of the evidence, CMS assesses the relative magnitude of an intervention or technology’s benefits and risk of harm to Medicare beneficiaries.

APPENDIX B
Medicare National Coverage Determinations Manual

This draft NCD is subject to formal revisions and formatting changes prior to the release of the final NCD contractor instructions and publication in the Medicare National Coverage Determinations Manual.

Table of Contents
(Rev.)

A. General

An implantable cardioverter defibrillator is an electronic device designed to diagnose and treat life-threatening ventricular tachyarrhythmias. The device consists of a pulse generator and electrodes for sensing and defibrillating. This therapy has been shown in trials to improve survival and reduce sudden cardiac death in patients with certain clinical characteristics.

B. Nationally Covered Indications

Effective for services performed on or after [Month/XX] [Day/XX], [20XX] CMS has determined that the evidence is sufficient to conclude that the use of implantable cardioverter defibrillators (ICDs, also referred to as defibrillators) is reasonable and necessary:

Covered Indications

1. Patients with a personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation. Patients must have demonstrated:
An episode of sustained ventricular tachyarrhythmia, either spontaneous or induced by an electrophysiology (EP) study, not associated with an acute myocardial infarction and not due to a transient or reversible cause; or
An episode of cardiac arrest due to ventricular fibrillation, not due to a transient or reversible cause.

2. Patients with a prior myocardial infarction and a measured left ventricular ejection fraction (LVEF) \( \leq 0.30 \). Patients must not have:
   - New York Heart Association (NYHA) classification IV heart failure;
   - Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
   - Had a myocardial infarction within the past 40 days; or
   - Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B2, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

3. Patients who have severe ischemic dilated cardiomyopathy but no personal history of sustained ventricular tachyarrhythmia or cardiac arrest due to ventricular fibrillation, and have New York Heart Association (NYHA) Class II or III heart failure, left ventricular ejection fraction (LVEF) \( \leq 35\% \). Additionally, patients must not have:
   - Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
   - Had a myocardial infarction within the past 40 days; or
   - Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B3, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

4. Patients who have severe non-ischemic dilated cardiomyopathy but no personal history of cardiac arrest or sustained ventricular tachyarrhythmia, New York Heart Association (NYHA) Class II or III heart failure, left ventricular ejection fraction (LVEF) \( \leq 35\% \), been on optimal medical therapy for at least 3 months. Additionally, patients must not have:
   - Had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months; or
   - Had a myocardial infarction within the past 40 days; or
   - Clinical symptoms and findings that would make them a candidate for coronary revascularization.

For these patients identified in B4, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.
5. Patients with documented familial, or genetic disorders with a high risk of life-threatening tachyarrhythmias (sustained ventricular tachycardia or ventricular fibrillation), to include, but not limited to, long QT syndrome or hypertrophic cardiomyopathy.

For these patients identified in B5, a formal shared decision making encounter must occur between the patient and a physician (as defined in Section 1861(r)(1)) or qualified non-physician practitioner (meaning a physician assistant, nurse practitioner, or clinical nurse specialist as defined in §1861(aa)(5)) using an evidence-based decision tool on ICDs prior to initial ICD implantation. The shared decision making encounter may occur at a separate visit.

6. Patients with an existing ICD may receive an ICD replacement if it is required due to the end of battery life, elective replacement indicator (ERI), or device/lead malfunction.

For each of these groups listed above, the following additional criteria must also be met:

1. Patients must be clinically stable (e.g., not in shock, from any etiology);
2. Left ventricular ejection fraction (LVEF) must be measured by echocardiography, radionuclide (nuclear medicine) imaging, cardiac magnetic resonance imaging (MRI), or catheter angiography;
3. Patients must not have:
   ◦ Significant, irreversible brain damage; or
   ◦ Any disease, other than cardiac disease (e.g., cancer, renal failure, liver failure) associated with a likelihood of survival less than 1 year; or
   ◦ Supraventricular tachycardia such as atrial fibrillation with a poorly controlled ventricular rate.

Exceptions to waiting periods for patients that have had a coronary artery bypass graft (CABG), or percutaneous coronary intervention (PCI) with angioplasty and/or stenting, within the past 3 months, or had a myocardial infarction within the past 40 days:

Cardiac Pacemakers: Patients who meet all CMS coverage requirements for cardiac pacemakers, and who meet the criteria in this national coverage determination for an ICD, may receive the combined devices in one procedure, at the time the pacemaker is clinically indicated;

Replacement of ICDs: Patients with an existing ICD may receive a ICD replacement if it is required due to the end of battery life, elective replacement indicator (ERI), or device/lead malfunction.

