

Product Performance Report

2018 First Edition



Letter from Abbott

As a world leader in the development of state-of-the-art technology for cardiac rhythm management devices, Abbott continuously strives to partner with physicians in reducing risks and facilitating the best possible patient outcomes. We understand that our products are implanted in people whose health and well-being depend on their performance. From product design through patient follow-up, Abbott employees are dedicated to product quality and patient safety.

In keeping with this commitment, we publish a Product Performance Report (PPR) semi-annually to ensure that the healthcare community and the patients it serves are informed about the overall performance of our cardiac devices, which include implantable cardiac monitors (ICMs), implantable cardioverter defibrillators (ICDs), implantable pacemakers, and implantable pacing and defibrillation leads. Abbott recognizes that such performance data must be transparent and consistent. In order to meet these goals we continue our commitment to the reporting methods described in the 2009 AdvaMed document [“Industry Guidance for Uniform Reporting of Clinical Performance of Cardiac Rhythm Management Pulse Generators and Leads”](#), which set new standards for lead performance reporting and specifically addressed the reporting of active registry performance data. Determined to provide the highest level of transparency, Abbott goes beyond the AdvaMed recommendations by identifying the root cause of each ICM, ICD, and pacemaker laboratory-confirmed malfunction and providing subcategories of laboratory-confirmed lead abrasion and fracture malfunctions.

Continuing within this edition of the PPR and consistent with previously published editions, Abbott reports on expanded data from actively monitored studies. Since 2007, the PPR has featured pacemaker, ICD, and lead data from the St. Jude Medical Product Longevity and Performance Registry (SCORE). Post-Approval studies are now standard practice for Abbott, providing a rich source of actively collected and continuously monitored reliability and performance data for cardiac rhythm management products. This PPR also features a product performance data set which includes OPTIMUM, SCORE and three Post-Approval Studies. This combined dataset encompasses more than 62,000 implants from multiple product families, including leads, ICDs and pacemakers, making it the most comprehensive actively monitored product performance dataset in the industry. We are continuing to expand the scope of confirmed product malfunction summaries with worldwide confirmed malfunctions in Durata™ and Optisure™ defibrillation lead models, our more recent ICD and pacemaker models and various low voltage and CRT leads which will further expand to different devices and leads in future additions.

In addition to providing performance data, a summary of advisories on implantable devices since 1999 can be found beginning on page 325.

As we continually strive to provide unbiased and reliable information on the performance of our products, Abbott is pleased to release the first edition of the 2018 Product Performance Report containing the latest performance information on our ICMs, ICDs, pacemakers and lead systems.

Sincerely,



Susan Jezior Slane
Divisional Vice President, Quality Assurance and Compliance

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Serving Our Mission

Abbott is advancing the treatment of heart and vascular disease through breakthrough medical technologies, allowing people to restore their health and get on with their lives. We focus on improving treatment options for coronary artery disease, cardiac rhythm management, atrial fibrillation, heart failure, structural heart and peripheral artery disease. We are here for the people we serve in their pursuit of healthy lives. This has been the way of Abbott for more than a century—passionately and thoughtfully translating science into lasting contributions to health.

Toward this mission, we maintain a rigorous approach to ensuring the quality of our products. The key elements of this effort include:

- Compliance with U.S. and international quality system standards, such as the U.S. FDA Quality Systems Regulation (21 CFR Part 820) and ISO 13485 (an international standard for the Quality Management System for medical devices)
- Thorough evaluation of product design, including extensive design verification and validation, as well as product qualification testing
- Rigorous control of the design and manufacturing processes
- Inspection and qualification of externally supplied components and materials
- Timely analysis of returned products, including extensive malfunction investigation
- Extensive internal auditing
- Post market surveillance
- Continuous improvement programs
- Ensuring the highest ethical standards

We continue to be committed to answering your questions and keeping you informed. If you have any questions or concerns, please contact your Abbott Representative or Abbott Technical Services at 1-800-722-3774. Thank you for your input and continued support, allowing Abbott to positively impact the lives of thousands of patients every year.

What's New in This Report

UPDATE ON FORTIFY™, FORTIFY ASSURA™, QUADRA ASSURA™, QUADRA ASSURA MP™, UNIFY™, UNIFY ASSURA™ AND UNIFY QUADRA™

ICD PREMATURE BATTERY DEPLETION ADVISORY

In order to provide our physician customers and patients the most up-to-date information, Abbott has included an update on the Fortify™, Fortify Assura™, Quadra Assura™, Quadra Assura MP™, Unify™, Unify Assura™ and Unify Quadra™ ICD premature battery depletion advisory in the Focus on Clinical Performance section (see pages 304-307). This section includes an update on the analysis of products returned to Abbott. Additionally, for advisory models with at least 500 active devices in service, Abbott provides a separate product performance data page.

UPDATE ON RIATA™ LEAD PERFORMANCE

Abbott continues to include an update on Riata lead performance in the Focus on Clinical Performance section (see pages 308-316). This section provides the latest Riata lead externalized conductor rates from the Abbott Riata Lead Evaluation Study, passive complaint and returns handling, and describes in considerable detail the rates of other types of Riata insulation abrasion failure mechanisms that Abbott has identified from returns analysis.

UPDATE ON DURATA™ LEAD PERFORMANCE

Durata lead performance continues to meet expectations by all measures. Our confidence in the Durata lead performance is based on combined data from three prospective, actively monitored registries that include approximately 11,000 Optim™ insulated defibrillation leads. Additionally, this section provides details on the very low rate of abrasion failures that have been identified on Optim insulated defibrillation leads. A statistical analysis of this registry data performed by PHRI, an independent, third-party, is presented in this special Focus on Clinical Performance section (see pages 317-322).

UPDATE ON OPTIM™ LEAD INSULATION

The Abbott Optim lead insulation combines the best characteristics of two established lead insulation materials, polyurethane and silicone. This novel insulation technology imparts lubricity, strength, and abrasion resistance while still maintaining flexibility and biostability. This Product Performance Report provides an up-to-date statistical assessment of the benefits of Optim lead insulation on Abbott defibrillation leads (see pages 323-324).

Customer Reported Performance Data

Product performance data derived from customer-initiated complaints and returned products is referred to as Customer Reported Performance Data. While Abbott strongly encourages the submission of any relevant complaints and product returns; this data is not proactively solicited or regularly monitored like data from the Post-Approval studies. Underreporting of events within customer reported performance data is recognized throughout our industry. Abbott is constantly improving the accuracy and utility of the data within this Product Performance Report.

SUMMARY INFORMATION

The Customer Reported Performance Data page for each model or model group includes a table of model-specific information. Several terms from this table that are relevant to performance data calculations are defined below:

Registered U.S. Implants - The total number of U.S. implanted devices for which patient and device information has been provided to Abbott. This total includes devices which have been explanted or are otherwise out of service.

Estimated Active U.S. Implants - The total number of U.S. registered implants that have not been identified to Abbott as explanted or otherwise out of service. An adjustment is made to account for the underreporting of patient mortality. Abbott performed an analysis of the data gathered from multiple clinical studies including some Abbott sponsored studies to determine the mortality rate within the pacemaker and ICD patient population and has factored this into the estimation of the number of active U.S. implants.

Estimated Longevity - The estimated number of years in which a device is expected to reach its Elective Replacement Indicator (ERI), as stated in the product literature. The estimate is based on battery life approximations and empirical battery performance distributions. It is strongly affected by many factors such as programmed parameters, percentage of time paced, internal impedance, etc. For example, the 9.2 year estimated longevity of an Accent™ DR model PM2110 pacemaker is based on the mean longevity (or 50%) value in the product literature corresponding to pacing at 60 ppm, 2.5V dual-chamber output at 0.4 ms pulse width, 500 ohm lead impedance, 100% DDD pacing, and Stored EGMs On. Actual performance can vary considerably, depending on the actual programmed settings and operations.

Normal Battery Depletion - The condition where a returned device met its electrical specification and reached its elective replacement indicator voltage (1) with an implant duration meeting or exceeding the nominal predicted longevity at default shipped settings, or (2) with an implant duration exceeding 75% of its estimated longevity, based on longevity calculations using information from device usage and the actual device settings. The quantity of normal battery depletions reported is determined directly from laboratory analysis and does not represent any adjustment to account for underreporting.

Introduction and Overview

SURVIVAL CALCULATION GENERAL METHODS

For ICDs, pacemakers, ICMs, and leads, we compile cumulative survival data based on the actuarial (or life-table) method of survival analysis, consistent with ISO 5841-2:2014(E) “Reporting of Clinical Performance of Populations of Pulse Generators and Leads” and the 2009 AdvaMed document “[Industry Guidance for Uniform Reporting of Clinical Performance of Cardiac Rhythm Management Pulse Generators and Leads](#)”. Product performance is plotted over a maximum range of 20 years, with a minimum of 500 registered implants required for inclusion in the report, and a minimum sample size for each reported time period of 200 devices. “Survival” refers to the proper function of the device, not the survival of the patient, and is intended to illustrate the calculated probability of device survival at a given point in time. A survival probability of 99% at five years, for example, indicates that at five years after implant, the system has a 1% risk of incurring a malfunction and/or normal battery depletion. All domestically implanted devices within each model family are included in the calculations.

With the large size of the U.S. data pool, and the same products are generally used both in the U.S. and internationally, we consider the data in this report to accurately represent each device’s performance, regardless of where in the world it was implanted.

The ISO 5841-2:2014(E) “Reporting of Clinical Performance of Populations of Pulse Generators and Leads” was revised in August 2014. The revision clarified survivor definitions and reporting methods, further standardizing product performance reporting across the cardiac rhythm management implantable device and lead manufacturers.

This revision of the ISO standard specifically excludes lead malfunctions confirmed through returned product analysis which were received with no accompanying complaint from the survival probability calculations. However, to provide the highest level of transparency, Abbott continues to include malfunctions not associated with a complaint in the survival probability calculations and in the tabular display of laboratory-confirmed malfunctions.

ICD, PACEMAKER, AND ICM SURVIVAL ANALYSIS

The data used for the analysis of ICDs, pacemakers, and ICMs includes up-to-date device registration information and the laboratory analysis of all domestically implanted devices returned to Abbott. The analysis measures device performance to specification, and does not reflect medical complications, such as infection, erosion, muscle stimulation or inhibition, or units implanted for fewer than 24 hours.

In accordance with ISO 5841-2:2014(E), the survival calculations for ICDs, pacemakers, and ICMs are adjusted to reduce the bias caused by underreporting of malfunctions and normal battery depletions. Abbott compared the malfunctions and normal battery depletion rates calculated from our actively monitored populations to the rates calculated from our passively monitored populations and have adjusted the survival calculations accordingly.

Survival data are presented in a single table and graph. The survival data is separated into “Including Normal Battery Depletion” and “Excluding Normal Battery Depletion” categories. The “Including Normal Battery Depletion” data reflects the frequency of device removal due to normal battery depletion and malfunction of any type. The “Excluding Normal Battery Depletion” category reflects the frequency of device removal due to malfunctions only.

Introduction and Overview

ICD, PACEMAKER, AND ICM MALFUNCTION REPORTING

The quantity and rate of malfunctions recorded for each ICD, pacemaker, and ICM model are presented in a tabular format on both the Customer Reported Performance Data and Actively Monitored Study Data pages. The root cause of all laboratory-confirmed malfunctions is classified into one of eight categories: Electrical Component, Electrical Interconnect, Battery, High Voltage Capacitor (ICDs and CRT-Ds only), Software/Firmware, Mechanical, Possible Early Battery Depletion, or Other. Note that in the rare cases where multiple malfunctions are identified in a single device, a single malfunction category will be selected with priority given in the order of the list above. Consistent with previous performance reports, ICD and Pacemaker malfunctions are further classified as with or without compromised therapy.

Malfunction Definitions

Malfunction - Having characteristics that are outside the performance limits established by the manufacturer while implanted and in service, as confirmed by laboratory analysis, except changes to characteristics due to normal battery depletion or induced malfunction. Device damage caused after or during explant is not considered a malfunction. Note that lead-related malfunctions of a pacemaker or ICD system are assigned to the lead.

Malfunction with Compromised Therapy - The condition when a device is found to have “malfunctioned,” as defined above, in a manner that compromised pacing or defibrillation therapy (including complete loss or partial degradation) while implanted and in service. Therapy is considered to have been compromised if no therapy is available or critical patient-protective pacing or defibrillation therapy is not available.

A malfunction with compromised therapy does not imply that a patient has actually experienced a serious complication or death as a result of the malfunction although it does imply that the potential for a serious complication or death did exist during the period of the malfunction.

Malfunction without Compromised Therapy - The condition when a device is found to have “malfunctioned,” as defined above, in a manner that did not compromise pacing or defibrillation therapy while implanted and in service, as confirmed by laboratory analysis. Therapy is not compromised as long as the critical patient-protective pacing and defibrillation therapies are available. Changes in device settings that occur as intended by the design (for example, reversion to a designed Safe Mode) that do not result in loss of critical patient-protective therapies but are the reported reasons for explant are categorized as a Malfunction without Compromised Therapy.

Malfunction Root Cause Category Definitions

Electrical Component - Findings linked to electrical components such as integrated circuits, resistors, low voltage capacitors, diodes, etc. Does not include high voltage capacitors or batteries as those are separately listed.

Electrical Interconnect - Findings linked to the connections between electrical components such as wires, solder joints, wire bonds, feedthroughs, etc.

Battery - Findings linked to the battery and its components.

High Voltage Capacitor - Findings linked to the high voltage capacitor and its components.

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Software/Firmware - Findings linked to software or firmware function.

Mechanical - Findings linked to mechanical components such as headers, setscrews, fluid seals, internal supports, the hermetic case, etc.

Possible Early Battery Depletion - Findings where the actual reported implant time is less than 75% of the expected longevity calculated using the available device setting information and no root cause was able to be identified. Additionally, in the absence of a specific root cause finding, returned devices with insufficient device setting information to determine conclusively if battery depletion was normal or premature are conservatively classified as Possible Early Battery Depletion malfunctions.

Other - Findings linked to other components such as packaging and accessories, and findings where analysis is inconclusive, as well as other complications not included above.

LEADS SURVIVAL ANALYSIS

Implanted cardiac leads are subjected to constant, complex flexural and torsional forces, interactions with other leads and/or the pulse generator device, plus other forces associated with cardiac contractions, patient physical activity, posture, and anatomy. Therefore, the functional lifetime of cardiac leads is limited and cannot be predicted with a high degree of confidence. Understanding these limitations, survival estimates are provided for all leads included in this report.

The data used for the survival analysis of leads includes up-to-date lead registration information, chronic complications (>30 days) reported by the field, and the laboratory analysis of all domestically implanted leads returned to Abbott. Complaints reported within 30 days of implant (acute observations), are considered to be related to factors other than lead malfunction, such as patient specific characteristics or implant technique, and are therefore excluded from the survival calculations, consistent with industry practice. If there is laboratory data that determines the lead to have exhibited a malfunction, and the lead is known to have been implanted for more than 24 hours, the lead is counted as a non-survivor. If a lead is the subject of a complaint report, and was implanted for more than 30 days (chronic complication), then the lead is counted as a non-survivor. These criteria are also followed for partial lead returns. This method for non-returned complications is used to ensure a conservative failure estimate for lead performance. Chronic complications commonly associated with non-returned leads and partial lead returns include, but are not limited to, reports of sensing, pacing, and capture anomalies, perforation, and dislodgement.

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LEADS OBSERVATION AND COMPLICATION REPORTING

Reporting for recently released lead models provides detail on specific chronic complications (more than 30 days implant), as well as acute observations (post implant to 30 days), that are reported to Abbott as complaints. Each complication and observation is categorized into one of the eleven categories below, irrespective of whether the lead has been returned for analysis. The quantity and rate of each complication and observation type is provided in a tabular format on the Customer Reported Performance Data page. Note that in the rare cases where multiple complaints are identified for a single lead, a single category will be selected with priority given in the order of the list below.

Cardiac Perforation - Penetration of the lead tip through the myocardium, clinically suspected and confirmed by chest x-ray, fluoroscopy, echocardiogram, or visual observation, which results in clinical symptoms, typically degradation of pacing/ICD lead electrical performance (high thresholds), chest pain, or tamponade.

Conductor Fracture - A mechanical break within a lead conductor (includes connectors, coils, cables and/or electrodes) observed visually, electrically, or radiographically.

Lead Dislodgement - Radiographic, electrical or electrocardiographic evidence of electrode displacement from the original implant site or electrode displacement that adversely affects pacing and/or lead performance.

Failure to Capture - Intermittent or complete failure to achieve cardiac stimulation (atrial or ventricular) at programmed output delivered outside of the cardiac refractory period. A sudden and significant increase in the pacing threshold value (elevated thresholds compared to previous measured value) at which 2:1 safety margin can no longer be achieved.

Oversensing - Misinterpretation of cardiac or non-cardiac events as cardiac depolarization, e.g. T-waves, skeletal muscle potentials, and extracardiac electromagnetic interference (EMI).

Failure to Sense (undersensing) - Intermittent or complete loss of sensing or failure to detect intended intrinsic cardiac signals (atrial or ventricular) during non-refractory periods at programmed sensitivity settings.

Insulation Breach - A disruption or break in lead insulation observed visually, electrically, or radiographically.

Abnormal Pacing Impedance - Pacing impedance is typically considered abnormal if a measurement is $< 200 \Omega$ or $> 2000 \Omega$ (based on lead model and measurement range of the device).

Abnormal Defibrillation Impedance - Defibrillation impedance is typically considered abnormal if a measurement is $< 20 \Omega$ or $> 200 \Omega$ (based on lead model and measurement range of the device).

Extracardiac Stimulation - Clinical observation of inadvertent nerve/muscle stimulation other than cardiac muscle.

Other - Specific proprietary lead mechanical attributes such as lead incorporated sensors, connectors or seal rings which affect a lead's ability to perform as designed or remain in service, as well as other complications not included above.

Introduction and Overview

LEADS MALFUNCTION REPORTING

As a supplement to the survival estimates, the categorization of lead malfunctions emphasizes the root cause of malfunction rather than a functional longevity prediction. In accordance with AdvaMed guidelines, laboratory analysis results of returned leads are categorized into one of the following five categories of malfunctions. The quantity and rate of each malfunction type is provided in a tabular format on the Customer Reported Performance Data and the Actively Monitored Study Data pages. Note that in the rare cases where multiple malfunctions are identified in a single lead, a single malfunction category will be selected with priority given in the order of the list below. The definition for each malfunction type is provided below:

Conductor Fracture - Conductor break with complete or intermittent loss of continuity that could interrupt current flow. This type of malfunction includes any conductor fracture such as those associated with flex-fatigue or clavicular crush damage.

In an effort to further increase customer understanding of Abbott defibrillation and left-heart lead performance, subcategories of conductor fracture are also provided. The definitions of these subcategories are provided below:

Clavicular Crush - Conductor fracture due to strong compression and bending at the approximation of the first rib and clavicle.

In the Pocket - Conductor fracture not within the vascular or cardiac systems, typically within the subcutaneous pocket or associated with the suture sleeve, excluding the mechanism of clavicular crush.

Intravascular - Conductor fracture within the vascular or cardiac systems.

Insulation Breach - Any lead insulation breach, such as: 1) proximal abrasion associated with lead-to-lead or lead-to-can contact in the pocket, 2) mid-lead insulation damage caused by clavicular crush or insulation wear in the region of vein insertion, 3) distal abrasion due to lead-to-lead interactions or contact with anatomic structures, and 4) externalized conductors in the distal region.

Subcategories of insulation breach for defibrillation and left-heart leads are also provided. The definitions of these subcategories are provided below:

Lead-to-Can Contact - Direct contact between the lead and the can (i.e. pacemaker, ICD, or CRT-D) combined with repetitive skeletal movement caused abrasion that resulted in a full thickness outer insulation breach.

Lead-to-Lead Contact - Repetitive contact between two leads caused abrasion that resulted in a full thickness outer insulation breach.

Clavicular Crush - Damage due to strong compression between the first rib and clavicle resulted in a full thickness outer insulation breach.

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Externalized Conductors - Abrasion resulted in an outer insulation breach within the vascular or cardiac systems allowing the normally contained conductors to become visible outside the lead body. Externalized conductors were described in our December 2010 and November 2011 communications regarding insulation abrasion failures on silicone Riata™ and Riata™ ST lead families (summary on pages 343-344) and in our April 2012 communication regarding insulation abrasion failures on QuickSite™ and QuickFlex™ lead families. Additional information regarding externalized conductors on Riata™ and Riata™ ST leads can be found at www.RiataCommunication.com.

Other (Insulation Breach) - Insulation breaches that resulted from a failure mode not represented by the other four categories. This includes a variety of failure modes, such as damage at the suture sleeve and contact with patient anatomy. Also includes insulation breaches for which analysis was unable to isolate a specific cause.

Crimps, Welds and Bonds - Any interruption in the conductor or lead body associated with a point of connection.

Other - Includes specific proprietary lead mechanical attributes, such as lead incorporated sensors, connectors, and seal rings, as well as other analysis results not included in the alternate categories.

Extrinsic Factors - The lead was implanted greater than 30 days, removed from service with an associated complaint and returned for analysis, however analysis was inconclusive because (1) only portions of the lead were available, or (2) the returned lead was damaged by the explantation process, or (3) lab analysis could not determine an out of specification condition (typically with complaints such as dislodgements, perforations, or failure to capture).

Actively Monitored Study Data

SUMMARY INFORMATION

Since 2007 the Product Performance Report has included data from the St. Jude Medical Product Longevity and Performance Registry (SCORE). This comprehensive study provided monitored performance data on pacemakers, ICDs, and leads. With product-specific, post-market registries being standard practice, Abbott continues to complement the SCORE registry with data from the SJ4 Post-Approval Study, the QuickFlex™ μ Lead Post-Approval Study, the Quadripolar CRT-D Post-Approval Study, and the OPTIMUM registry. These actively monitored study data now represent >62,000 implanted devices, and continues to be a very powerful source of product performance information which complements the data collected from Customer Reported Performance Data. Actively monitored study data is not susceptible to underreporting and provides the most accurate understanding of product performance. The many sites participating in these actively monitored studies are individually providing data on the performance of Abbott cardiac rhythm management products using common definitions and criteria. In addition, each of these sites is regularly audited by Abbott personnel to ensure comprehensive reporting.