C. Other Indications

For patients that are candidates for heart transplantation on the United Network for Organ Sharing (UNOS)

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transplant list awaiting a donor heart, coverage of ICDs, as with cardiac resynchronization therapy, as a bridge to transplant to prolong survival until a donor becomes available is determined by the local Medicare Administrative Contractors (MACs).

All other indications for ICDs not currently covered in accordance with this decision may be covered under Category B IDE trials (42 CFR 405.201).


A. General

The implantable automatic defibrillator is an electronic device designed to detect and treat life-threatening tachyarrhythmias. The device consists of a pulse generator and electrodes for sensing and defibrillating.

B. Covered Indications

1. Documented episode of cardiac arrest due to ventricular fibrillation (VF), not due to a transient or reversible cause (effective July 1, 1991).

2. Documented sustained ventricular tachyarrhythmia (VT), either spontaneous or induced by an electrophysiology (EP) study, not associated with an acute myocardial infarction (MI) and not due to a transient or reversible cause (effective July 1, 1999).

3. Documented familial or inherited conditions with a high risk of life-threatening VT, such as long QT syndrome or hypertrophic cardiomyopathy (effective July 1, 1999).
Additional indications effective for services performed on or after October 1, 2003:

4. Coronary artery disease with a documented prior MI, a measured left ventricular ejection fraction (LVEF) ≤ 0.35, and inducible, sustained VT or VF at EP study. (The MI must have occurred more than 40 days prior to defibrillator insertion. The EP test must be performed more than 4 weeks after the qualifying MI.)

5. Documented prior MI and a measured LVEF ≤ 0.30 and a QRS duration of >120 milliseconds (the QRS restriction does not apply to services performed on or after January 27, 2005). Patients must not have:
   a. New York Heart Association (NYHC) classification IV;
   b. Cardiogenic shock or symptomatic hypotension while in a stable baseline rhythm;
   c. Had a coronary artery bypass graft (CABG) or percutaneous transluminal coronary angioplasty (PTCA) within past 3 months;
   d. Had an enzyme positive MI within past month (Effective for services on or after January 27, 2005, patients must not have an acute MI in the past 40 days);
   e. Clinical symptoms or findings that would make them a candidate for coronary revascularization; or
   f. Any disease, other than cardiac disease (e.g., cancer, uremia, liver failure), associated with a likelihood of survival less than 1 year.

Additional indications effective for services performed on or after January 27, 2005:

6. Patients with ischemic dilated cardiomyopathy (IDCM), documented prior MI, NYHA Class II and III heart failure, and measured LVEF ≤ 35%;

7. Patients with non-ischemic dilated cardiomyopathy (NIDCM) >9 months, NYHA Class II and III heart failure, and measured LVEF ≤ 35%;

8. Patients who meet all current Centers for Medicare & Medicaid Services (CMS) coverage requirements for a cardiac resynchronization therapy (CRT) device and have NYHA Class IV heart failure;
All indications must meet the following criteria:

a. Patients must not have irreversible brain damage from preexisting cerebral disease;

b. MIs must be documented and defined according to the consensus document of the Joint European Society of Cardiology/American College of Cardiology Committee for the Redefinition of Myocardial Infarction; ¹

Indications 3 - 8 (primary prevention of sudden cardiac death) must also meet the following criteria:

a. Patients must be able to give informed consent;

b. Patients must not have:

¹ Alpert and Thygesen et al., 2000.
   Criteria for acute, evolving or recent MI.

Either one of the following criteria satisfies the diagnosis for an acute, evolving or recent MI:

1) Typical rise and gradual fall (troponin) or more rapid rise and fall (CK-MB) of biochemical markers of myocardial necrosis with at least one of the following:

   a) ischemic symptoms;
   b) development of pathologic Q waves on the ECG;
   c) ECG changes indicative of ischemia (ST segment elevation or depression); or
   d) coronary artery intervention (e.g., coronary angioplasty).

2) Pathologic findings of an acute MI.

Criteria for established MI.

Any one of the following criteria satisfies the diagnosis for established MI:

1) Development of new pathologic Q waves on serial ECGs. The patient may or may not remember previous symptoms. Biochemical markers of myocardial necrosis may have normalized, depending on the length of time that has passed since the infarct developed.