Introduction and Overview

	STUDY DESCRIPTION	STUDY INITIATED	# SITES	# PATIENTS	PRODUCT TYPES/FAMILIES
SCORE (ST. JUDE MEDICAL PRODUCT LONGEVITY AND PERFORMANCE REGISTRY)	Prospective, actively monitored, multicenter registry to evaluate the long-term performance of Abbott market-released cardiac rhythm management products.	September 2007	80	11,247	Pacemakers, ICDs, CRT-Ds, Leads (all types)
SJ4 POST-APPROVAL STUDY	Prospective, actively monitored, multicenter study to evaluate the acute and chronic performance of the Abbott SJ4/DF4 connector and SJ4/DF4 defibrillation leads.	June 2009	58	1,701	ICDs, CRT-Ds, Leads (all types)
QUICKFLEX™ μ POST-APPROVAL STUDY	Prospective, actively monitored, multicenter study to evaluate the acute and chronic performance of the Abbott QuickFlex™ μ 1258T left ventricular leads.	September 2010	76	1,930	CRT-Ds, Leads (all types)
QUADRIPOlar CRT-D POST-APPROVAL STUDY	Prospective, actively monitored, multicenter study to evaluate the acute and chronic performance of the Abbott Quadripolar CRT-D system.	February 2012	71	1,970	Unify Quadra™ and Quadra Assura™ CRT-Ds, Leads (all types)
OPTIMUM REGISTRY	Prospective, actively monitored, multicenter registry to evaluate the long-term performance of market-released Abbott leads with Optim™ insulation material.	August 2006	241	14,120	Leads (any model with Optim™ Insulation)

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The models included in the actively monitored dataset are listed below:

ICDS

Current™ + DR (Model CD2211-36)
Current™ + DR (Model CD2211-36Q)
Current™ + VR (Model CD1211-36Q)
Current™ DR RF (Model 2207-36)
Current™ VR RF (Model 1207-36)
Fortify™ DR (Model CD2231-40)
Fortify™ DR (Model CD2231-40Q)
Fortify™ VR (Model CD1231-40Q)
Promote™ + CRT-D (Model CD3211-36)
Promote™ + CRT-D (Model CD3211-36Q)
Promote™ RF CRT-D (Model 3207-36)
Quadra Assura™ CRT-D (Model CD3265-40)*
Quadra Assura™ CRT-D (Model CD3265-40Q)
Quadra Assura™ CRT-D (Model CD3365-40Q)
Unify Assura™ CRT-D (Model CD3357-40C)
Unify Assura™ CRT-D (Model CD3357-40Q)
Unify Quadra™ CRT-D (Model CD3249-40)
Unify Quadra™ CRT-D (Model CD3249-40Q)
Unify™ CRT-D (Model CD3231-40)
Unify™ CRT-D (Model CD3231-40Q)

DEFIBRILLATION LEADS

Durata™ (Model 7122)
Durata™ (Models 7120/7121)
Durata™ DF4 (Model 7122Q)
Durata™ DF4 (Models 7120Q/7121Q)
Durata™ DF4 (Models 7170Q/7171Q)
Riata™ (Models 1580/1581)
Riata™ ST (Models 7000/7001)
Riata™ ST Optim™ (Models 7020/7021)
Riata™ ST Optim™ (Models 7070/7071)

CRT LEADS

Quartet™ (Model 1458Q)
QuickFlex™ (Model 1156T)
QuickFlex™ XL (Model 1158T)
QuickFlex™ μ (Model 1258T)
QuickSite™ (Model 1056T)
QuickSite™ XL (Model 1058T)

PACEMAKERS

Accent™ DR (Model PM2110)
Accent™ DR RF (Model PM2210)
Accent™ SR RF (Model PM1210)
Anthem™ RF CRT-P (Model PM3210)
Identity ADx™ XL DR (Model 5386)
Victory™ XL DR (Model 5816)
Zephyr™ DR (Model 5820)
Zephyr™ XL DR (Model 5826)
Zephyr™ XL SR (Model 5626)

PACING LEADS

IsoFlex™ Optim™ (Model 1944)
IsoFlex™ Optim™ (Model 1948)
IsoFlex™ S (Model 1646)
OptiSense™ (Model 1699)
OptiSense™ (Model 1999)
Tendril™ (Model 1782)
Tendril™ (Model 1788)
Tendril™ SDX (Model 1488)
Tendril™ SDX (Model 1688)
Tendril™ ST Optim™ (Model 1882)
Tendril™ ST Optim™ (Model 1888)
Tendril™ STS (Model 2088)

**New for 2018 Second Edition*

Introduction and Overview

QUALIFYING COMPLICATIONS

When abnormal performance is suspected of an actively monitored study device, the related clinical event and any resulting clinical action is reported to Abbott. A Qualifying Complication is defined to have occurred if the report identifies one of the following Clinical Events that resulted in one of the following Clinical Actions. Any Clinical Event without a related Clinical Action is not considered a Qualifying Complication.

QUALIFYING CLINICAL EVENTS

- Abnormal Defibrillation Impedance
- Abnormal Pacing Impedance
- Cardiac Perforation
- Conductor Fracture
- Extracardiac Stimulation
- Failure to Capture
- Failure to Sense
- Inappropriate Shock
- Insulation Breach
- Lead Dislodgement
- Loss of Telemetry
- Oversensing
- Pericardial Effusion
- Premature Battery Depletion
- Skin Erosion

QUALIFYING CLINICAL ACTION

- Generator Pacing Mode Changed
- Lead Electrically Abandoned/Capped
- Lead/Generator Explanted
- Lead/Generator Replaced
- Lead Polarity Changed
- Lead Surgically Abandoned/Capped
- Lead Surgically Repositioned

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SURVIVAL CALCULATION METHODS

Survival calculations for actively monitored studies are made in a manner consistent with the ISO 5841-2:2014(E) method used for Customer Reported Performance Data. A minimum of 100 devices are required to have been enrolled, with the latest interval to be reported having a minimum of 50 devices which have been followed for at least six months. Any device with a Qualifying Complication is defined as a non-survivor. Consistent with industry practice, Qualifying Complications for leads are included in the survival calculations for events with an implant duration greater than 30 days. For pacemakers and ICDs, Qualifying Complications are included in the survival calculations for events with an implant duration more than 24 hours. Medical complications unrelated to device performance are not considered as Qualified Complications. Devices included in the actively monitored studies are excluded from the Customer Reported Performance Data. Certain devices and leads, including any which transferred from Customer Reported Performance Data into Actively Monitored Study Data are subsequently excluded from the Customer Reported Performance Data and subject to these Survival Calculation methods. Actively monitored study performance data includes both advisory and non-advisory devices.

MALFUNCTION REPORTING

The Actively Monitored Study Data page contains a table of all device malfunctions. The type, quantity, and rate of all laboratory-confirmed malfunctions are listed using the same categories reported in Customer Reported Performance Data. The malfunction data is not utilized in the actively monitored study survival calculations, but does provide important supplementary information about product performance and reliability.

Medical Advisory Board Review

Abbott has an established and independent Medical Advisory Board (MAB) focused on cardiac rhythm management systems, including pulse generators and leads. One of the important tasks assigned to the MAB is the review of the performance data contained in this report prior to its release and publication on a semi-annual basis. MAB members and their location of practice include:

Dr. Anne Curtis, Buffalo, New York

Dr. Roger Freedman, Salt Lake City, Utah

Dr. Christoph Geller, Bad Berka, Germany

Dr. Thomas Mattioni, Paradise Valley, Arizona

Dr. Raymond Schaerf, Burbank, California

Dr. Bruce Wilkoff, Cleveland, Ohio

Returning Devices to Abbott

To maintain the continued accuracy of our performance reporting, Abbott strongly encourages physicians to notify our Patient Records department (888-SJM-2763) each time a device is removed from service for any reason. Additionally, all explanted products are requested to be returned to Abbott for laboratory evaluation whether or not a malfunction is suspected. To facilitate the return of explanted devices, Abbott offers a no-cost Returned Products Kit comprised of a postage paid explant box with a shipping address label, a removed device information form, a biohazard bag, and biohazard labels to seal the explant box. This kit, #N0004, can be ordered free of charge by contacting Abbott Customer Service (888-SJM-2763).

Contact Us

The Abbott team is always ready to respond to questions, comments or suggestions as well as receive product performance feedback. You can reach us by phone at 888-SJM-2763, on the web at www.SJM.com, or by contacting your local Abbott representative.

Cardiac Resynchronization Therapy (CRT) ICDs

Cardiac Resynchronization Therapy (CRT) ICDs

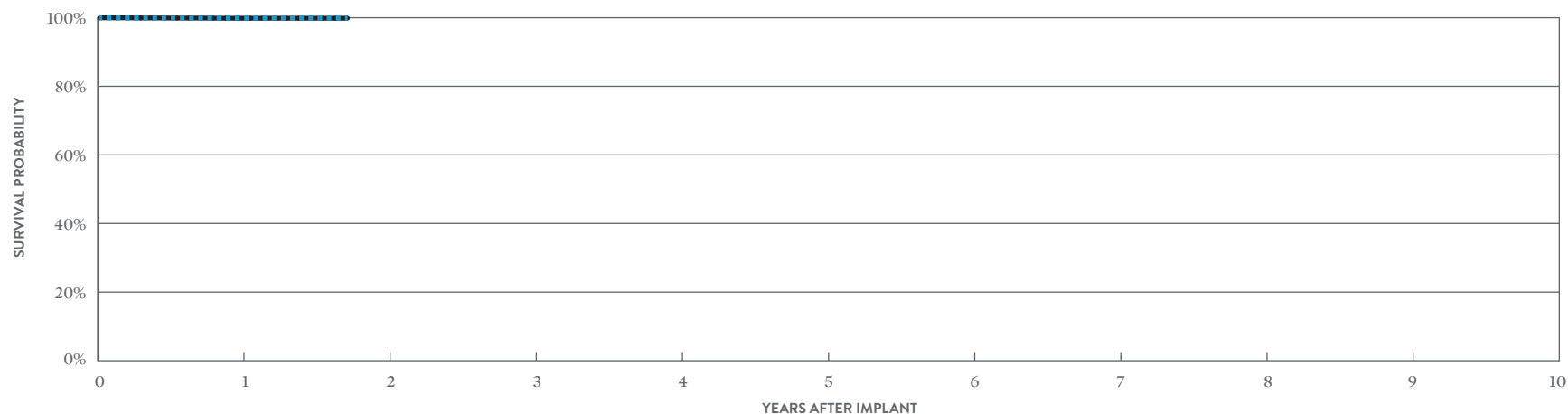
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura MP™ CRT-D

MODEL CD3369-40Q*

US Regulatory Approval	February 2016
Registered US Implants	22,164
Estimated Active US Implants	19,995
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	0
Max. Delivered Energy	40 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	1	<0.01%
Electrical Interconnect	3	0.01%	0	0.00%
Battery	0	0.00%	1	<0.01%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	3	0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	5	0.02%	5	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	AT 21 MONTHS
SURVIVAL PROBABILITY	99.86%	99.86%
± 1 STANDARD ERROR	0.03%	0.03%
SAMPLE SIZE	14,980	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	AT 21 MONTHS
SURVIVAL PROBABILITY	99.86%	99.86%
± 1 STANDARD ERROR	0.03%	0.03%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

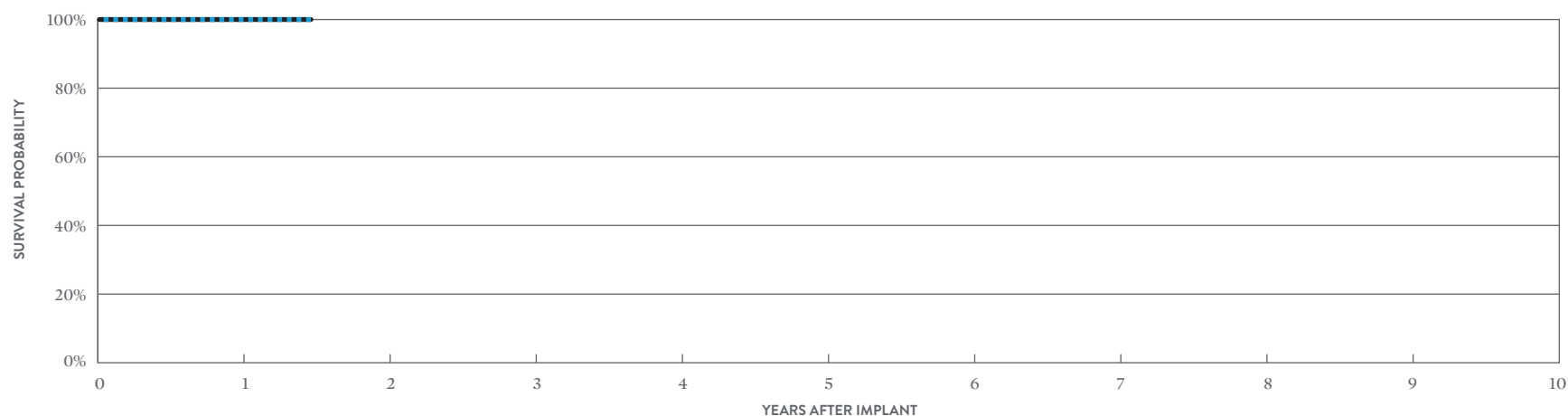
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura MP™ CRT-D

MODEL CD3369-40C*

US Regulatory Approval	February 2016
Registered US Implants	3,217
Estimated Active US Implants	2,906
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	0
Max. Delivered Energy	40 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	AT 18 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%
SAMPLE SIZE	2,190	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	AT 18 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%

*Parylene coating.

Cardiac Resynchronization Therapy (CRT) ICDs

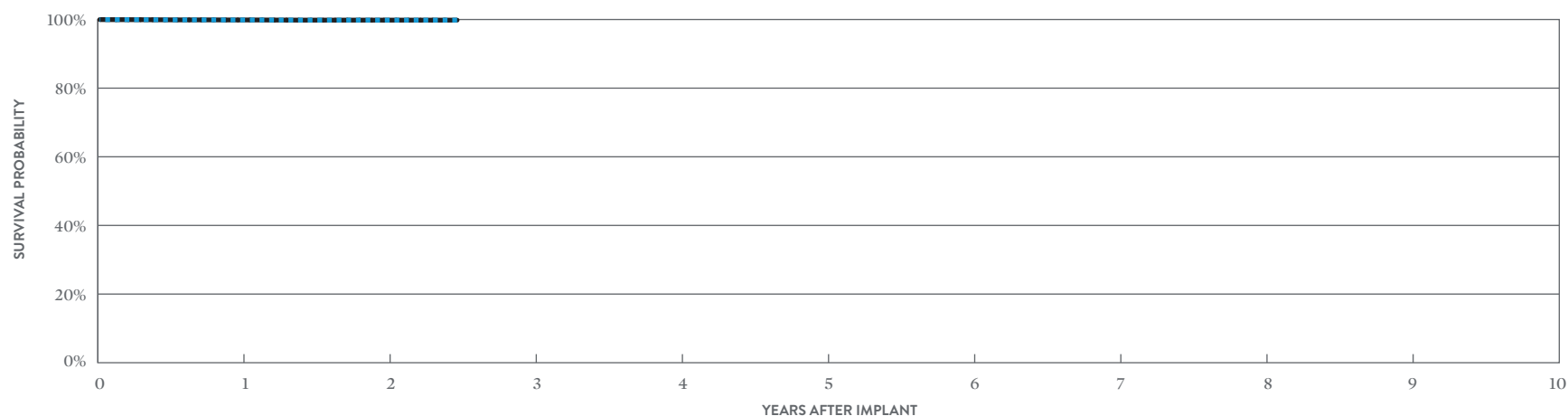
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura™ CRT-D

MODEL CD3365-40Q* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	14,525
Estimated Active US Implants	12,025
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	0
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	3	0.02%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	1	<0.01%	0	0.00%
Mechanical	0	0.00%	4	0.03%
Possible Early Battery Depletion	1	<0.01%	0	0.00%
Other	1	<0.01%	1	<0.01%
Total	6	0.04%	5	0.03%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 30 MONTHS
SURVIVAL PROBABILITY	99.87%	99.82%	99.82%
± 1 STANDARD ERROR	0.03%	0.04%	0.04%
SAMPLE SIZE	13,070	7,970	360

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 30 MONTHS
SURVIVAL PROBABILITY	99.87%	99.82%	99.82%
± 1 STANDARD ERROR	0.03%	0.04%	0.04%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

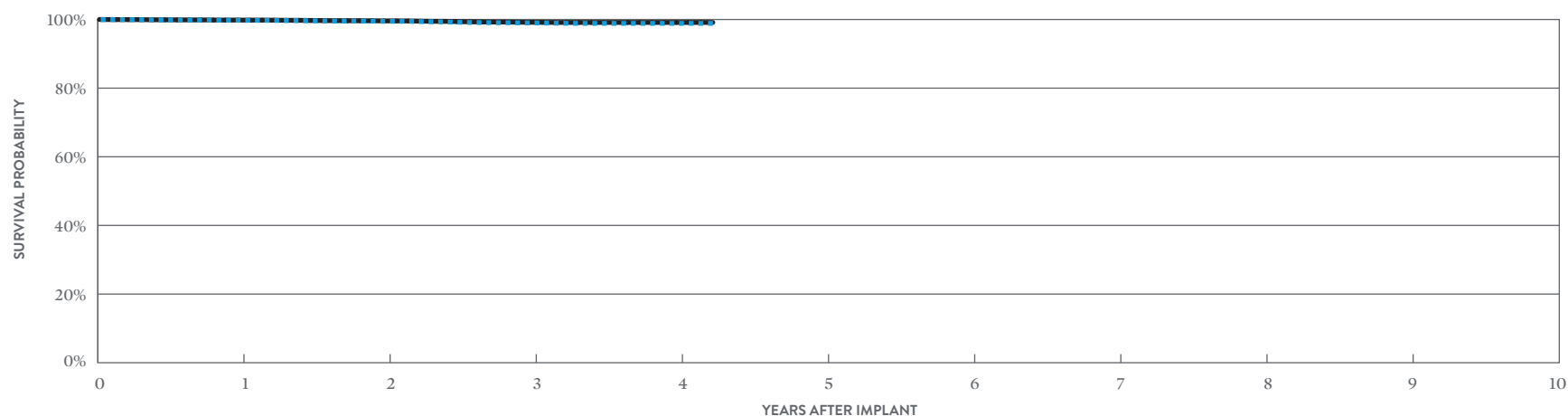
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura™ CRT-D

MODEL CD3365-40Q* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	24,082
Estimated Active US Implants	15,742
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	17
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	4	0.02%	7	0.03%
Electrical Interconnect	7	0.03%	0	0.00%
Battery	2	<0.01%	4	0.02%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	1	<0.01%	3	0.01%
Mechanical	0	0.00%	2	<0.01%
Possible Early Battery Depletion	12	0.05%	14	0.06%
Other	5	0.02%	2	<0.01%
Total	31	0.13%	32	0.13%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 51 MONTHS
SURVIVAL PROBABILITY	99.78%	99.47%	99.00%	98.87%	98.87%
± 1 STANDARD ERROR	0.03%	0.05%	0.07%	0.08%	0.08%
SAMPLE SIZE	22,660	19,860	14,190	5,470	460

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 51 MONTHS
SURVIVAL PROBABILITY	99.83%	99.59%	99.19%	99.14%	99.14%
± 1 STANDARD ERROR	0.03%	0.04%	0.07%	0.07%	0.07%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

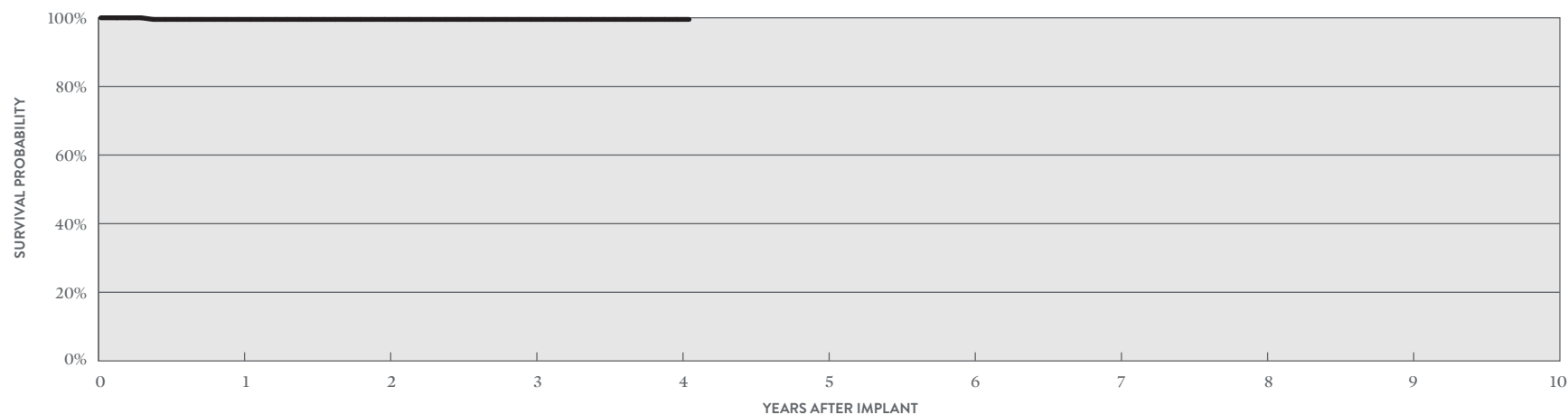
Quadra Assura™ CRT-D

MODEL CD3365-40Q*

US Regulatory Approval	June 2013
Number of Devices Enrolled in Study	222
Active Devices Enrolled in Study	129
Cumulative Months of Follow-up	6,580
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Skin Erosion	1	0.45%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	AT 49 MONTHS
SURVIVAL PROBABILITY	99.53%	99.53%	99.53%	99.53%	99.53%
± 1 STANDARD ERROR	0.47%	0.47%	0.47%	0.47%	0.47%
SAMPLE SIZE	210	160	110	80	60

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

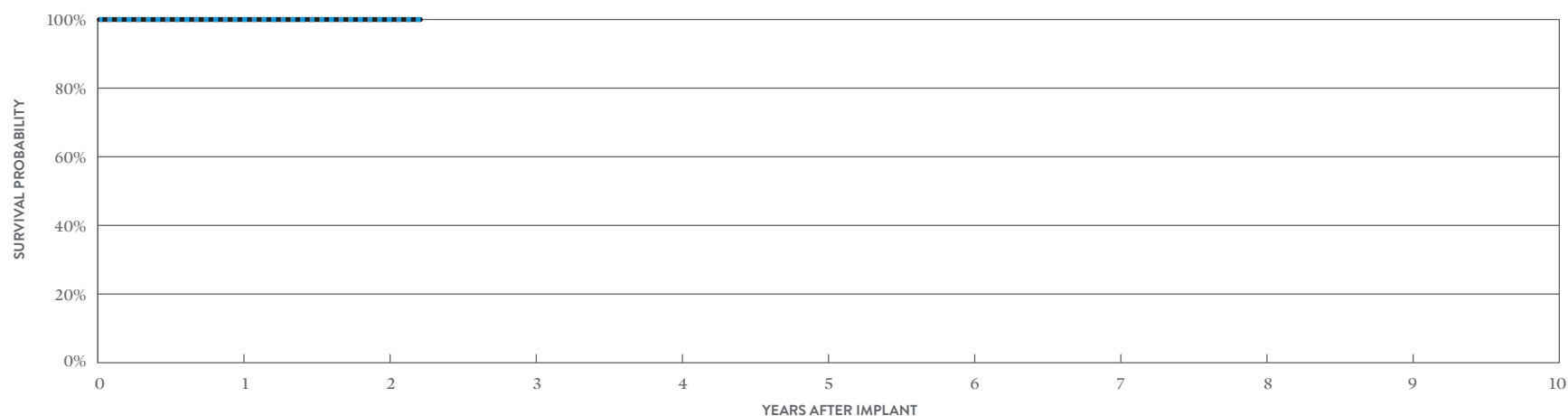
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura™ CRT-D

MODEL CD3365-40C* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	2,315
Estimated Active US Implants	1,943
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	0
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 27 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%
SAMPLE SIZE	1,960	1,020	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 27 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%

*Parylene coating.

Cardiac Resynchronization Therapy (CRT) ICDs

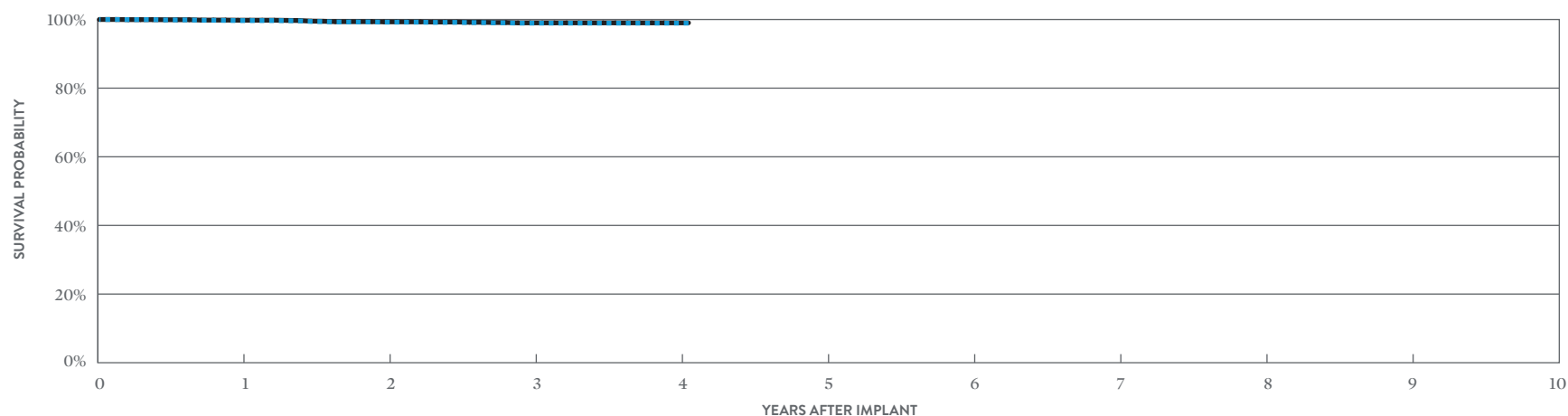
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura™ CRT-D

MODEL CD3365-40C* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	5,626
Estimated Active US Implants	3,517
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	2
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	5	0.09%	2	0.04%
Electrical Interconnect	2	0.04%	0	0.00%
Battery	1	0.02%	1	0.02%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	2	0.04%	2	0.04%
Other	2	0.04%	1	0.02%
Total	12	0.21%	6	0.11%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 49 MONTHS
SURVIVAL PROBABILITY	99.69%	99.22%	98.94%	98.94%	98.94%
± 1 STANDARD ERROR	0.07%	0.12%	0.16%	0.16%	0.16%
SAMPLE SIZE	5,250	4,400	2,940	1,110	260

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 49 MONTHS
SURVIVAL PROBABILITY	99.79%	99.32%	99.03%	99.03%	99.03%
± 1 STANDARD ERROR	0.06%	0.12%	0.16%	0.16%	0.16%

*Parylene coating.

Cardiac Resynchronization Therapy (CRT) ICDs

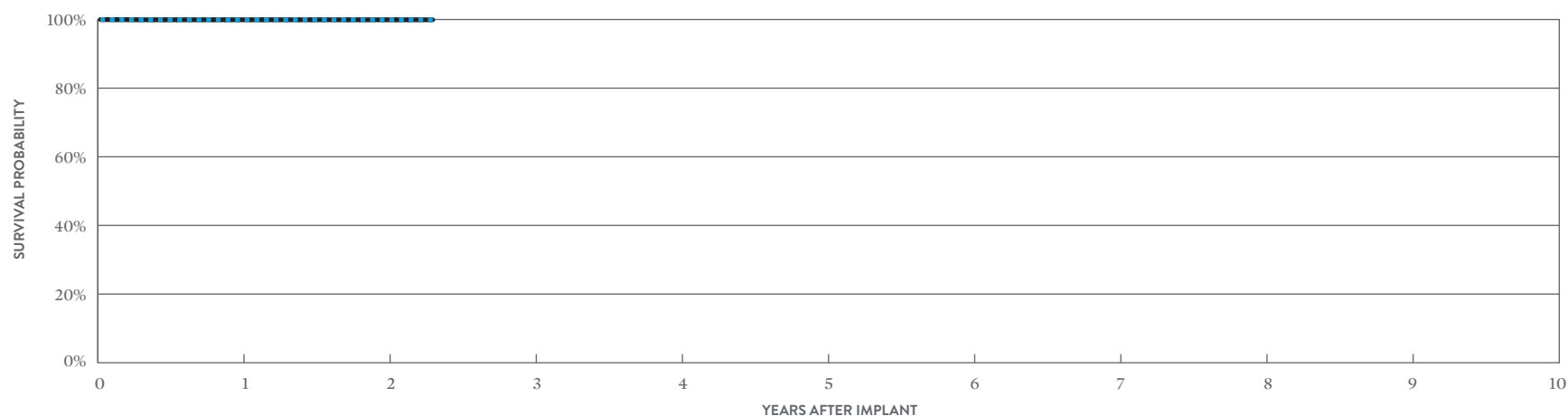
CUSTOMER REPORTED PERFORMANCE DATA

Unify Assura™ CRT-D

MODEL CD3357-40Q* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	8,366
Estimated Active US Implants	7,251
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	0
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	1	0.01%
Total	0	0.00%	3	0.04%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 28 MONTHS
SURVIVAL PROBABILITY	99.92%	99.92%	99.92%
± 1 STANDARD ERROR	0.03%	0.03%	0.03%
SAMPLE SIZE	6,320	2,480	260

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 28 MONTHS
SURVIVAL PROBABILITY	99.92%	99.92%	99.92%
± 1 STANDARD ERROR	0.03%	0.03%	0.03%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

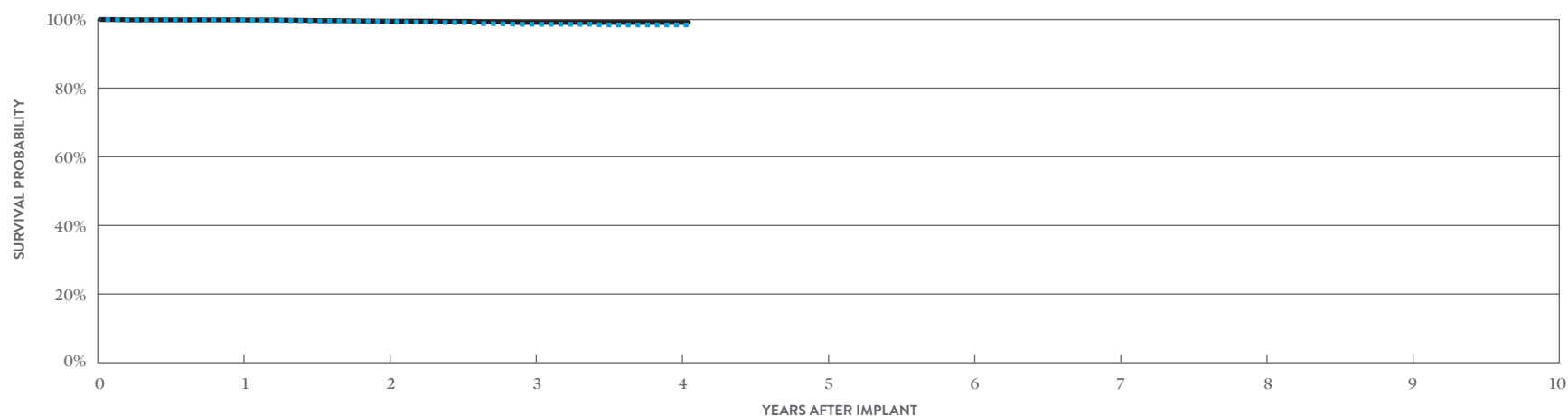
CUSTOMER REPORTED PERFORMANCE DATA

Unify Assura™ CRT-D

MODEL CD3357-40Q* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	5,343
Estimated Active US Implants	3,381
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	9
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	0.02%	0	0.00%
Electrical Interconnect	2	0.04%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	2	0.04%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	7	0.13%	1	0.02%
Other	0	0.00%	1	0.02%
Total	12	0.22%	2	0.04%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 49 MONTHS
SURVIVAL PROBABILITY	99.78%	99.40%	98.63%	98.42%	98.42%
± 1 STANDARD ERROR	0.06%	0.11%	0.21%	0.25%	0.25%
SAMPLE SIZE	4,980	4,160	2,700	960	230

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 49 MONTHS
SURVIVAL PROBABILITY	99.90%	99.52%	99.16%	99.16%	99.16%
± 1 STANDARD ERROR	0.04%	0.10%	0.16%	0.16%	0.16%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

Unify Assura™ CRT-D

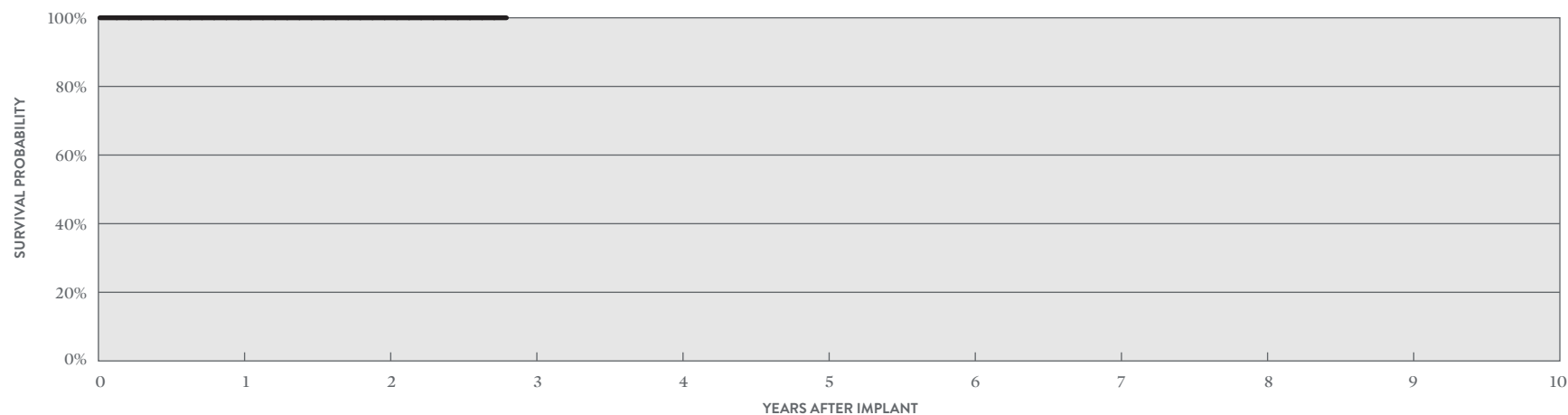
MODEL CD3357-40Q*

US Regulatory Approval	June 2013
Number of Devices Enrolled in Study	219
Active Devices Enrolled in Study	167
Cumulative Months of Follow-up	4,837
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	AT 34 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%
SAMPLE SIZE	200	130	50

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

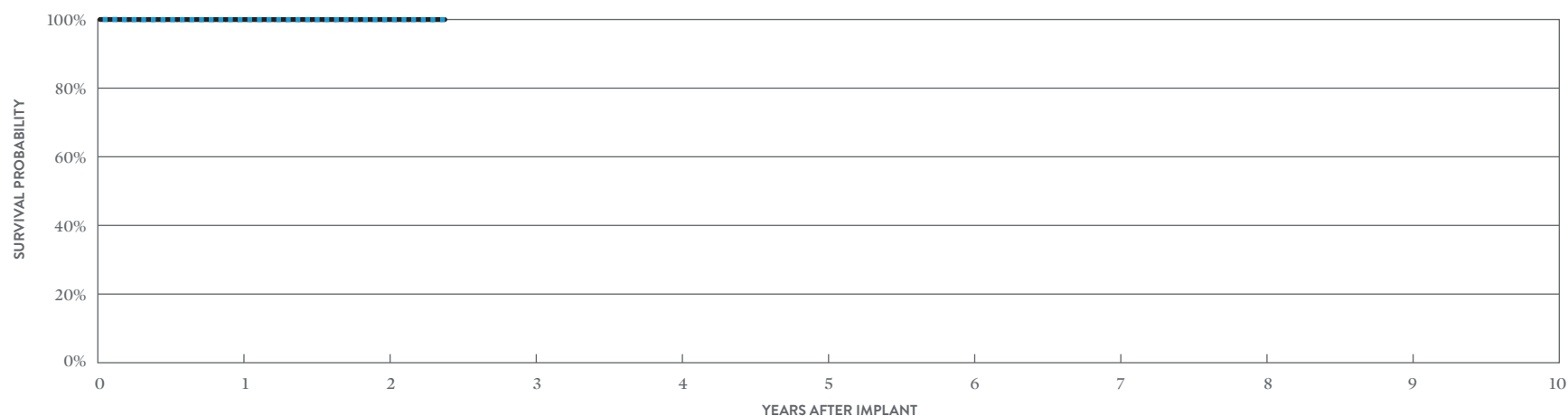
CUSTOMER REPORTED PERFORMANCE DATA

Unify Assura™ CRT-D

MODEL CD3357-40C* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	9,184
Estimated Active US Implants	7,876
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	0
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	1	0.01%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	1	0.01%	0	0.00%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	99.98%	99.98%	99.98%
± 1 STANDARD ERROR	0.02%	0.02%	0.02%
SAMPLE SIZE	7,240	3,300	320

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	99.98%	99.98%	99.98%
± 1 STANDARD ERROR	0.02%	0.02%	0.02%

*Parylene coating.

Cardiac Resynchronization Therapy (CRT) ICDs

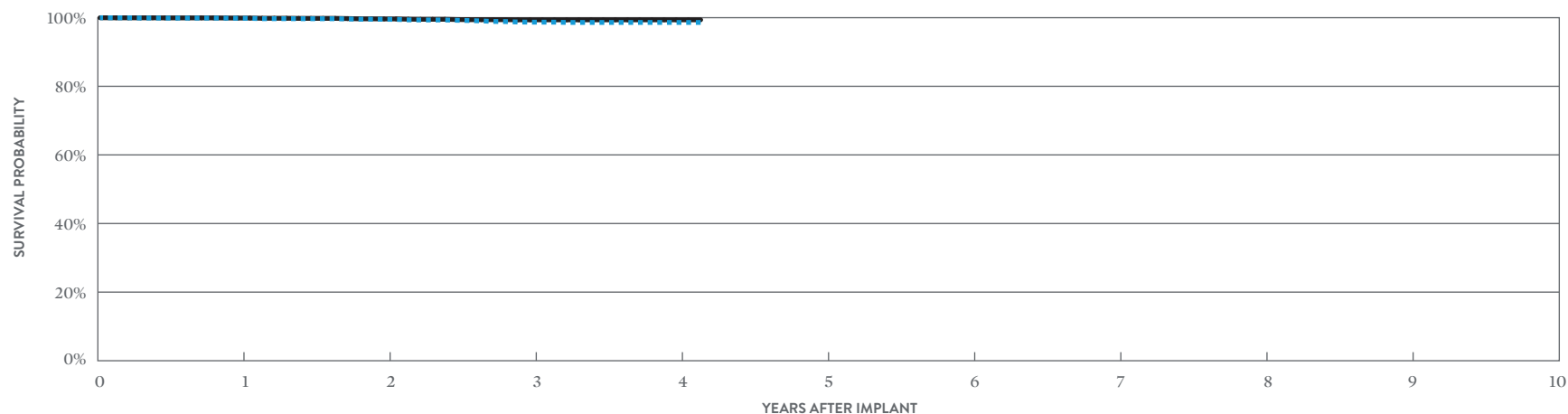
CUSTOMER REPORTED PERFORMANCE DATA

Unify Assura™ CRT-D

MODEL CD3357-40C* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	9,599
Estimated Active US Implants	6,101
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	17
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.02%	2	0.02%
Electrical Interconnect	2	0.02%	0	0.00%
Battery	0	0.00%	1	0.01%
High Voltage Capacitor	1	0.01%	0	0.00%
Software/Firmware	0	0.00%	1	0.01%
Mechanical	0	0.00%	1	0.01%
Possible Early Battery Depletion	5	0.05%	2	0.02%
Other	0	0.00%	3	0.03%
Total	10	0.10%	10	0.10%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.83%	99.53%	98.69%	98.61%	98.61%
± 1 STANDARD ERROR	0.04%	0.07%	0.14%	0.16%	0.16%
SAMPLE SIZE	9,010	7,770	5,310	1,930	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.89%	99.64%	99.38%	99.38%	99.38%
± 1 STANDARD ERROR	0.03%	0.06%	0.09%	0.09%	0.09%

*Parylene coating.

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

Unify Assura™ CRT-D

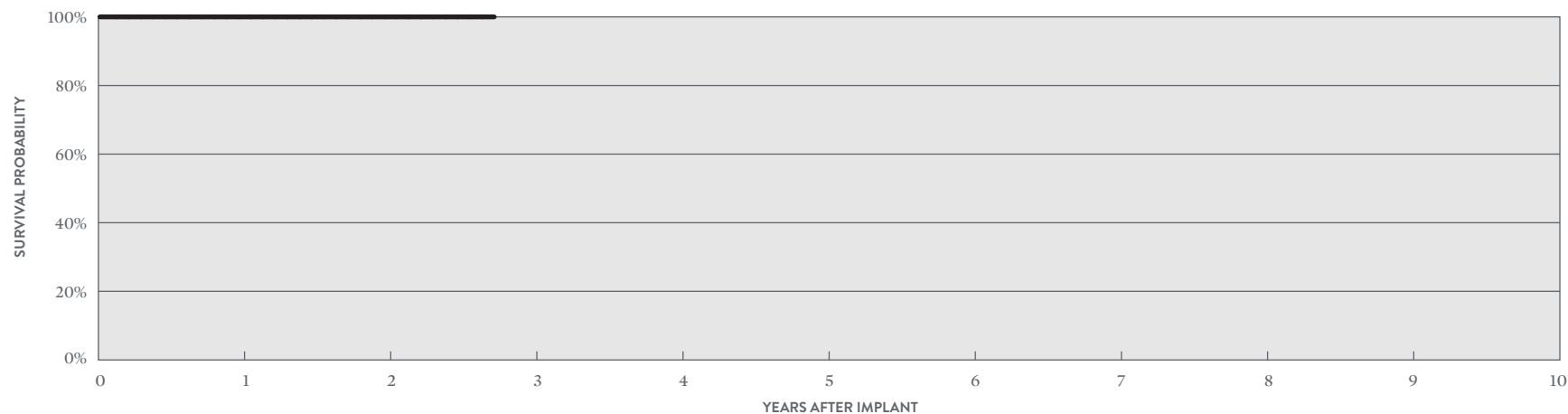
MODEL CD3357-40C*

US Regulatory Approval	June 2013
Number of Devices Enrolled in Study	185
Active Devices Enrolled in Study	115
Cumulative Months of Follow-up	4,079
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	AT 33 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%
SAMPLE SIZE	160	110	50

*Parylene coating.