2) Pathologic findings of a healed or healing MI.
• Cardiogenic shock or symptomatic hypotension while in a stable baseline rhythm;
• Had a CABG or PTCA within the past 3 months;
• Had an acute MI within the past 40 days;
• Clinical symptoms or findings that would make them a candidate for coronary revascularization;
• Any disease, other than cardiac disease (e.g., cancer, uremia, liver failure), associated with a likelihood of survival less than 1 year;

c. Ejection fractions must be measured by angiography, radionuclide scanning, or echocardiography;

d. The beneficiary receiving the defibrillator implantation for primary prevention is enrolled in either a Food and Drug Administration (FDA)-approved category B investigational device exemption (IDE) clinical trial (42 CFR §405.201), a trial under the CMS Clinical Trial Policy (National Coverage Determination (NCD) Manual §310.1) or a qualifying data collection system including approved clinical trials and registries. Initially, an implantable cardiac defibrillator (ICD) database will be maintained using a data submission mechanism that is already in use by Medicare participating hospitals to submit data to the Iowa Foundation for Medical Care (IFMC) a Quality Improvement Organization (QIO) contractor for determination of reasonable and necessary and quality improvement. Initial hypothesis and data elements are specified in this decision (Appendix VI) and are the minimum necessary to ensure that the device is reasonable and necessary. Data collection will be completed using the ICDA (ICD Abstraction Tool) and transmitted via QNet (Quality Network Exchange) to the IFMC who will collect and maintain the database. Additional stakeholder-developed data collection systems to augment or replace the initial QNet system, addressing at a minimum the hypotheses specified in this decision, must meet the following basic criteria:

- Written protocol on file;
- Institutional review board review and approval;
- Scientific review and approval by two or more qualified individuals who are not part of the research team;
- Certification that investigators have not been disqualified.

For purposes of this coverage decision, CMS will determine whether specific registries or clinical trials meet these criteria.

f. Providers must be able to justify the medical necessity of devices other than single lead devices. This justification should be available in the patient's medical record.

9. Patients with NIDCM >3 months, NYHA Class II or III heart failure, and measured LVEF ≤ 35%, only if the following additional criteria are also met:

a. Patients must be able to give informed consent;

b. Patients must not have:

- Cardiogenic shock or symptomatic hypotension while in a stable baseline rhythm;
- Had a CABG or PTCA within the past 3 months;
- Had an acute MI within the past 40 days;
- Clinical symptoms or findings that would make them a candidate for coronary revascularization;
- Irreversible brain damage from preexisting cerebral disease;
- Any disease, other than cardiac disease (e.g. cancer, uremia, liver failure), associated with a likelihood of survival less than 1 year;

c. Ejection fractions must be measured by angiography, radionuclide scanning, or echocardiography;
e. The beneficiary receiving the defibrillator implantation for this indication is enrolled in either an FDA-approved category B IDE clinical trial (42 CFR §405.201), a trial under the CMS Clinical Trial Policy (NCD Manual §310.1), or a prospective data collection system meeting the following basic criteria:

- Written protocol on file;
- Institutional Review Board review and approval;
- Scientific review and approval by two or more qualified individuals who are not part of the research team;
- Certification that investigators have not been disqualified.

For purposes of this coverage decision, CMS will determine whether specific registries or clinical trials meet these criteria.

d. Providers must be able to justify the medical necessity of devices other than single lead devices. This justification should be available in the patient's medical record.

C. Other Indications

All other indications for implantable automatic defibrillators not currently covered in accordance with this decision will continue to be covered under Category B IDE trials (42 CFR §405.201) and the CMS routine clinical trials policy (NCD §310.1).

2 Ibid

(This NCD last reviewed February 2005.)

APPENDIX D – NYHA Class Symptoms

<table>
<thead>
<tr>
<th>Class</th>
<th>Functional Capacity: How a patient with cardiac disease feels during physical activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>No limitation of physical activity. Ordinary physical activity does not cause undue fatigue, palpitation, dyspnea (shortness of breath).</td>
</tr>
</tbody>
</table>
### Class Objective Assessment

<table>
<thead>
<tr>
<th>Class</th>
<th>Objective Assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>No objective evidence of cardiovascular disease. No symptoms and no limitation in ordinary physical activity.</td>
</tr>
<tr>
<td>C</td>
<td>Objective evidence of moderately severe cardiovascular disease. Marked limitation in activity due to symptoms, even during less-than-ordinary activity. Comfortable only at rest.</td>
</tr>
<tr>
<td>D</td>
<td>Objective evidence of severe cardiovascular disease. Severe limitations. Experiences symptoms even while at rest.</td>
</tr>
</tbody>
</table>

*Retrieved from AHA website, [https://www.heart.org/HEARTORG/Conditions/HeartFailure/AboutHeartFailure/Classes-of-Heart-Failure_UCM_306328_Article.jsp?appName=MobileApp](https://www.heart.org/HEARTORG/Conditions/HeartFailure/AboutHeartFailure/Classes-of-Heart-Failure_UCM_306328_Article.jsp?appName=MobileApp). Current as of November 14, 2017.*

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