Cardiac Resynchronization Therapy (CRT) ICDs

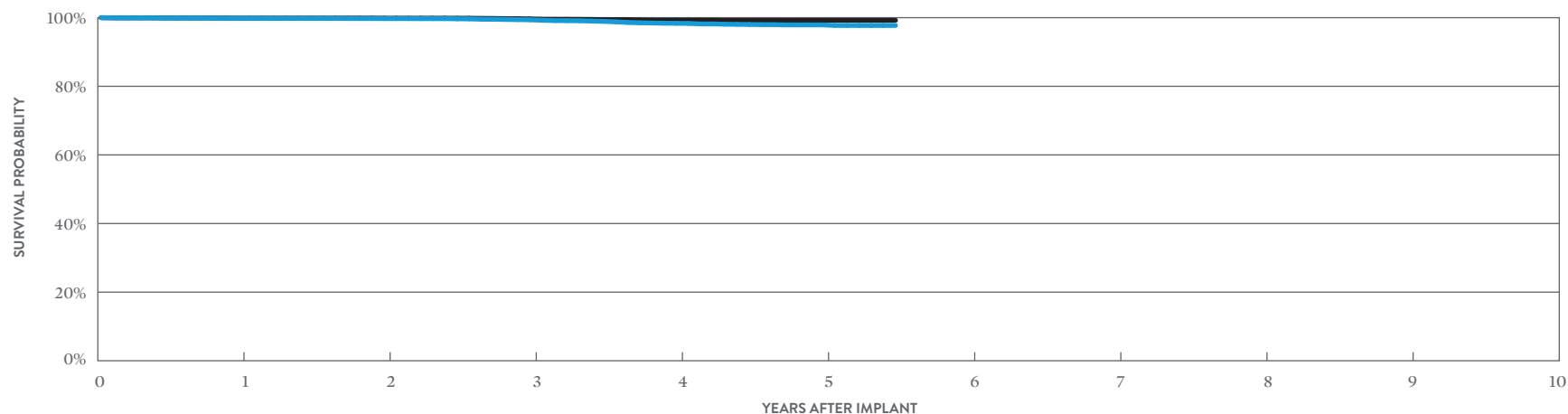
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura™ CRT-D

MODEL CD3265-40Q* (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	13,540
Estimated Active US Implants	7,742
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	46
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.01%	5	0.04%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	1	<0.01%	2	0.01%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	2	0.01%
Possible Early Battery Depletion	12	0.09%	5	0.04%
Other	1	<0.01%	0	0.00%
Total	17	0.13%	15	0.11%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 66 MONTHS
SURVIVAL PROBABILITY	99.83%	99.74%	99.38%	98.36%	97.90%	97.74%
± 1 STANDARD ERROR	0.04%	0.04%	0.07%	0.13%	0.16%	0.20%
SAMPLE SIZE	12,750	11,380	10,200	8,430	4,300	230

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 66 MONTHS
SURVIVAL PROBABILITY	99.87%	99.85%	99.64%	99.27%	99.23%	99.23%
± 1 STANDARD ERROR	0.03%	0.03%	0.05%	0.09%	0.09%	0.09%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

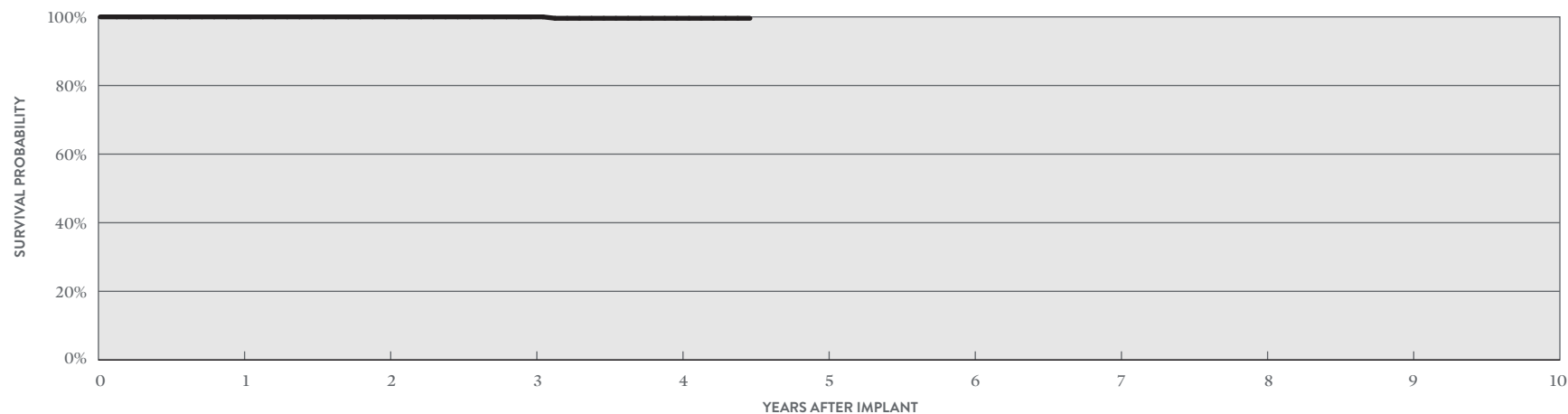
Quadra Assura™ CRT-D

MODEL CD3265-40Q*

US Regulatory Approval	May 2012
Number of Devices Enrolled in Study	421
Active Devices Enrolled in Study	182
Cumulative Months of Follow-up	14,966
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Premature Battery Depletion	1	0.24%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	1	0.24%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	1	0.24%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	AT 54 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	99.59%	99.59%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.41%	0.41%
SAMPLE SIZE	390	330	270	220	70

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

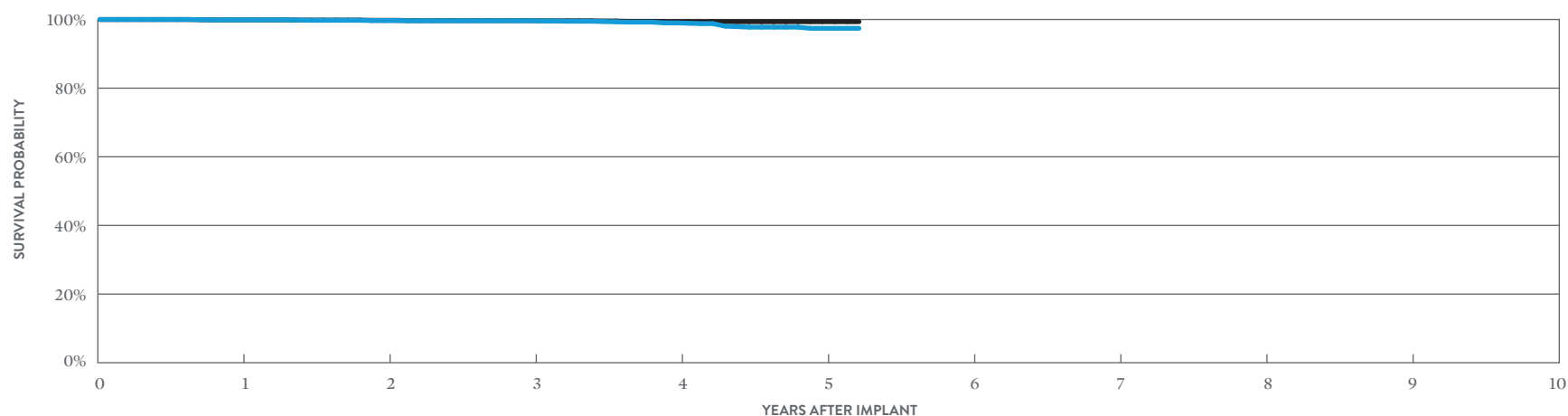
CUSTOMER REPORTED PERFORMANCE DATA

Quadra Assura™ CRT-D

MODEL CD3265-40 (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	3,926
Estimated Active US Implants	2,187
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	14
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	1	0.03%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.03%
Possible Early Battery Depletion	2	0.05%	0	0.00%
Other	4	0.10%	1	0.03%
Total	7	0.18%	2	0.05%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 63 MONTHS
SURVIVAL PROBABILITY	99.89%	99.70%	99.64%	99.01%	97.43%	97.43%
± 1 STANDARD ERROR	0.06%	0.09%	0.10%	0.19%	0.41%	0.41%
SAMPLE SIZE	3,680	3,270	2,930	2,370	1,200	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 63 MONTHS
SURVIVAL PROBABILITY	99.89%	99.76%	99.70%	99.37%	99.37%	99.37%
± 1 STANDARD ERROR	0.06%	0.08%	0.10%	0.15%	0.15%	0.15%

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

Quadra Assura™ CRT-D

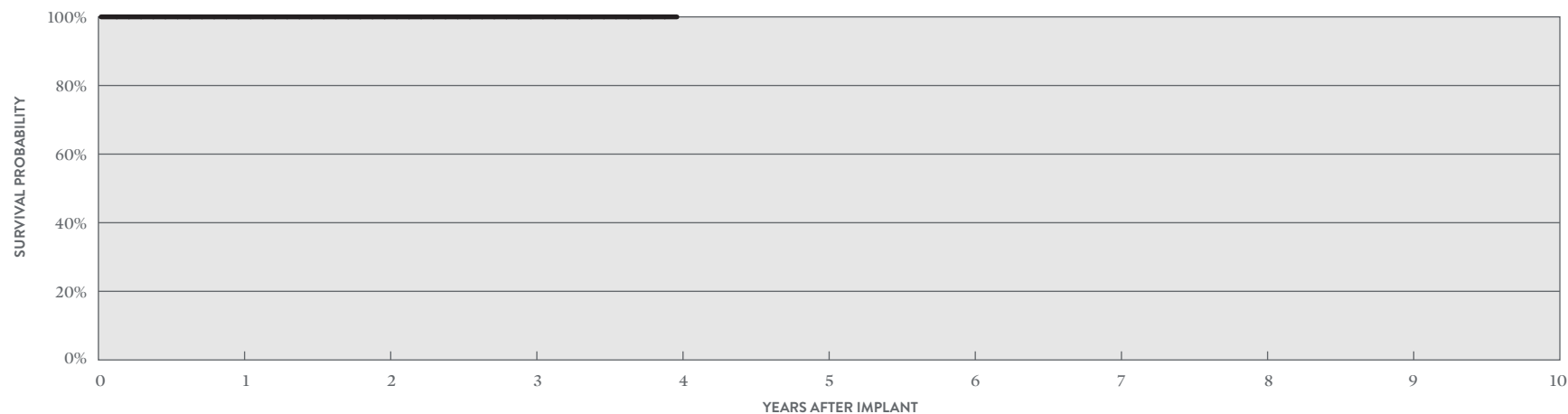
MODEL CD3265-40

US Regulatory Approval	May 2012
Number of Devices Enrolled in Study	100
Active Devices Enrolled in Study	48
Cumulative Months of Follow-up	3,649
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%
SAMPLE SIZE	100	80	60	50

Cardiac Resynchronization Therapy (CRT) ICDs

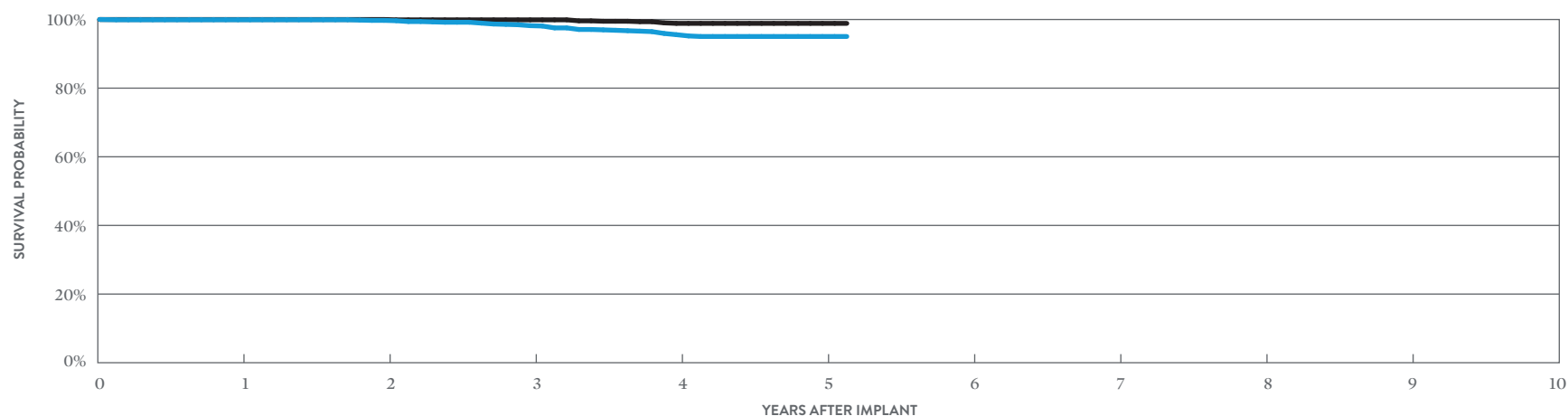
CUSTOMER REPORTED PERFORMANCE DATA

Unify Assura™ CRT-D

MODEL CD3257-40Q* (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	2,716
Estimated Active US Implants	1,408
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	29
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	1	0.04%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	1	0.04%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	3	0.11%	3	0.11%
Other	0	0.00%	0	0.00%
Total	4	0.15%	4	0.15%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 62 MONTHS
SURVIVAL PROBABILITY	99.92%	99.74%	98.20%	95.58%	95.05%	95.05%
± 1 STANDARD ERROR	0.05%	0.11%	0.27%	0.47%	0.53%	0.53%
SAMPLE SIZE	2,530	2,210	1,950	1,570	790	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 62 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	99.90%	98.87%	98.87%	98.87%
± 1 STANDARD ERROR	0.00%	0.00%	0.07%	0.25%	0.27%	0.27%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

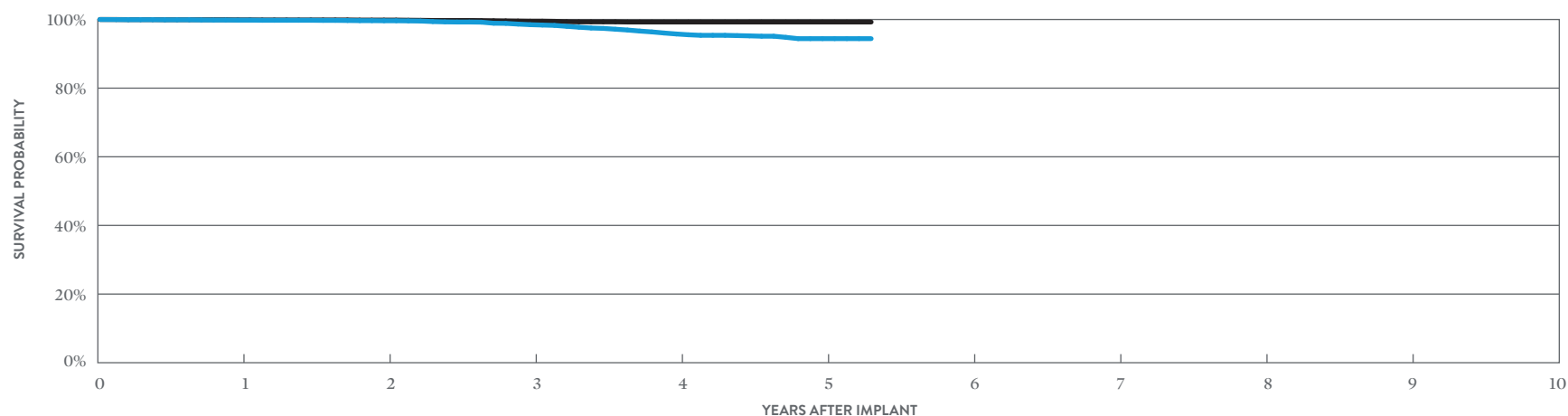
CUSTOMER REPORTED PERFORMANCE DATA

Unify Assura™ CRT-D

MODEL CD3257-40 (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	6,744
Estimated Active US Implants	3,580
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	74
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	4	0.06%	3	0.04%
Electrical Interconnect	1	0.01%	0	0.00%
Battery	0	0.00%	1	0.01%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	4	0.06%	3	0.04%
Other	1	0.01%	1	0.01%
Total	10	0.15%	8	0.12%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 64 MONTHS
SURVIVAL PROBABILITY	99.81%	99.63%	98.49%	95.78%	94.42%	94.42%
± 1 STANDARD ERROR	0.05%	0.08%	0.16%	0.29%	0.40%	0.40%
SAMPLE SIZE	6,320	5,590	4,970	4,000	2,080	300

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 64 MONTHS
SURVIVAL PROBABILITY	99.90%	99.83%	99.51%	99.26%	99.26%	99.26%
± 1 STANDARD ERROR	0.03%	0.05%	0.09%	0.12%	0.12%	0.12%

Cardiac Resynchronization Therapy (CRT) ICDs

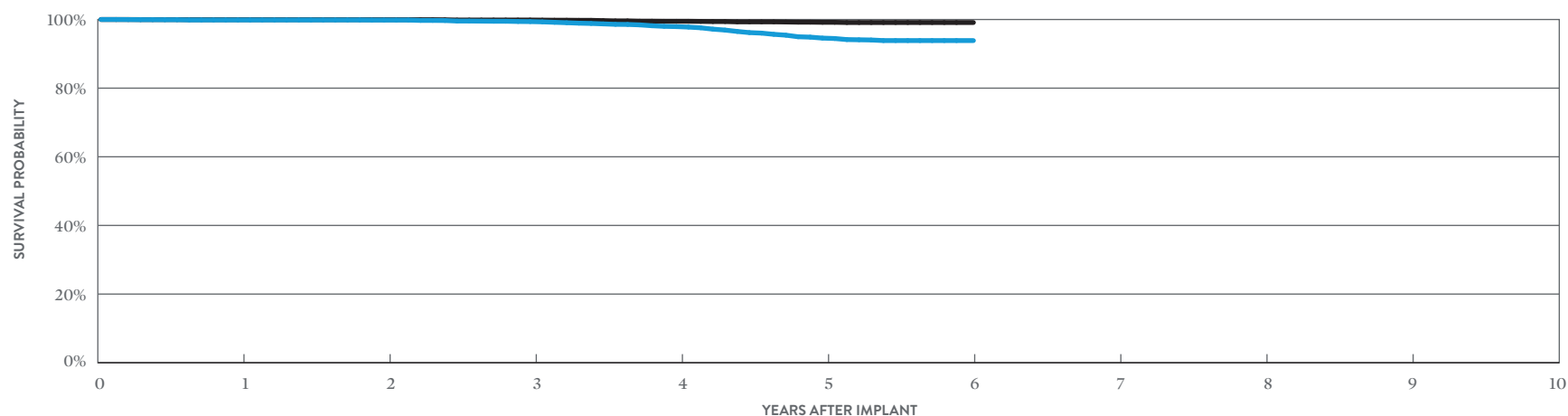
CUSTOMER REPORTED PERFORMANCE DATA

Unify Quadra™ CRT-D

MODEL CD3249-40Q* (ADVISORY POPULATION)

US Regulatory Approval	November 2011
Registered US Implants	8,948
Estimated Active US Implants	4,610
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	112
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.03%	3	0.03%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	0.01%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.01%
Possible Early Battery Depletion	9	0.10%	4	0.04%
Other	2	0.02%	0	0.00%
Total	15	0.17%	8	0.09%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6
SURVIVAL PROBABILITY	99.87%	99.84%	99.39%	97.93%	94.58%	93.86%
± 1 STANDARD ERROR	0.04%	0.04%	0.09%	0.17%	0.29%	0.33%
SAMPLE SIZE	8,410	7,500	6,790	6,090	4,860	390

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6
SURVIVAL PROBABILITY	99.95%	99.95%	99.85%	99.48%	99.18%	99.12%
± 1 STANDARD ERROR	0.02%	0.02%	0.05%	0.09%	0.11%	0.12%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

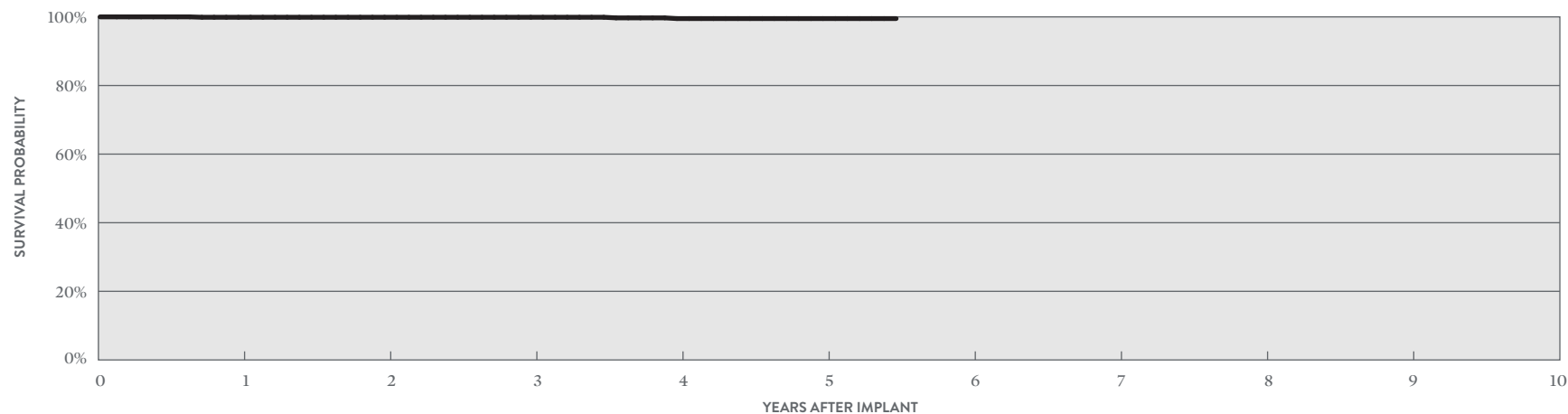
Unify Quadra™ CRT-D

MODEL CD3249-40Q*

US Regulatory Approval	November 2011
Number of Devices Enrolled in Study	990
Active Devices Enrolled in Study	431
Cumulative Months of Follow-up	39,827
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Premature Battery Depletion	2	0.20%
Skin Erosion	1	0.10%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	1	0.10%	1	0.10%
Other	0	0.00%	0	0.00%
Total	1	0.10%	1	0.10%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	AT 66 MONTHS
SURVIVAL PROBABILITY	99.89%	99.89%	99.89%	99.51%	99.51%	99.51%
± 1 STANDARD ERROR	0.11%	0.11%	0.11%	0.21%	0.29%	0.29%
SAMPLE SIZE	920	780	660	560	350	50

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

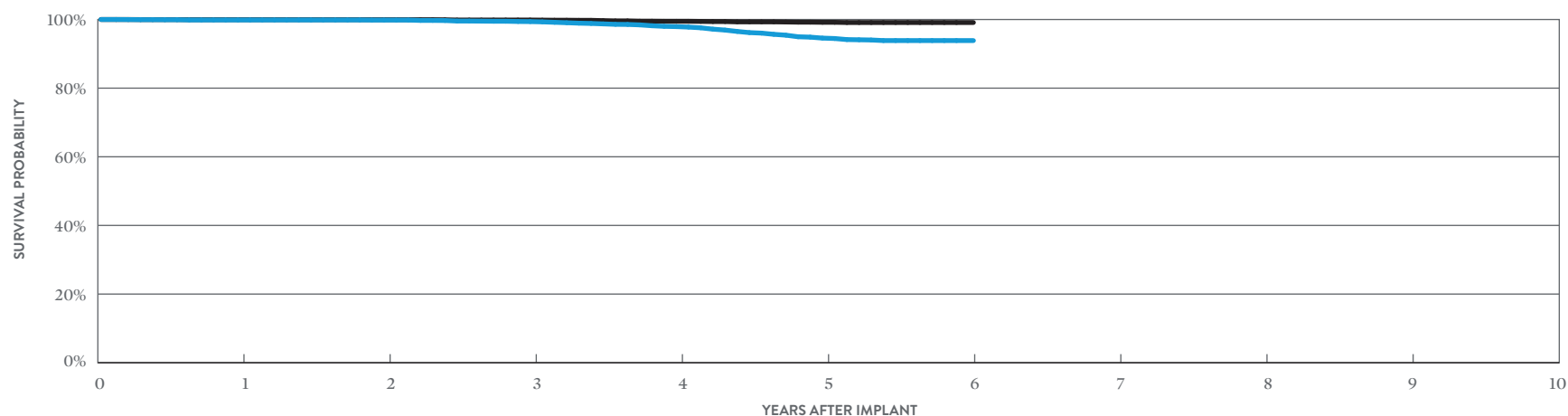
CUSTOMER REPORTED PERFORMANCE DATA

Unify Quadra™ CRT-D

MODEL CD3249-40 (ADVISORY POPULATION)

US Regulatory Approval	November 2011
Registered US Implants	2,524
Estimated Active US Implants	1,243
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	38
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.04%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	1	0.04%	0	0.00%
Total	1	0.04%	1	0.04%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 71 MONTHS
SURVIVAL PROBABILITY	99.92%	99.92%	99.60%	98.02%	94.65%	93.83%
± 1 STANDARD ERROR	0.06%	0.06%	0.12%	0.32%	0.58%	0.63%
SAMPLE SIZE	2,370	2,090	1,870	1,680	1,350	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 71 MONTHS
SURVIVAL PROBABILITY	99.92%	99.92%	99.92%	99.81%	99.81%	99.81%
± 1 STANDARD ERROR	0.06%	0.06%	0.06%	0.10%	0.10%	0.10%

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

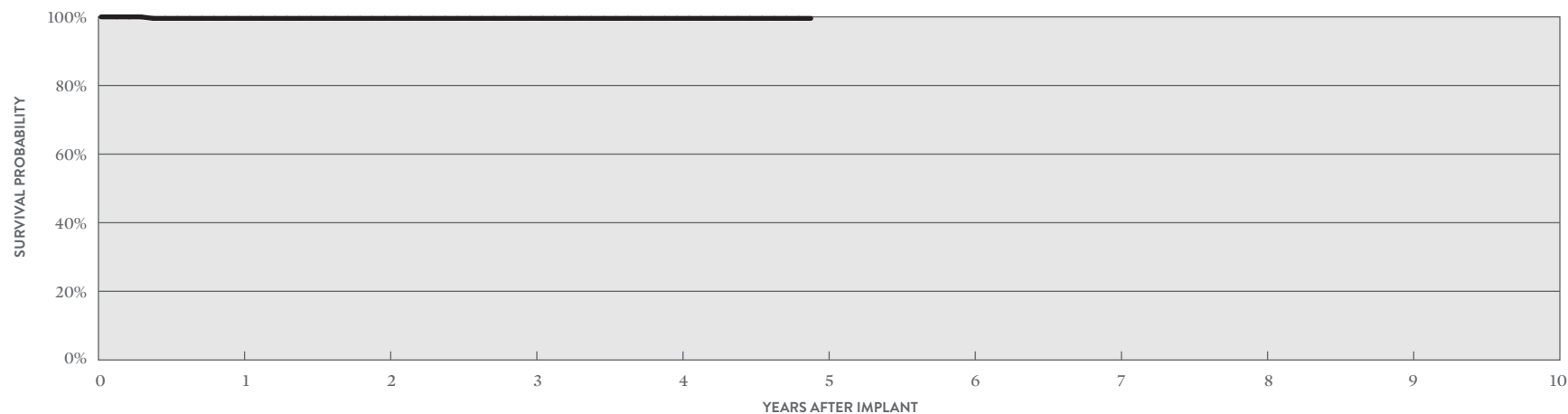
Unify Quadra™ CRT-D

MODEL CD3249-40

US Regulatory Approval	November 2011
Number of Devices Enrolled in Study	244
Active Devices Enrolled in Study	78
Cumulative Months of Follow-up	9,335
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Skin Erosion	1	0.41%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	AT 59 MONTHS
SURVIVAL PROBABILITY	99.57%	99.57%	99.57%	99.57%	99.57%
± 1 STANDARD ERROR	0.43%	0.43%	0.43%	0.43%	0.43%
SAMPLE SIZE	230	190	160	130	50

Cardiac Resynchronization Therapy (CRT) ICDs

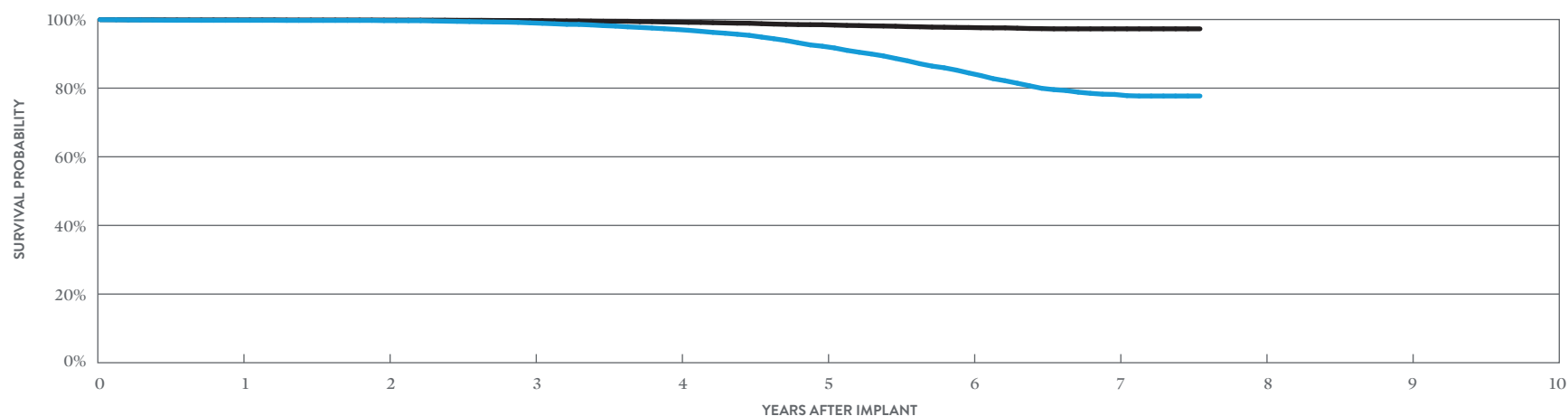
CUSTOMER REPORTED PERFORMANCE DATA

Unify™ CRT-D

MODEL CD3231-40Q (ADVISORY POPULATION)

US Regulatory Approval	May 2010
Registered US Implants	19,029
Estimated Active US Implants	7,306
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	727
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.01%	4	0.02%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	12	0.06%	6	0.03%
High Voltage Capacitor	14	0.07%	2	0.01%
Software/Firmware	0	0.00%	2	0.01%
Mechanical	1	<0.01%	2	0.01%
Possible Early Battery Depletion	49	0.26%	14	0.07%
Other	6	0.03%	3	0.02%
Total	85	0.45%	33	0.17%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 91 MONTHS
SURVIVAL PROBABILITY	99.76%	99.67%	99.02%	97.10%	92.21%	84.43%	78.16%	77.71%
± 1 STANDARD ERROR	0.04%	0.04%	0.07%	0.14%	0.24%	0.34%	0.45%	0.48%
SAMPLE SIZE	17,770	15,690	14,100	12,620	10,990	8,590	4,520	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 91 MONTHS
SURVIVAL PROBABILITY	99.88%	99.83%	99.68%	99.24%	98.48%	97.66%	97.28%	97.28%
± 1 STANDARD ERROR	0.03%	0.03%	0.04%	0.07%	0.11%	0.14%	0.17%	0.17%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

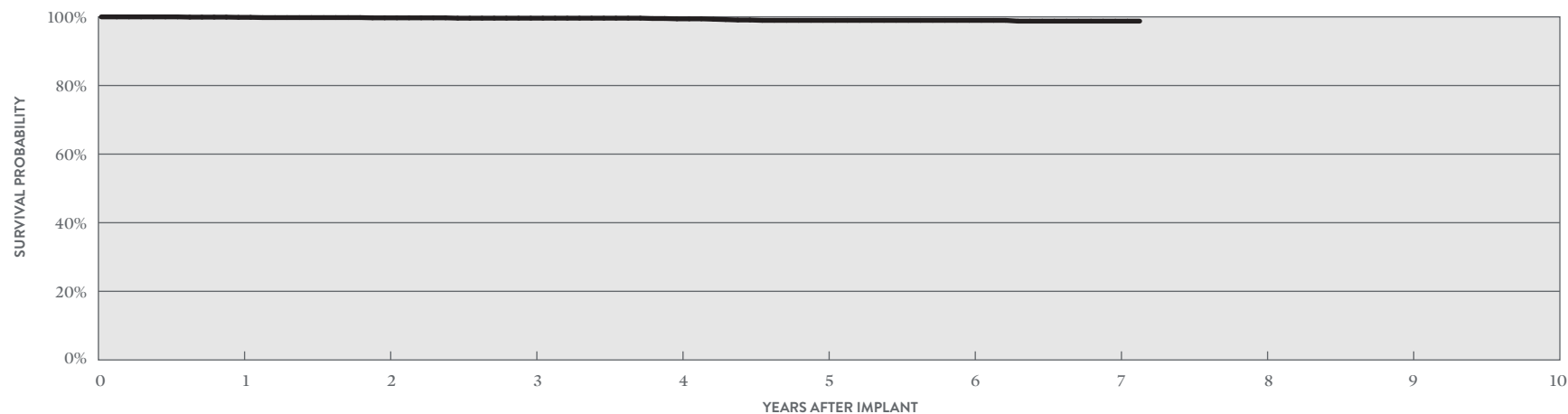
Unify™ CRT-D

MODEL CD3231-40Q

US Regulatory Approval	May 2010
Number of Devices Enrolled in Study	1,679
Active Devices Enrolled in Study	725
Cumulative Months of Follow-up	83,509
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Inappropriate Shock	2	0.12%
Premature Battery Depletion	9	0.54%
Skin Erosion	1	0.06%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.06%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	0.06%	1	0.06%
High Voltage Capacitor	1	0.06%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.06%
Possible Early Battery Depletion	11	0.66%	1	0.06%
Other	2	0.12%	0	0.00%
Total	15	0.89%	4	0.24%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	AT 86 MONTHS
SURVIVAL PROBABILITY	99.87%	99.72%	99.63%	99.43%	98.98%	98.98%	98.76%	98.76%
± 1 STANDARD ERROR	0.07%	0.14%	0.16%	0.19%	0.31%	0.31%	0.38%	0.38%
SAMPLE SIZE	1,570	1,360	1,180	1,020	860	680	340	60

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

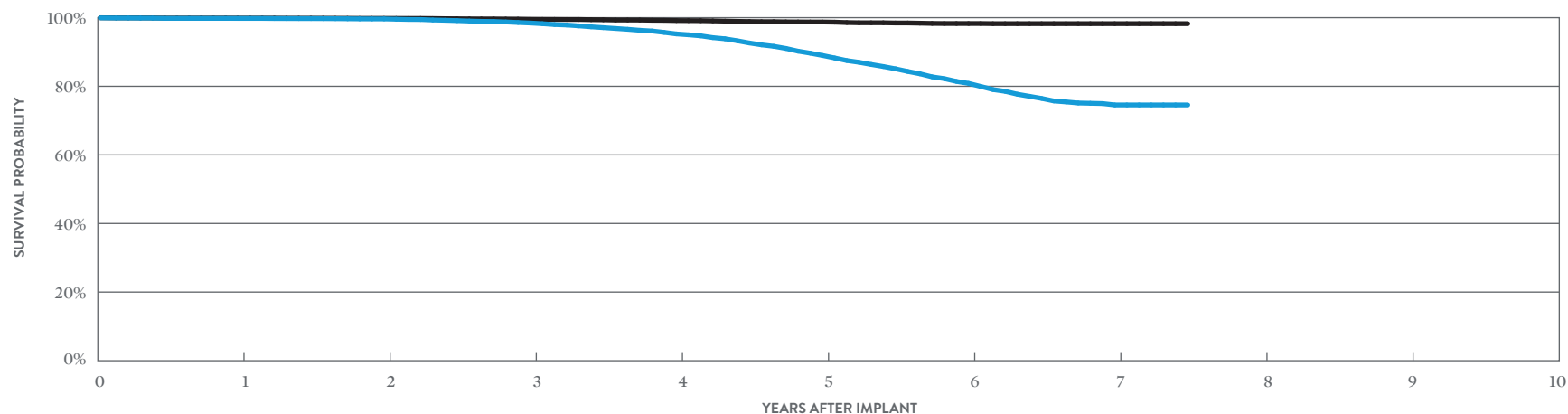
CUSTOMER REPORTED PERFORMANCE DATA

Unify™ CRT-D

MODEL CD3231-40 (ADVISORY POPULATION)

US Regulatory Approval	May 2010
Registered US Implants	20,501
Estimated Active US Implants	7,797
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	909
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	10	0.05%	4	0.02%
Electrical Interconnect	3	0.01%	0	0.00%
Battery	9	0.04%	3	0.01%
High Voltage Capacitor	5	0.02%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	1	<0.01%	0	0.00%
Possible Early Battery Depletion	25	0.12%	7	0.03%
Other	10	0.05%	11	0.05%
Total	63	0.31%	25	0.12%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 90 MONTHS
SURVIVAL PROBABILITY	99.79%	99.64%	98.45%	95.27%	88.99%	80.84%	74.57%	74.57%
± 1 STANDARD ERROR	0.03%	0.04%	0.09%	0.17%	0.27%	0.37%	0.49%	0.51%
SAMPLE SIZE	19,140	16,760	14,860	13,090	11,030	7,750	3,520	310

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 90 MONTHS
SURVIVAL PROBABILITY	99.88%	99.80%	99.53%	99.17%	98.75%	98.30%	98.26%	98.26%
± 1 STANDARD ERROR	0.02%	0.03%	0.05%	0.07%	0.10%	0.12%	0.12%	0.12%

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

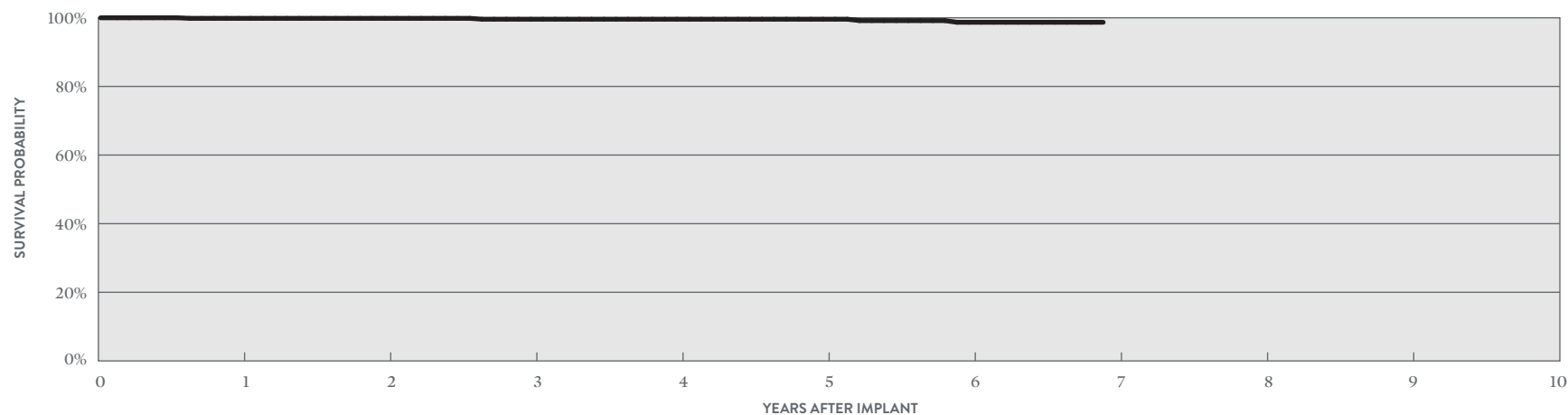
Unify™ CRT-D

MODEL CD3231-40

US Regulatory Approval	May 2010
Number of Devices Enrolled in Study	687
Active Devices Enrolled in Study	208
Cumulative Months of Follow-up	29,989
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Premature Battery Depletion	3	0.44%
Skin Erosion	1	0.15%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	0.15%	0	0.00%
Electrical Interconnect	1	0.15%	0	0.00%
Battery	1	0.15%	2	0.29%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	0.15%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	1	0.15%	0	0.00%
Other	0	0.00%	0	0.00%
Total	4	0.58%	3	0.44%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	AT 83 MONTHS
SURVIVAL PROBABILITY	99.84%	99.84%	99.59%	99.59%	99.59%	98.69%	98.69%
± 1 STANDARD ERROR	0.16%	0.16%	0.29%	0.29%	0.29%	0.70%	0.70%
SAMPLE SIZE	630	510	410	350	280	220	60

Cardiac Resynchronization Therapy (CRT) ICDs

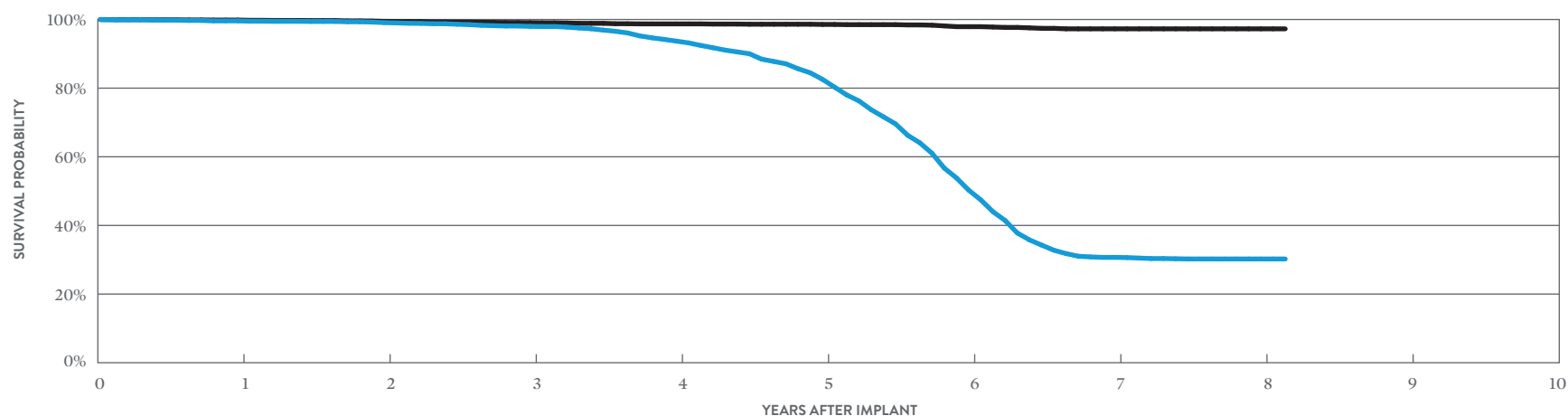
CUSTOMER REPORTED PERFORMANCE DATA

Promote™ + CRT-D

MODEL CD3211-36Q*

US Regulatory Approval	February 2009
Registered US Implants	6,902
Estimated Active US Implants	1,167
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	1,080
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	4	0.06%	3	0.04%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	9	0.13%	5	0.07%
High Voltage Capacitor	1	0.01%	0	0.00%
Software/Firmware	0	0.00%	9	0.13%
Mechanical	1	0.01%	0	0.00%
Possible Early Battery Depletion	2	0.03%	0	0.00%
Other	5	0.07%	6	0.09%
Total	22	0.32%	23	0.33%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 98 MONTHS
SURVIVAL PROBABILITY	99.59%	99.09%	97.99%	93.70%	82.56%	50.30%	30.69%	30.23%	30.23%
± 1 STANDARD ERROR	0.08%	0.11%	0.19%	0.34%	0.55%	0.79%	0.73%	0.73%	0.73%
SAMPLE SIZE	6,380	5,530	4,940	4,370	3,680	2,650	1,560	710	230

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 98 MONTHS
SURVIVAL PROBABILITY	99.84%	99.46%	99.08%	98.72%	98.56%	97.91%	97.28%	97.28%	97.28%
± 1 STANDARD ERROR	0.05%	0.09%	0.13%	0.16%	0.17%	0.24%	0.31%	0.31%	0.31%

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

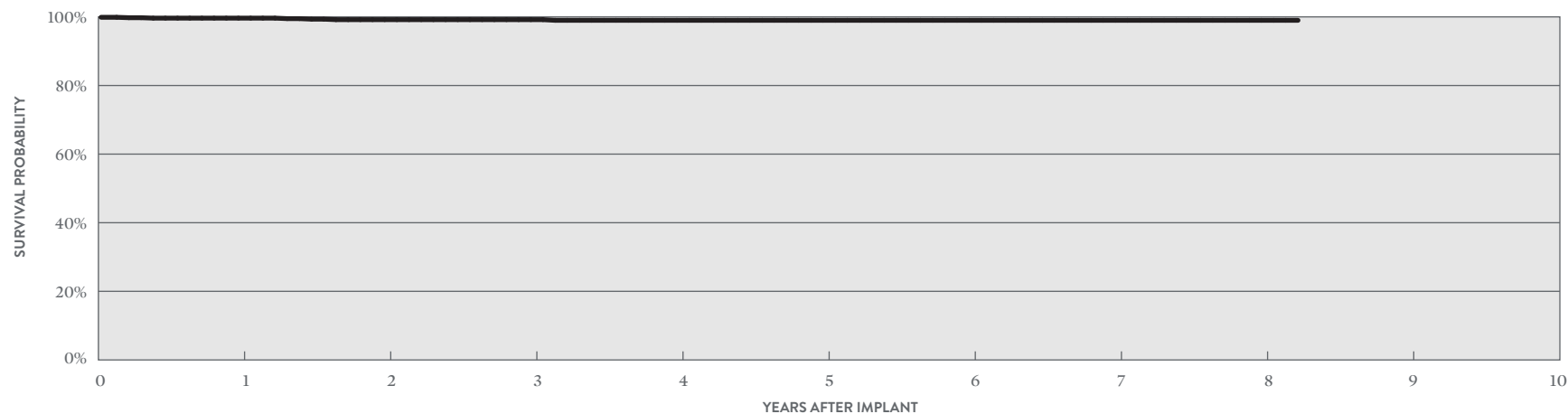
Promote™ + CRT-D

MODEL CD3211-36Q*

US Regulatory Approval	February 2009
Number of Devices Enrolled in Study	856
Active Devices Enrolled in Study	240
Cumulative Months of Follow-up	43,584
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	36 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Inappropriate Shock	3	0.35%
Premature Battery Depletion	2	0.23%
Skin Erosion	2	0.23%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	0.12%	1	0.12%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	0.12%	1	0.12%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.12%
Possible Early Battery Depletion	2	0.23%	0	0.00%
Other	0	0.00%	0	0.00%
Total	4	0.47%	3	0.35%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	8	AT 99 MONTHS
SURVIVAL PROBABILITY	99.64%	99.19%	99.19%	99.00%	99.00%	99.00%	99.00%	99.00%	99.00%
± 1 STANDARD ERROR	0.21%	0.33%	0.33%	0.38%	0.38%	0.38%	0.38%	0.38%	0.38%
SAMPLE SIZE	790	680	580	480	380	300	260	180	70

*DF4-LLHH connector type.

Cardiac Resynchronization Therapy (CRT) ICDs

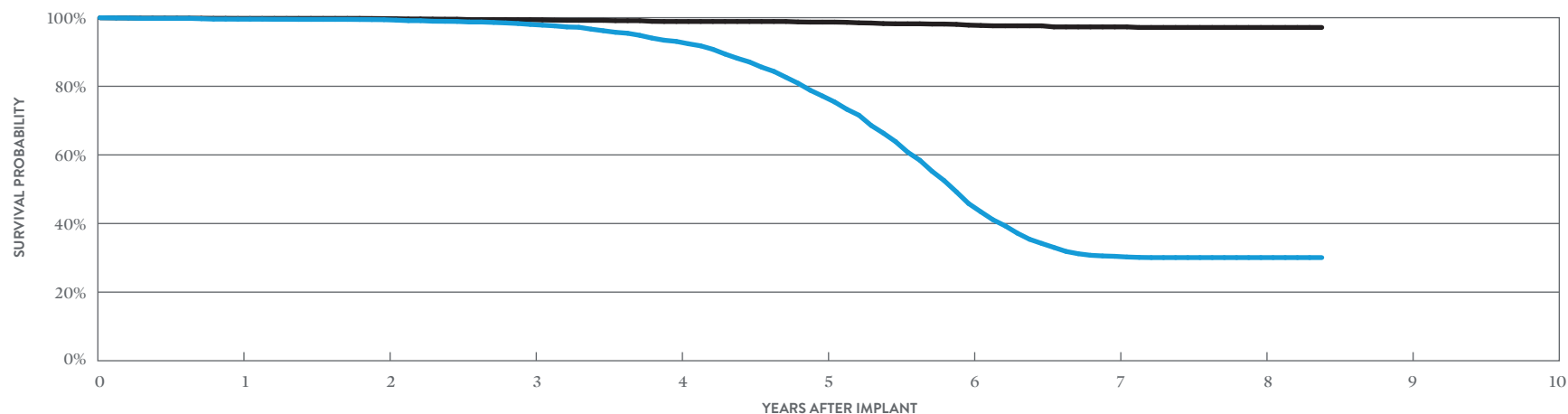
CUSTOMER REPORTED PERFORMANCE DATA

Promote™ + CRT-D

MODEL CD3211-36

US Regulatory Approval	February 2009
Registered US Implants	8,644
Estimated Active US Implants	1,414
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	1,261
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.03%	3	0.03%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	11	0.13%	3	0.03%
High Voltage Capacitor	2	0.02%	0	0.00%
Software/Firmware	1	0.01%	11	0.13%
Mechanical	0	0.00%	1	0.01%
Possible Early Battery Depletion	5	0.06%	1	0.01%
Other	5	0.06%	3	0.03%
Total	27	0.31%	22	0.25%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	99.56%	99.41%	98.04%	93.08%	77.16%	45.82%	30.44%	30.07%	30.07%
± 1 STANDARD ERROR	0.07%	0.09%	0.16%	0.33%	0.58%	0.74%	0.68%	0.68%	0.68%
SAMPLE SIZE	7,970	6,860	6,040	5,200	4,170	2,850	1,690	920	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	99.79%	99.73%	99.39%	98.89%	98.71%	97.83%	97.31%	97.15%	97.15%
± 1 STANDARD ERROR	0.05%	0.06%	0.10%	0.14%	0.15%	0.21%	0.29%	0.31%	0.31%

Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

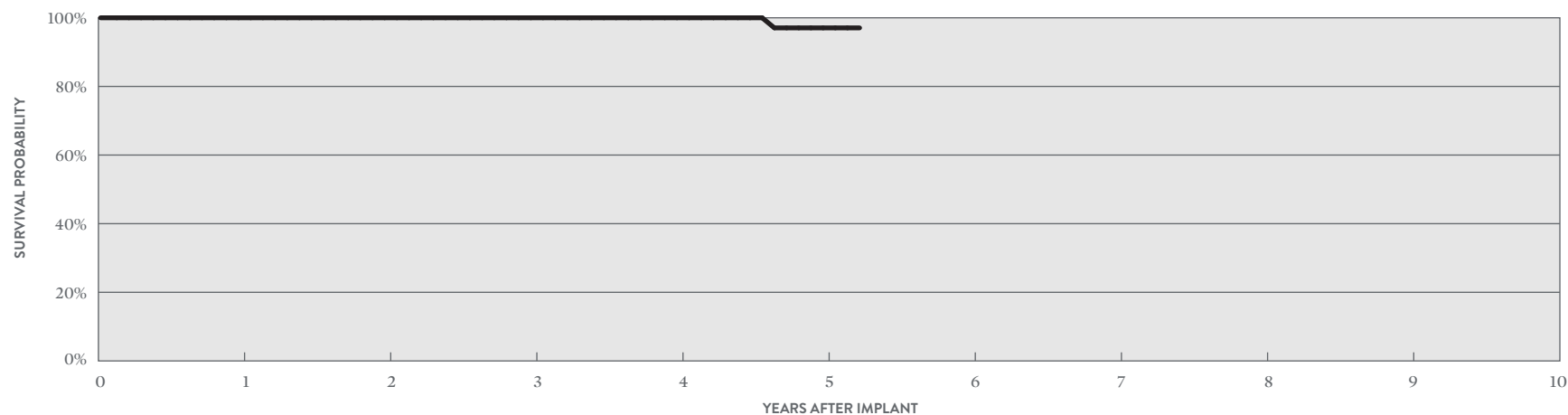
Promote™ + CRT-D

MODEL CD3211-36

US Regulatory Approval	February 2009
Number of Devices Enrolled in Study	223
Active Devices Enrolled in Study	22
Cumulative Months of Follow-up	9,268
Estimated Longevity	(see table on page 51)
Max. Delivered Energy	36 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Skin Erosion	2	0.90%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	2	0.90%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	2	0.90%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	AT 63 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%	97.06%	97.06%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%	2.05%	2.05%
SAMPLE SIZE	210	170	130	100	70	50

Cardiac Resynchronization Therapy (CRT) ICDs

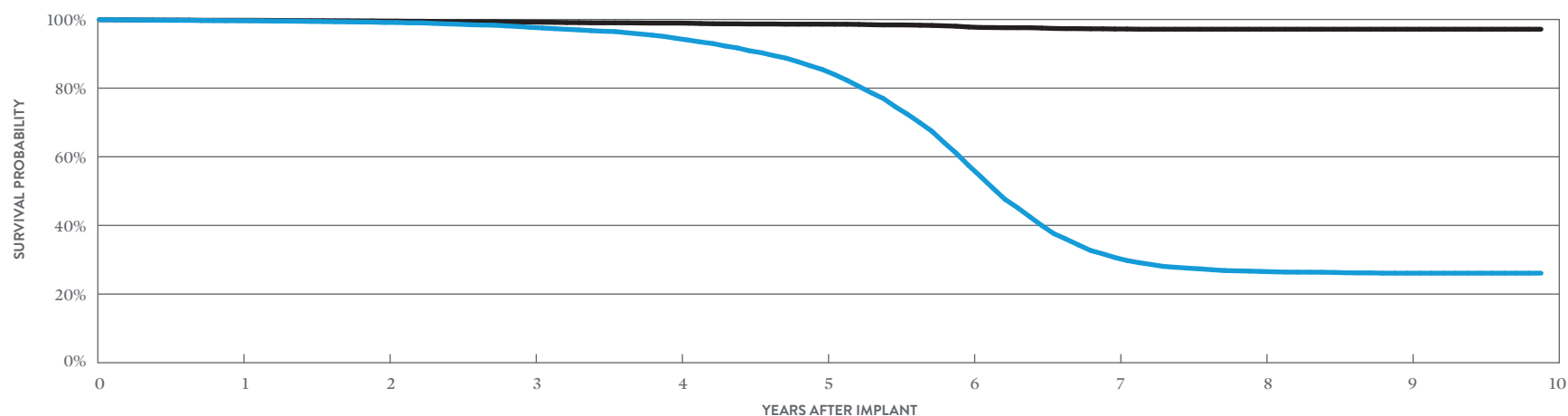
CUSTOMER REPORTED PERFORMANCE DATA

Promote™ RF CRT-D

MODEL 3207-36

US Regulatory Approval	September 2007
Registered US Implants	24,004
Estimated Active US Implants	2,794
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	3,234
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	4	0.02%	6	0.02%
Electrical Interconnect	5	0.02%	3	0.01%
Battery	18	0.07%	9	0.04%
High Voltage Capacitor	5	0.02%	1	<0.01%
Software/Firmware	0	0.00%	15	0.06%
Mechanical	3	0.01%	10	0.04%
Possible Early Battery Depletion	10	0.04%	6	0.02%
Other	17	0.07%	17	0.07%
Total	62	0.26%	67	0.28%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.65%	99.14%	97.72%	94.50%	85.42%	57.46%	30.66%	26.59%	26.09%	26.09%
± 1 STANDARD ERROR	0.04%	0.06%	0.11%	0.17%	0.30%	0.46%	0.44%	0.42%	0.42%	0.42%
SAMPLE SIZE	22,100	18,790	16,200	13,880	11,360	8,110	4,810	3,000	1,910	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.77%	99.53%	99.23%	98.95%	98.64%	97.81%	97.23%	97.17%	97.17%	97.17%
± 1 STANDARD ERROR	0.03%	0.04%	0.06%	0.08%	0.09%	0.13%	0.17%	0.18%	0.18%	0.18%

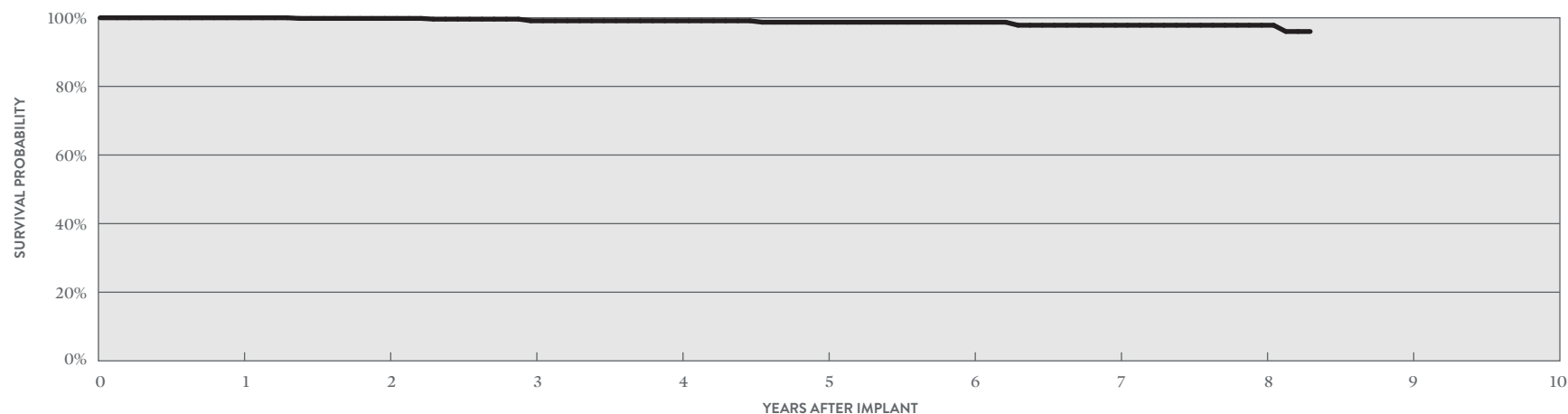
Cardiac Resynchronization Therapy (CRT) ICDs

ACTIVELY MONITORED STUDY DATA

Promote™ RF CRT-D

MODEL 3207-36

		QUALIFYING COMPLICATIONS		MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY			
			QTY	RATE	QTY	RATE	QTY	RATE	
US Regulatory Approval	September 2007	Inappropriate Shock	1	0.15%	Electrical Component	0	0.00%	1	0.15%
Number of Devices Enrolled in Study	673	Premature Battery Depletion	3	0.45%	Electrical Interconnect	0	0.00%	0	0.00%
Active Devices Enrolled in Study	59	Skin Erosion	3	0.45%	Battery	0	0.00%	1	0.15%
Cumulative Months of Follow-up	30,734				High Voltage Capacitor	0	0.00%	0	0.00%
Estimated Longevity	(see table on page 51)				Software/Firmware	0	0.00%	1	0.15%
Max. Delivered Energy	36 joules				Mechanical	0	0.00%	0	0.00%
					Possible Early Battery Depletion	0	0.00%	1	0.15%
					Other	2	0.30%	1	0.15%
					Total	2	0.30%	5	0.74%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	8	AT 100 MONTHS
SURVIVAL PROBABILITY	100.00%	99.82%	99.12%	99.12%	98.71%	98.71%	97.83%	97.83%	96.00%
± 1 STANDARD ERROR	0.00%	0.18%	0.28%	0.44%	0.60%	0.60%	1.06%	1.06%	2.09%
SAMPLE SIZE	630	550	450	340	240	170	100	60	50

Cardiac Resynchronization Therapy (CRT) ICDs

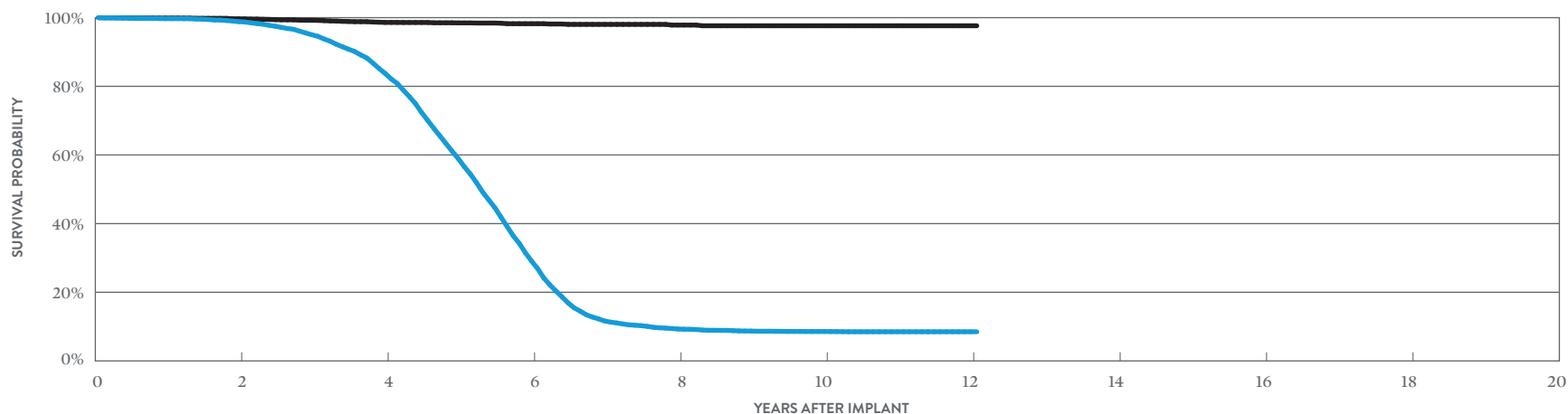
CUSTOMER REPORTED PERFORMANCE DATA

Atlas™ + HF CRT-D

MODEL V-343

US Regulatory Approval	November 2004
Registered US Implants	18,777
Estimated Active US Implants	821
Estimated Longevity	(see table on page 51)
Normal Battery Depletion	3,441
Max. Delivered Energy	36 joules
Number of US Advisories (see pgs. 330, 331)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.02%	1	<0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	40	0.21%	4	0.02%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	7	0.04%	11	0.06%
Other	10	0.05%	4	0.02%
Total	60	0.32%	22	0.12%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 145 MONTHS
SURVIVAL PROBABILITY	98.84%	83.76%	29.13%	9.29%	8.57%	8.50%	8.50%
± 1 STANDARD ERROR	0.08%	0.33%	0.49%	0.30%	0.28%	0.28%	0.28%
SAMPLE SIZE	14,880	9,930	3,970	1,100	820	400	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 145 MONTHS
SURVIVAL PROBABILITY	99.66%	98.61%	98.24%	97.84%	97.64%	97.64%	97.64%
± 1 STANDARD ERROR	0.05%	0.10%	0.14%	0.22%	0.27%	0.27%	0.27%

BATTERY LONGEVITY SUMMARY
**Cardiac Resynchronization
Therapy (CRT) ICDs**

Cardiac Resynchronization Therapy (CRT) ICDs

Battery Longevity (years)

MODELS	FAMILY	NO PACING	25% PACING	50% PACING	100% PACING
CD3369-40Q	Quadra Assura MP [™] CRT-D*	11.1	9.9	8.9	7.4
CD3369-40C	Quadra Assura MP [™] CRT-D*	11.1	9.9	8.9	7.4
CD3365-40Q	Quadra Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3365-40C	Quadra Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3357-40Q	Unify Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3357-40C	Unify Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3265-40Q	Quadra Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3265-40	Quadra Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3257-40Q	Unify Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3257-40	Unify Assura [™] CRT-D*	11.1	9.9	8.9	7.4
CD3249-40Q	Unify Quadra [™] CRT-D*	10.2	9.0	8.1	6.7
CD3249-40	Unify Quadra [™] CRT-D*	10.2	9.0	8.1	6.7
CD3231-40Q	Unify [™] CRT-D*	10.1	9.0	8.1	6.7
CD3231-40	Unify [™] CRT-D*	10.1	9.0	8.1	6.7
CD3211-36Q	Promote [™] + CRT-D**	8.2	7.2	6.5	5.4
CD3211-36	Promote [™] + CRT-D**	8.2	7.2	6.5	5.4
3207-36	Promote [™] RF CRT-D**	8.2	7.2	6.5	5.4
V-343	Atlas [™] + HF CRT-D**	7.9	7.1	6.4	5.4

Pacing parameters: DDD-BiV, RV 2.5V, LV 2.5V, A 2.5V, 0.5 ms, 60 ppm, 500 ohms

*Battery voltage range: 3.20-2.59. Three maximum charges per year.

**Battery voltage range: 3.20-2.45. Four maximum charges per year as well as monthly charging during the battery's mid-life voltage range.

SUMMARY INFORMATION

Cardiac Resynchronization
Therapy (CRT) ICDs

Cardiac Resynchronization Therapy (CRT) ICDs

Survival Probability Summary

INCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
CD3369-40Q	Quadra Assura MP [™] CRT-D	99.86%									
CD3369-40C	Quadra Assura MP [™] CRT-D	100.00%									
CD3365-40Q	Quadra Assura [™] CRT-D	99.87%	99.82%								
CD3365-40Q	Quadra Assura [™] CRT-D†	99.78%	99.47%	99.00%	98.87%						
CD3365-40C	Quadra Assura [™] CRT-D	100.00%	100.00%								
CD3365-40C	Quadra Assura [™] CRT-D†	99.69%	99.22%	98.94%	98.94%						
CD3357-40Q	Unify Assura [™] CRT-D	99.92%	99.92%								
CD3357-40Q	Unify Assura [™] CRT-D†	99.78%	99.40%	98.63%	98.42%						
CD3357-40C	Unify Assura [™] CRT-D	99.98%	99.98%								
CD3357-40C	Unify Assura [™] CRT-D†	99.83%	99.53%	98.69%	98.61%						
CD3265-40Q	Quadra Assura [™] CRT-D†	99.83%	99.74%	99.38%	98.36%	97.90%					
CD3265-40	Quadra Assura [™] CRT-D†	99.89%	99.70%	99.64%	99.01%	97.43%					
CD3257-40Q	Unify Assura [™] CRT-D†	99.92%	99.74%	98.20%	95.58%	95.05%					
CD3257-40	Unify Assura [™] CRT-D†	99.81%	99.63%	98.49%	95.78%	94.42%					
CD3249-40Q	Unify Quadra [™] CRT-D†	99.87%	99.84%	99.39%	97.93%	94.58%	93.86%				
CD3249-40	Unify Quadra [™] CRT-D†	99.92%	99.92%	99.60%	98.02%	94.65%					
CD3231-40Q	Unify [™] CRT-D†	99.76%	99.67%	99.02%	97.10%	92.21%	84.43%	78.16%			
CD3231-40	Unify [™] CRT-D†	99.79%	99.64%	98.45%	95.27%	88.99%	80.84%	74.57%			
CD3211-36Q	Promote [™] + CRT-D	99.59%	99.09%	97.99%	93.70%	82.56%	50.30%	30.69%	30.23%		
CD3211-36	Promote [™] + CRT-D	99.56%	99.41%	98.04%	93.08%	77.16%	45.82%	30.44%	30.07%		
3207-36	Promote [™] RF CRT-D	99.65%	99.14%	97.72%	94.50%	85.42%	57.46%	30.66%	26.59%	26.09%	
V-343	Atlas [™] + HF CRT-D	99.71%	98.84%	95.01%	83.76%	58.92%	29.13%	11.62%	9.29%	8.70%	8.57%

†Premature battery depletion advisory population.

Cardiac Resynchronization Therapy (CRT) ICDs

Survival Probability Summary

EXCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
CD3369-40Q	Quadra Assura MP [™] CRT-D	99.86%									
CD3369-40C	Quadra Assura MP [™] CRT-D	100.00%									
CD3365-40Q	Quadra Assura [™] CRT-D	99.87%	99.82%								
CD3365-40Q	Quadra Assura [™] CRT-D†	99.83%	99.59%	99.19%	99.14%						
CD3365-40C	Quadra Assura [™] CRT-D	100.00%	100.00%								
CD3365-40C	Quadra Assura [™] CRT-D†	99.79%	99.32%	99.03%	99.03%						
CD3357-40Q	Unify Assura [™] CRT-D	99.92%	99.92%								
CD3357-40Q	Unify Assura [™] CRT-D†	99.90%	99.52%	99.16%	99.16%						
CD3357-40C	Unify Assura [™] CRT-D	99.98%	99.98%								
CD3357-40C	Unify Assura [™] CRT-D†	99.89%	99.64%	99.38%	99.38%						
CD3265-40Q	Quadra Assura [™] CRT-D†	99.87%	99.85%	99.64%	99.27%	99.23%					
CD3265-40	Quadra Assura [™] CRT-D†	99.89%	99.76%	99.70%	99.37%	99.37%					
CD3257-40Q	Unify Assura [™] CRT-D†	100.00%	100.00%	99.90%	98.87%	98.87%					
CD3257-40	Unify Assura [™] CRT-D†	99.90%	99.83%	99.51%	99.26%	99.26%					
CD3249-40Q	Unify Quadra [™] CRT-D†	99.95%	99.95%	99.85%	99.48%	99.18%	99.12%				
CD3249-40	Unify Quadra [™] CRT-D†	99.92%	99.92%	99.92%	99.81%	99.81%					
CD3231-40Q	Unify [™] CRT-D†	99.88%	99.83%	99.68%	99.24%	98.48%	97.66%	97.28%			
CD3231-40	Unify [™] CRT-D†	99.88%	99.80%	99.53%	99.17%	98.75%	98.30%	98.26%			
CD3211-36Q	Promote [™] + CRT-D	99.84%	99.46%	99.08%	98.72%	98.56%	97.91%	97.28%	97.28%		
CD3211-36	Promote [™] + CRT-D	99.79%	99.73%	99.39%	98.89%	98.71%	97.83%	97.31%	97.15%		
3207-36	Promote [™] RF CRT-D	99.77%	99.53%	99.23%	98.95%	98.64%	97.81%	97.23%	97.17%	97.17%	
V-343	Atlas [™] + HF CRT-D	99.88%	99.66%	99.24%	98.61%	98.47%	98.24%	98.04%	97.84%	97.64%	97.64%

†Premature battery depletion advisory population.

Cardiac Resynchronization Therapy (CRT) ICDs

US Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD3369-40Q	Quadra Assura MP ⁺ CRT-D	22,164	0.70%	2	<0.01%	3	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	5	0.02%
CD3369-40C	Quadra Assura MP ⁺ CRT-D	3,217	0.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3365-40Q	Quadra Assura ⁻ CRT-D	14,525	1.70%	0	0.00%	3	0.02%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	1	<0.01%	6	0.04%
CD3365-40Q	Quadra Assura ⁻ CRT-D [†]	24,082	10.70%	4	0.02%	7	0.03%	2	<0.01%	0	0.00%	1	<0.01%	0	0.00%	12	0.05%	5	0.02%	31	0.13%
CD3365-40C	Quadra Assura ⁻ CRT-D	2,315	1.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3365-40C	Quadra Assura ⁻ CRT-D [†]	5,626	14.70%	5	0.09%	2	0.04%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	2	0.04%	2	0.04%	12	0.21%
CD3357-40Q	Unify Assura ⁻ CRT-D	8,366	1.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3357-40Q	Unify Assura ⁻ CRT-D [†]	5,343	13.40%	1	0.02%	2	0.04%	0	0.00%	2	0.04%	0	0.00%	0	0.00%	7	0.13%	0	0.00%	12	0.22%
CD3357-40C	Unify Assura ⁻ CRT-D	9,184	1.10%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.01%
CD3357-40C	Unify Assura ⁻ CRT-D [†]	9,599	13.10%	2	0.02%	2	0.02%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	5	0.05%	0	0.00%	10	0.10%
CD3265-40Q	Quadra Assura ⁻ CRT-D [†]	13,540	10.90%	2	0.01%	1	<0.01%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	12	0.09%	1	<0.01%	17	0.13%
CD3265-40	Quadra Assura ⁻ CRT-D [†]	3,926	13.10%	0	0.00%	1	0.03%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.05%	4	0.10%	7	0.18%
CD3257-40Q	Unify Assura ⁻ CRT-D [†]	2,716	15.80%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	3	0.11%	0	0.00%	4	0.15%
CD3257-40	Unify Assura ⁻ CRT-D [†]	6,744	13.10%	4	0.06%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	0.06%	1	0.01%	10	0.15%
CD3249-40Q	Unify Quadra ⁻ CRT-D [†]	8,948	11.70%	3	0.03%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	9	0.10%	2	0.02%	15	0.17%
CD3249-40	Unify Quadra ⁻ CRT-D [†]	2,524	13.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.04%	1	0.04%
CD3231-40Q	Unify ⁻ CRT-D [†]	19,029	14.80%	2	0.01%	1	<0.01%	12	0.06%	14	0.07%	0	0.00%	1	<0.01%	49	0.26%	6	0.03%	85	0.45%
CD3231-40	Unify ⁻ CRT-D [†]	20,501	16.50%	10	0.05%	3	0.01%	9	0.04%	5	0.02%	0	0.00%	1	<0.01%	25	0.12%	10	0.05%	63	0.31%
CD3211-36Q	Promote ⁻ + CRT-D	6,902	24.40%	4	0.06%	0	0.00%	9	0.13%	1	0.01%	0	0.00%	1	0.01%	2	0.03%	5	0.07%	22	0.32%
CD3211-36	Promote ⁻ + CRT-D	8,644	25.50%	3	0.03%	0	0.00%	11	0.13%	2	0.02%	1	0.01%	0	0.00%	5	0.06%	5	0.06%	27	0.31%
3207-36	Promote ⁻ RF CRT-D	24,004	26.30%	4	0.02%	5	0.02%	18	0.07%	5	0.02%	0	0.00%	3	0.01%	10	0.04%	17	0.07%	62	0.26%
V-343	Atlas ⁻ + HF CRT-D	18,777	24.90%	3	0.02%	0	0.00%	40	0.21%	0	0.00%	0	0.00%	0	0.00%	7	0.04%	10	0.05%	60	0.32%

Definitions of malfunction categories can be found on [pages 5-6](#).

†Premature battery depletion advisory population.

Cardiac Resynchronization Therapy (CRT) ICDs

US Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD3369-40Q	Quadra Assura MP ⁺ CRT-D	22,164	0.70%	1	<0.01%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	3	0.01%	0	0.00%	0	0.00%	5	0.02%
CD3369-40C	Quadra Assura MP ⁺ CRT-D	3,217	0.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3365-40Q	Quadra Assura ⁻ CRT-D	14,525	1.70%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	0.03%	0	0.00%	1	<0.01%	5	0.03%
CD3365-40Q	Quadra Assura ⁻ CRT-D [†]	24,082	10.70%	7	0.03%	0	0.00%	4	0.02%	0	0.00%	3	0.01%	2	<0.01%	14	0.06%	2	<0.01%	32	0.13%
CD3365-40C	Quadra Assura ⁻ CRT-D	2,315	1.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3365-40C	Quadra Assura ⁻ CRT-D [†]	5,626	14.70%	2	0.04%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	2	0.04%	1	0.02%	6	0.11%
CD3357-40Q	Unify Assura ⁻ CRT-D	8,366	1.10%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.01%	0	0.00%	1	0.01%	3	0.04%
CD3357-40Q	Unify Assura ⁻ CRT-D [†]	5,343	13.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.02%	1	0.02%	2	0.04%
CD3357-40C	Unify Assura ⁻ CRT-D	9,184	1.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3357-40C	Unify Assura ⁻ CRT-D [†]	9,599	13.10%	2	0.02%	0	0.00%	1	0.01%	0	0.00%	1	0.01%	1	0.01%	2	0.02%	3	0.03%	10	0.10%
CD3265-40Q	Quadra Assura ⁻ CRT-D [†]	13,540	10.90%	5	0.04%	0	0.00%	2	0.01%	0	0.00%	1	<0.01%	2	0.01%	5	0.04%	0	0.00%	15	0.11%
CD3265-40	Quadra Assura ⁻ CRT-D [†]	3,926	13.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.03%	0	0.00%	1	0.03%	2	0.05%
CD3257-40Q	Unify Assura ⁻ CRT-D [†]	2,716	15.80%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	0	0.00%	0	0.00%	3	0.11%	0	0.00%	4	0.15%
CD3257-40	Unify Assura ⁻ CRT-D [†]	6,744	13.10%	3	0.04%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	3	0.04%	1	0.01%	8	0.12%
CD3249-40Q	Unify Quadra ⁻ CRT-D [†]	8,948	11.70%	3	0.03%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.01%	4	0.04%	0	0.00%	8	0.09%
CD3249-40	Unify Quadra ⁻ CRT-D [†]	2,524	13.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	0	0.00%	1	0.04%
CD3231-40Q	Unify ⁻ CRT-D [†]	19,029	14.80%	4	0.02%	0	0.00%	6	0.03%	2	0.01%	2	0.01%	2	0.01%	14	0.07%	3	0.02%	33	0.17%
CD3231-40	Unify ⁻ CRT-D [†]	20,501	16.50%	4	0.02%	0	0.00%	3	0.01%	0	0.00%	0	0.00%	0	0.00%	7	0.03%	11	0.05%	25	0.12%
CD3211-36Q	Promote ⁻ + CRT-D	6,902	24.40%	3	0.04%	0	0.00%	5	0.07%	0	0.00%	9	0.13%	0	0.00%	0	0.00%	6	0.09%	23	0.33%
CD3211-36	Promote ⁻ + CRT-D	8,644	25.50%	3	0.03%	0	0.00%	3	0.03%	0	0.00%	11	0.13%	1	0.01%	1	0.01%	3	0.03%	22	0.25%
3207-36	Promote ⁻ RF CRT-D	24,004	26.30%	6	0.02%	3	0.01%	9	0.04%	1	<0.01%	15	0.06%	10	0.04%	6	0.02%	17	0.07%	67	0.28%
V-343	Atlas ⁻ + HF CRT-D	18,777	24.90%	1	<0.01%	0	0.00%	4	0.02%	0	0.00%	1	<0.01%	1	<0.01%	11	0.06%	4	0.02%	22	0.12%

Definitions of malfunction categories can be found on [pages 5-6](#).

†Premature battery depletion advisory population.

Cardiac Resynchronization Therapy (CRT) ICDs

Worldwide Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD3369-40Q	Quadra Assura MP [™] CRT-D	22,996	0.78%	2	<0.01%	3	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	5	0.02%
CD3369-40C	Quadra Assura MP [™] CRT-D	3,386	0.80%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3365-40Q	Quadra Assura [™] CRT-D	38,968	7.63%	4	0.01%	10	0.03%	2	<0.01%	0	0.00%	2	<0.01%	0	0.00%	13	0.03%	6	0.02%	37	0.09%
CD3365-40C	Quadra Assura [™] CRT-D	8,031	11.41%	5	0.06%	2	0.02%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	2	0.02%	2	0.02%	12	0.15%
CD3357-40Q	Unify Assura [™] CRT-D	14,174	6.29%	1	<0.01%	2	0.01%	0	0.00%	2	0.01%	0	0.00%	0	0.00%	7	0.05%	0	0.00%	12	0.08%
CD3357-40C	Unify Assura [™] CRT-D	19,286	7.65%	2	0.01%	3	0.02%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	5	0.03%	0	0.00%	11	0.06%
CD3265-40Q	Quadra Assura [™] CRT-D	13,958	11.48%	2	0.01%	2	0.01%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	12	0.09%	1	<0.01%	18	0.13%
CD3265-40	Quadra Assura [™] CRT-D	4,047	13.71%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.05%	4	0.10%	7	0.17%
CD3257-40Q	Unify Assura [™] CRT-D	2,728	16.68%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	3	0.11%	0	0.00%	4	0.15%
CD3257-40	Unify Assura [™] CRT-D	6,723	13.62%	4	0.06%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	0.06%	1	0.01%	10	0.15%
CD3249-40Q	Unify Quadra [™] CRT-D	10,696	11.39%	4	0.04%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	10	0.09%	2	0.02%	17	0.16%
CD3249-40	Unify Quadra [™] CRT-D	3,717	11.19%	1	0.03%	1	0.03%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.03%	3	0.08%
CD3231-40Q	Unify [™] CRT-D	20,963	15.39%	3	0.01%	1	<0.01%	13	0.06%	15	0.07%	0	0.00%	1	<0.01%	60	0.29%	8	0.04%	101	0.48%
CD3231-40	Unify [™] CRT-D	22,380	16.12%	11	0.05%	4	0.02%	10	0.04%	5	0.02%	0	0.00%	1	<0.01%	27	0.12%	10	0.04%	68	0.30%
CD3211-36Q	Promote [™] + CRT-D	15,991	13.04%	13	0.08%	0	0.00%	13	0.08%	7	0.04%	1	<0.01%	2	0.01%	6	0.04%	5	0.03%	47	0.29%
CD3211-36	Promote [™] + CRT-D	21,005	11.58%	13	0.06%	2	<0.01%	15	0.07%	4	0.02%	1	<0.01%	0	0.00%	8	0.04%	11	0.05%	54	0.26%
3207-36	Promote [™] RF CRT-D	25,838	26.22%	5	0.02%	5	0.02%	21	0.08%	5	0.02%	0	0.00%	3	0.01%	10	0.04%	20	0.08%	69	0.27%
V-343	Atlas [™] + HF CRT-D	19,292	24.73%	3	0.02%	0	0.00%	41	0.21%	0	0.00%	0	0.00%	0	0.00%	7	0.04%	10	0.05%	61	0.32%

Definitions of malfunction categories can be found on [pages 5-6](#).

Cardiac Resynchronization Therapy (CRT) ICDs

Worldwide Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD3369-40Q	Quadra Assura MP [™] CRT-D	22,996	0.78%	1	<0.01%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	3	0.01%	0	0.00%	0	0.00%	5	0.02%
CD3369-40C	Quadra Assura MP [™] CRT-D	3,386	0.80%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3365-40Q	Quadra Assura [™] CRT-D	38,968	7.63%	7	0.02%	0	0.00%	4	0.01%	0	0.00%	3	<0.01%	6	0.02%	14	0.04%	3	<0.01%	37	0.09%
CD3365-40C	Quadra Assura [™] CRT-D	8,031	11.41%	2	0.02%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	2	0.02%	1	0.01%	6	0.07%
CD3357-40Q	Unify Assura [™] CRT-D	14,174	6.29%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%	2	0.01%	5	0.04%
CD3357-40C	Unify Assura [™] CRT-D	19,286	7.65%	2	0.01%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	1	<0.01%	2	0.01%	3	0.02%	10	0.05%
CD3265-40Q	Quadra Assura [™] CRT-D	13,958	11.48%	5	0.04%	0	0.00%	2	0.01%	0	0.00%	1	<0.01%	2	0.01%	5	0.04%	0	0.00%	15	0.11%
CD3265-40	Quadra Assura [™] CRT-D	4,047	13.71%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	1	0.02%	2	0.05%
CD3257-40Q	Unify Assura [™] CRT-D	2,728	16.68%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	0	0.00%	0	0.00%	3	0.11%	0	0.00%	4	0.15%
CD3257-40	Unify Assura [™] CRT-D	6,723	13.62%	3	0.04%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	3	0.04%	1	0.01%	8	0.12%
CD3249-40Q	Unify Quadra [™] CRT-D	10,696	11.39%	3	0.03%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	5	0.05%	1	<0.01%	10	0.09%
CD3249-40	Unify Quadra [™] CRT-D	3,717	11.19%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.03%	0	0.00%	0	0.00%	1	0.03%
CD3231-40Q	Unify [™] CRT-D	20,963	15.39%	5	0.02%	0	0.00%	7	0.03%	2	<0.01%	2	<0.01%	3	0.01%	15	0.07%	3	0.01%	37	0.18%
CD3231-40	Unify [™] CRT-D	22,380	16.12%	5	0.02%	0	0.00%	5	0.02%	0	0.00%	1	<0.01%	0	0.00%	7	0.03%	11	0.05%	29	0.13%
CD3211-36Q	Promote [™] + CRT-D	15,991	13.04%	6	0.04%	0	0.00%	7	0.04%	0	0.00%	11	0.07%	2	0.01%	3	0.02%	9	0.06%	38	0.24%
CD3211-36	Promote [™] + CRT-D	21,005	11.58%	7	0.03%	0	0.00%	4	0.02%	0	0.00%	16	0.08%	2	<0.01%	2	<0.01%	6	0.03%	37	0.18%
3207-36	Promote [™] RF CRT-D	25,838	26.22%	7	0.03%	3	0.01%	10	0.04%	1	<0.01%	17	0.07%	10	0.04%	7	0.03%	18	0.07%	73	0.28%
V-343	Atlas [™] + HF CRT-D	19,292	24.73%	1	<0.01%	0	0.00%	4	0.02%	0	0.00%	1	<0.01%	1	<0.01%	11	0.06%	4	0.02%	22	0.11%

Definitions of malfunction categories can be found on [pages 5-6](#).

Cardiac Resynchronization Therapy (CRT) ICDs

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	INAPPROPRIATE SHOCK		LOSS OF TELEMETRY		PERICARDIAL EFFUSION		PREMATURE BATTERY DEPLETION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD3365-40Q	222	129	6,580	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.45%	1	0.45%
CD3357-40Q	219	167	4,837	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3357-40C	185	115	4,079	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3265-40Q	421	182	14,966	0	0.00%	0	0.00%	0	0.00%	1	0.24%	0	0.00%	1	0.24%
CD3265-40	100	48	3,649	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3249-40Q	990	431	39,827	0	0.00%	0	0.00%	0	0.00%	2	0.20%	1	0.10%	3	0.30%
CD3249-40	244	78	9,335	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.41%	1	0.41%
CD3231-40Q	1,679	725	83,509	2	0.12%	0	0.00%	0	0.00%	9	0.54%	1	0.06%	12	0.71%
CD3231-40	687	208	29,989	0	0.00%	0	0.00%	0	0.00%	3	0.44%	1	0.15%	4	0.58%
CD3211-36Q	856	240	43,584	3	0.35%	0	0.00%	0	0.00%	2	0.23%	2	0.23%	7	0.82%
CD3211-36	223	22	9,268	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.90%	2	0.90%
3207-36	673	59	30,734	1	0.15%	0	0.00%	0	0.00%	3	0.45%	3	0.45%	7	1.04%

A list of of complications can be found on [page 12](#).

Cardiac Resynchronization Therapy (CRT) ICDs

Actively Monitored Study Data Summary

MALFUNCTIONS WITH COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/ FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD3365-40Q	Quadra Assura™ CRT-D	222	12.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3357-40Q	Unify Assura™ CRT-D	219	10.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3357-40C	Unify Assura™ CRT-D	185	12.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3265-40Q	Quadra Assura™ CRT-D	421	16.60%	0	0.00%	1	0.24%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.24%
CD3265-40	Quadra Assura™ CRT-D	100	10.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3249-40Q	Unify Quadra™ CRT-D	990	13.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.10%	0	0.00%	0	0.00%	1	0.10%
CD3249-40	Unify Quadra™ CRT-D	244	20.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3231-40Q	Unify™ CRT-D	1,679	17.80%	0	0.00%	0	0.00%	1	0.06%	1	0.06%	0	0.00%	0	0.00%	11	0.66%	2	0.12%	0	0.00%	15	0.89%
CD3231-40	Unify™ CRT-D	687	20.20%	1	0.15%	1	0.15%	1	0.15%	0	0.00%	0	0.00%	0	0.00%	1	0.15%	0	0.00%	0	0.00%	4	0.58%
CD3211-36Q	Promote™ + CRT-D	856	29.60%	1	0.12%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	0	0.00%	2	0.23%	0	0.00%	0	0.00%	4	0.47%
CD3211-36	Promote™ + CRT-D	223	23.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
3207-36	Promote™ RF CRT-D	673	34.60%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.30%	0	0.00%	2	0.30%

MALFUNCTIONS WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/ FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD3365-40Q	Quadra Assura™ CRT-D	222	12.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3357-40Q	Unify Assura™ CRT-D	219	10.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3357-40C	Unify Assura™ CRT-D	185	12.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3265-40Q	Quadra Assura™ CRT-D	421	16.60%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3265-40	Quadra Assura™ CRT-D	100	10.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3249-40Q	Unify Quadra™ CRT-D	990	13.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.10%	0	0.00%	0	0.00%	1	0.10%
CD3249-40	Unify Quadra™ CRT-D	244	20.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD3231-40Q	Unify™ CRT-D	1,679	17.80%	1	0.06%	0	0.00%	1	0.06%	0	0.00%	0	0.00%	1	0.06%	1	0.06%	0	0.00%	0	0.00%	4	0.24%
CD3231-40	Unify™ CRT-D	687	20.20%	0	0.00%	0	0.00%	2	0.29%	0	0.00%	1	0.15%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	3	0.44%
CD3211-36Q	Promote™ + CRT-D	856	29.60%	1	0.12%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	0	0.00%	3	0.35%
CD3211-36	Promote™ + CRT-D	223	23.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.90%
3207-36	Promote™ RF CRT-D	673	34.60%	1	0.15%	0	0.00%	1	0.15%	0	0.00%	1	0.15%	0	0.00%	1	0.15%	1	0.15%	0	0.00%	5	0.74%

Definitions of malfunction categories can be found on [pages 5-6](#).

Cardiac Resynchronization Therapy (CRT) Pacemakers

Cardiac Resynchronization Therapy (CRT) Pacemakers

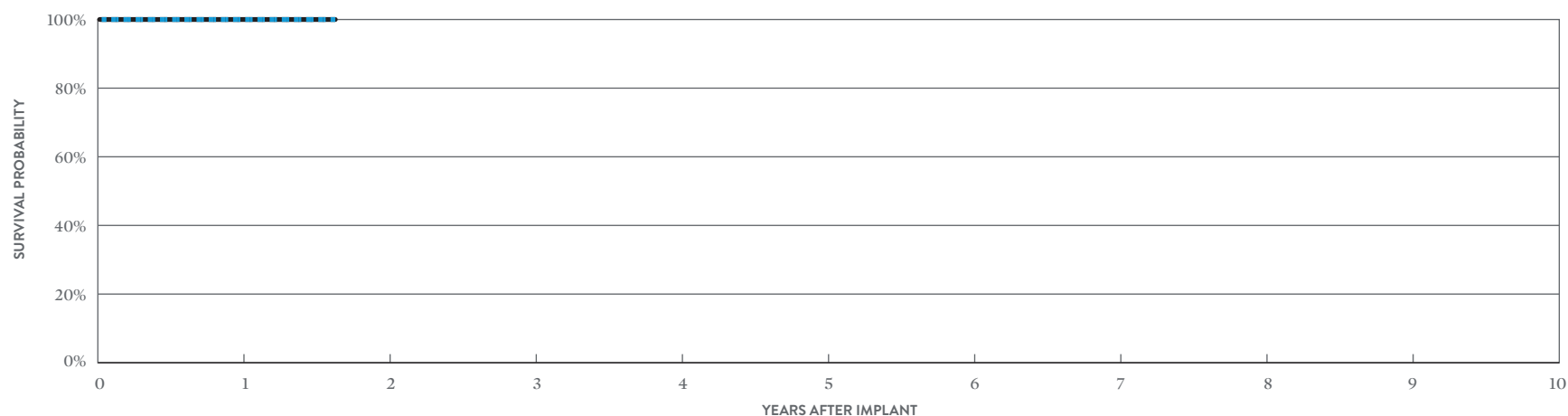
CUSTOMER REPORTED PERFORMANCE DATA

Allure Quadra MP™ CRT-P

MODEL PM3262

US Regulatory Approval	February 2016
Registered US Implants	10,812
Estimated Active US Implants	9,743
Estimated Longevity	8 Years
Normal Battery Depletion	0
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	AT 20 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%
SAMPLE SIZE	7,180	280

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	AT 20 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%

Cardiac Resynchronization Therapy (CRT) Pacemakers

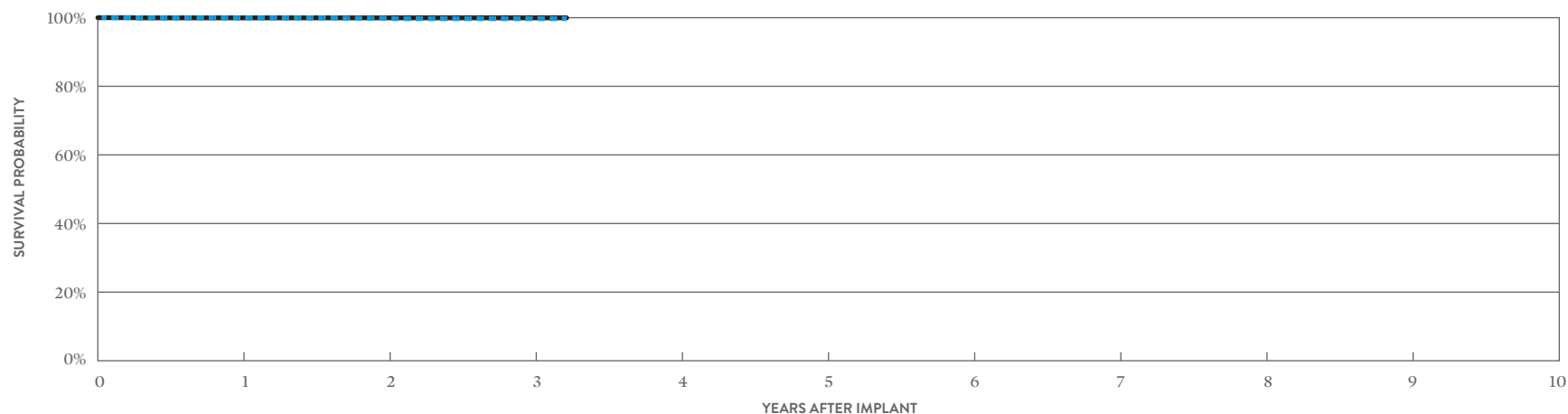
CUSTOMER REPORTED PERFORMANCE DATA

Allure™ RF CRT-P

MODEL PM3222

US Regulatory Approval	March 2014
Registered US Implants	4,925
Estimated Active US Implants	3,964
Estimated Longevity	8 Years
Normal Battery Depletion	3
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.02%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	1	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 39 MONTHS
SURVIVAL PROBABILITY	99.95%	99.85%	99.71%	99.71%
± 1 STANDARD ERROR	0.03%	0.08%	0.13%	0.13%
SAMPLE SIZE	3,910	2,200	930	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 39 MONTHS
SURVIVAL PROBABILITY	99.95%	99.95%	99.95%	99.95%
± 1 STANDARD ERROR	0.03%	0.03%	0.03%	0.03%

Cardiac Resynchronization Therapy (CRT) Pacemakers

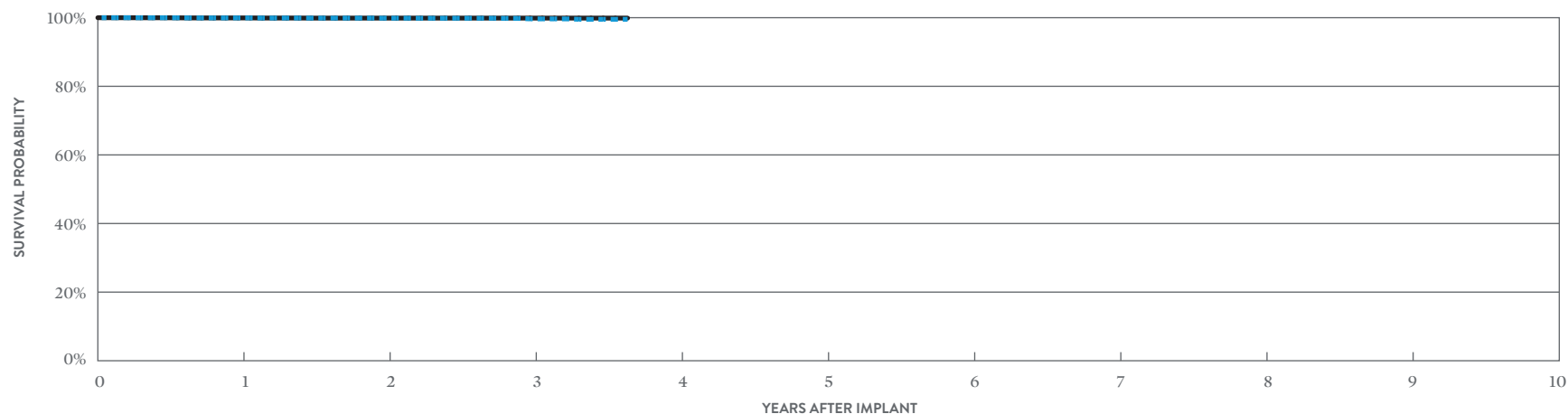
CUSTOMER REPORTED PERFORMANCE DATA

Allure Quadra™ RF CRT-P

MODEL PM3242

US Regulatory Approval	March 2014
Registered US Implants	17,718
Estimated Active US Implants	13,406
Estimated Longevity	8 Years
Normal Battery Depletion	6
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	1	<0.01%	7	0.04%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	1	<0.01%	7	0.04%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 44 MONTHS
SURVIVAL PROBABILITY	99.93%	99.86%	99.65%	99.53%
± 1 STANDARD ERROR	0.02%	0.03%	0.05%	0.12%
SAMPLE SIZE	16,290	12,380	6,560	420

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 44 MONTHS
SURVIVAL PROBABILITY	99.93%	99.88%	99.88%	99.88%
± 1 STANDARD ERROR	0.02%	0.03%	0.03%	0.03%

Cardiac Resynchronization Therapy (CRT) Pacemakers

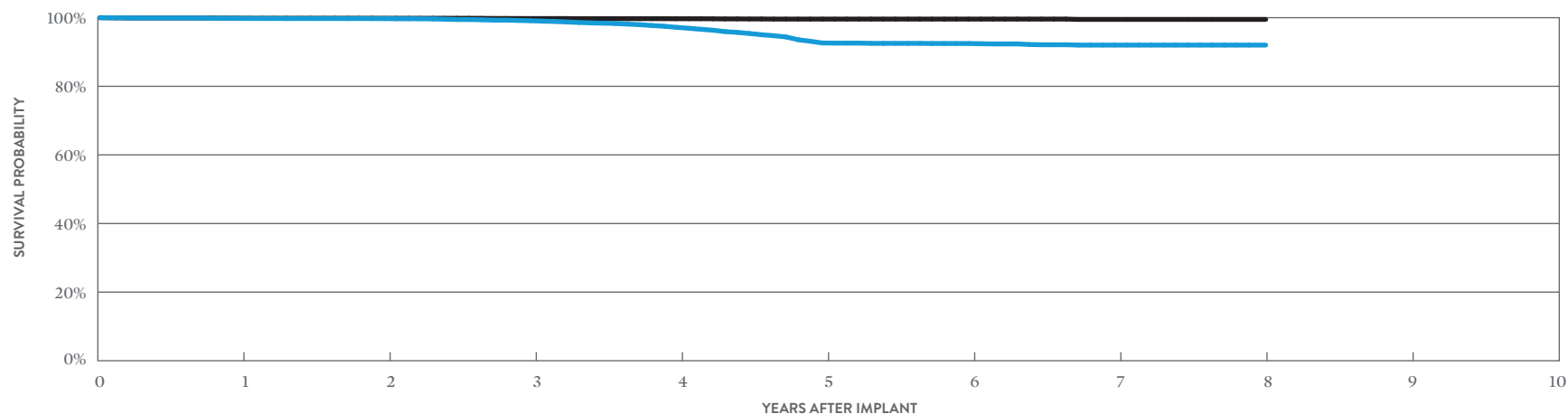
CUSTOMER REPORTED PERFORMANCE DATA

Anthem™ RF CRT-P

MODEL PM3210

US Regulatory Approval	July 2009
Registered US Implants	20,447
Estimated Active US Implants	9,525
Estimated Longevity	8 Years
Normal Battery Depletion	282
Number of US Advisories (see pgs. 334, 336)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.01%	4	0.02%
Electrical Interconnect	3	0.01%	1	<0.01%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	6	0.03%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	1	<0.01%	3	0.01%
Other	0	0.00%	7	0.03%
Total	7	0.03%	21	0.10%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8
SURVIVAL PROBABILITY	99.81%	99.72%	99.11%	97.19%	92.62%	92.47%	92.01%	92.01%
± 1 STANDARD ERROR	0.03%	0.04%	0.07%	0.14%	0.26%	0.28%	0.30%	0.30%
SAMPLE SIZE	18,820	16,180	14,250	11,720	8,190	4,900	2,410	260

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8
SURVIVAL PROBABILITY	99.87%	99.83%	99.75%	99.69%	99.59%	99.59%	99.49%	99.49%
± 1 STANDARD ERROR	0.03%	0.03%	0.04%	0.04%	0.06%	0.06%	0.09%	0.09%

Cardiac Resynchronization Therapy (CRT) Pacemakers

ACTIVELY MONITORED STUDY DATA

Anthem™ RF CRT-P

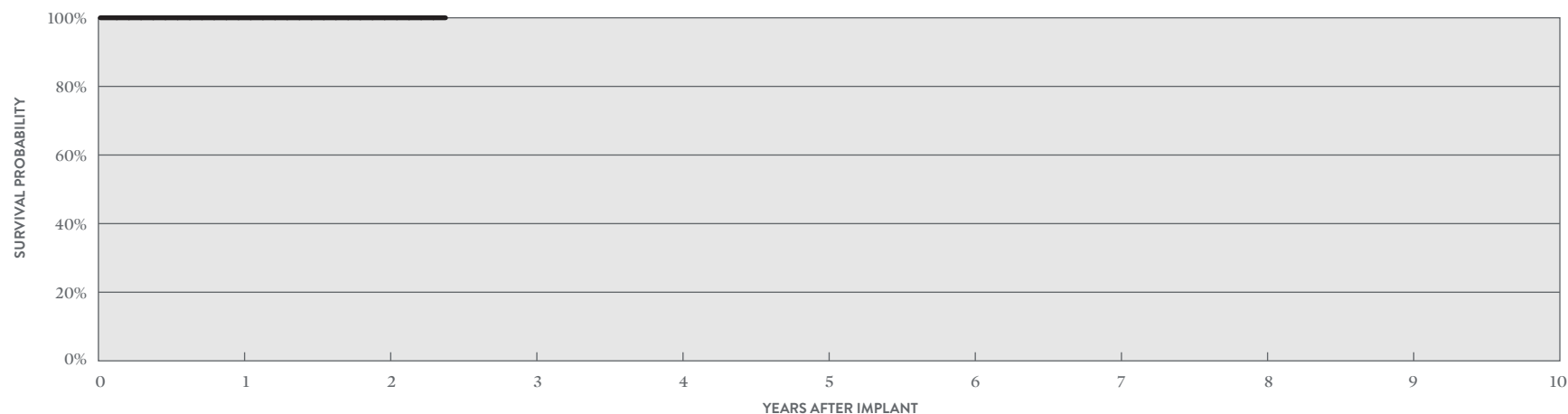
MODEL PM3210

US Regulatory Approval	July 2009
Number of Devices Enrolled in Study	201
Active Devices Enrolled in Study	26
Cumulative Months of Follow-up	5,053
Estimated Longevity	8 Years

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%
SAMPLE SIZE	170	100	50

Cardiac Resynchronization Therapy (CRT) Pacemakers

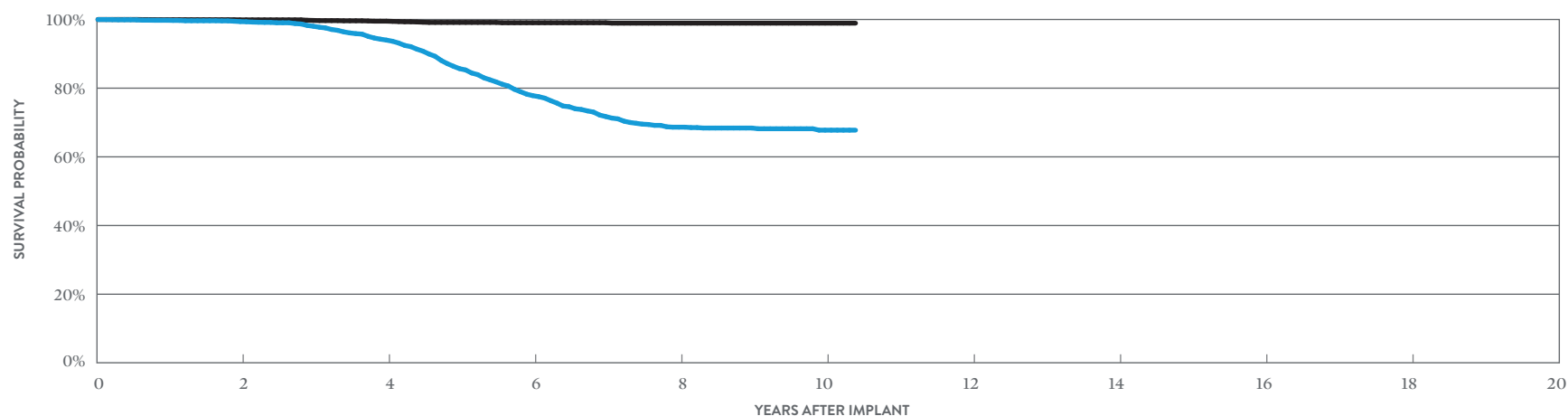
CUSTOMER REPORTED PERFORMANCE DATA

Frontier™ II CRT-P

MODEL 5586

US Regulatory Approval	August 2004
Registered US Implants	6,910
Estimated Active US Implants	1,016
Estimated Longevity	6.5 Years
Normal Battery Depletion	379
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	7	0.10%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	7	0.10%
Other	1	0.01%	3	0.04%
Total	1	0.01%	17	0.25%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 125 MONTHS
SURVIVAL PROBABILITY	99.38%	94.00%	77.83%	68.64%	67.77%	67.77%
± 1 STANDARD ERROR	0.10%	0.36%	0.72%	0.87%	0.93%	0.93%
SAMPLE SIZE	5,180	3,740	2,450	1,410	480	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 125 MONTHS
SURVIVAL PROBABILITY	99.89%	99.51%	99.05%	98.93%	98.93%	98.93%
± 1 STANDARD ERROR	0.03%	0.11%	0.16%	0.18%	0.18%	0.18%

SUMMARY INFORMATION

Cardiac Resynchronization
Therapy (CRT) Pacemakers

Cardiac Resynchronization Therapy (CRT) Pacemakers

Survival Probability Summary

INCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
PM3262	Allure Quadra MP [™] CRT-P	100.00%									
PM3222	Allure [™] RF CRT-P	99.95%	99.85%	99.71%							
PM3242	Allure Quadra [™] RF CRT-P	99.93%	99.86%	99.65%							
PM3210	Anthem [™] RF CRT-P	99.81%	99.72%	99.11%	97.19%	92.62%	92.47%	92.01%	92.01%		
5586	Frontier [™] II CRT-P	99.76%	99.38%	98.03%	94.00%	85.68%	77.83%	71.73%	68.64%	68.39%	67.77%

EXCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
PM3262	Allure Quadra MP [™] CRT-P	100.00%									
PM3222	Allure [™] RF CRT-P	99.95%	99.95%	99.95%							
PM3242	Allure Quadra [™] RF CRT-P	99.93%	99.88%	99.88%							
PM3210	Anthem [™] RF CRT-P	99.87%	99.83%	99.75%	99.69%	99.59%	99.59%	99.49%	99.49%		
5586	Frontier [™] II CRT-P	99.93%	99.89%	99.72%	99.51%	99.14%	99.05%	99.05%	98.93%	98.93%	98.93%

Cardiac Resynchronization Therapy (CRT) Pacemakers

US Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM3262	Allure Quadra MP [™] CRT-P	10,812	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM3222	Allure [™] RF CRT-P	4,925	0.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM3242	Allure Quadra [™] RF CRT-P	17,718	0.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
PM3210	Anthem [™] RF CRT-P	20,447	4.30%	3	0.01%	3	0.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	7	0.03%
5586	Frontier [™] II CRT-P	6,910	14.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.01%	1	0.01%	1	0.01%

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM3262	Allure Quadra MP [™] CRT-P	10,812	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM3222	Allure [™] RF CRT-P	4,925	0.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	1	0.02%
PM3242	Allure Quadra [™] RF CRT-P	17,718	0.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	7	0.04%	0	0.00%	0	0.00%	0	0.00%	7	0.04%
PM3210	Anthem [™] RF CRT-P	20,447	4.30%	4	0.02%	1	<0.01%	0	0.00%	6	0.03%	0	0.00%	3	0.01%	7	0.03%	0	0.00%	21	0.10%
5586	Frontier [™] II CRT-P	6,910	14.30%	7	0.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	7	0.10%	3	0.04%	0	0.00%	17	0.25%

Definitions of malfunction categories can be found on [pages 5-6](#).

Cardiac Resynchronization Therapy (CRT) Pacemakers Worldwide Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM3262	Allure Quadra MP [™] CRT-P	22,955	0.83%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM3222	Allure [™] RF CRT-P	15,912	1.22%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
PM3242	Allure Quadra [™] RF CRT-P	34,245	2.06%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
PM3210	Anthem [™] RF CRT-P	21,092	10.07%	3	0.01%	3	0.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	7	0.03%

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM3262	Allure Quadra MP [™] CRT-P	22,955	0.83%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM3222	Allure [™] RF CRT-P	15,912	1.22%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
PM3242	Allure Quadra [™] RF CRT-P	34,245	2.06%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	9	0.03%	1	<0.01%	1	<0.01%	1	<0.01%	12	0.04%
PM3210	Anthem [™] RF CRT-P	21,092	10.07%	3	0.01%	1	<0.01%	0	0.00%	6	0.03%	0	0.00%	3	0.01%	7	0.03%	0	0.00%	20	0.09%

Definitions of malfunction categories can be found on [pages 5-6](#).

Cardiac Resynchronization Therapy (CRT) Pacemakers

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	LOSS OF TELEMETRY		PERICARDIAL EFFUSION		PREMATURE BATTERY DEPLETION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM3210	201	26	5,053	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

MALFUNCTIONS WITH COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM3210	Anthem [™] RF	201	6.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

MALFUNCTIONS WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM3210	Anthem [™] RF	201	6.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

A list of complications can be found on [page 12](#).

Definitions of malfunction categories can be found on [pages 5-6](#).

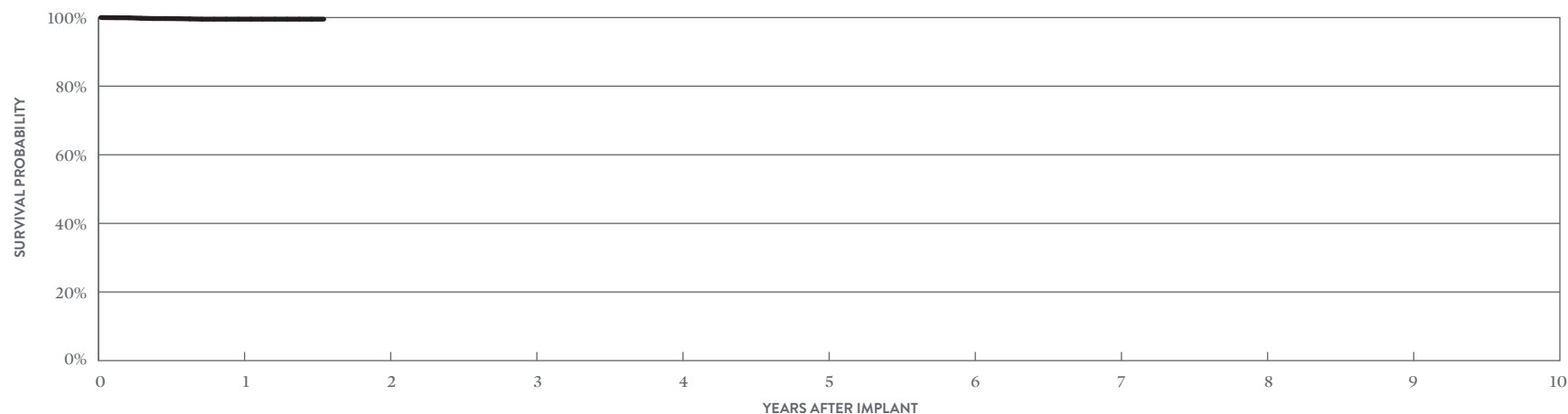
Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

Quartet™

MODEL 1458QL

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	October 2015	Cardiac Perforation	0	0.00%	0	0.00%	Conductor Fracture	0	0.00%
Registered US Implants	4,706	Conductor Fracture	0	0.00%	0	0.00%	Clavicular Crush	0	0.00%
Estimated Active US Implants	4,224	Lead Dislodgement	8	0.17%	13	0.28%	In the Pocket	0	0.00%
Insulation	Optim™*	Failure to Capture	8	0.17%	5	0.11%	Intravascular	0	0.00%
Type and/or Fixation	S-Curve	Oversensing	0	0.00%	0	0.00%	Insulation Breach	0	0.00%
Polarity	Quadpolar	Failure to Sense	0	0.00%	0	0.00%	Lead-to-Can Contact	0	0.00%
Steroid	Yes	Insulation Breach	0	0.00%	0	0.00%	Lead-to-Lead Contact	0	0.00%
Number of US Advisories	None	Abnormal Pacing Impedance	1	0.02%	0	0.00%	Clavicular Crush	0	0.00%
		Extracardiac Stimulation	9	0.19%	11	0.23%	Externalized Conductors	0	0.00%
		Other	2	0.04%	0	0.00%	Other	0	0.00%
		Total	28	0.59%	29	0.62%	Crimps, Welds & Bonds	0	0.00%
		Total Returned for Analysis	4		11		Other	0	0.00%
							Extrinsic Factors	10	0.21%
							Total	10	0.21%



YEAR	1	AT 19 MONTHS
SURVIVAL PROBABILITY	99.53%	99.53%
± 1 STANDARD ERROR	0.12%	0.12%
SAMPLE SIZE	3,180	320

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

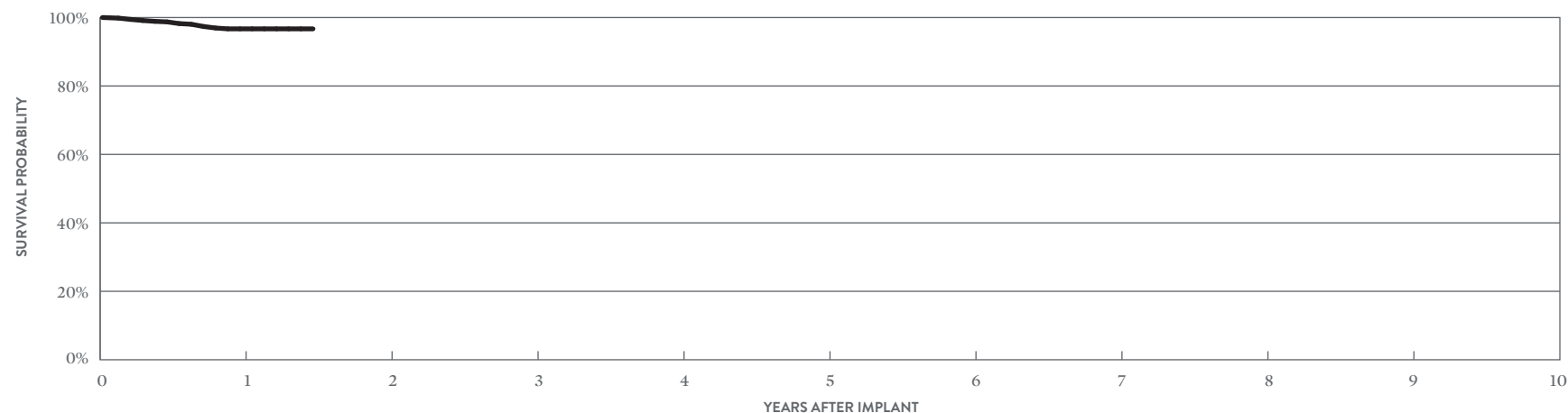
Quartet™

MODEL 1457Q

US Regulatory Approval	March 2017
Registered US Implants	1,430
Estimated Active US Implants	1,270
Insulation	Optim*
Type and/or Fixation	S-Curve
Polarity	Quadpolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	0	0.00%
Conductor Fracture	0	0.00%	0	0.00%
Lead Dislodgement	8	0.56%	12	0.84%
Failure to Capture	0	0.00%	2	0.14%
Oversensing	1	0.07%	0	0.00%
Failure to Sense	0	0.00%	0	0.00%
Insulation Breach	0	0.00%	0	0.00%
Abnormal Pacing Impedance	0	0.00%	0	0.00%
Extracardiac Stimulation	11	0.77%	5	0.35%
Other	2	0.14%	2	0.14%
Total	22	1.54%	21	1.47%
Total Returned for Analysis	2		8	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	0	0.00%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	13	0.91%
Total	13	0.91%



YEAR	1	AT 18 MONTHS
SURVIVAL PROBABILITY	96.69%	96.69%
± 1 STANDARD ERROR	0.74%	0.74%
SAMPLE SIZE	890	200

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

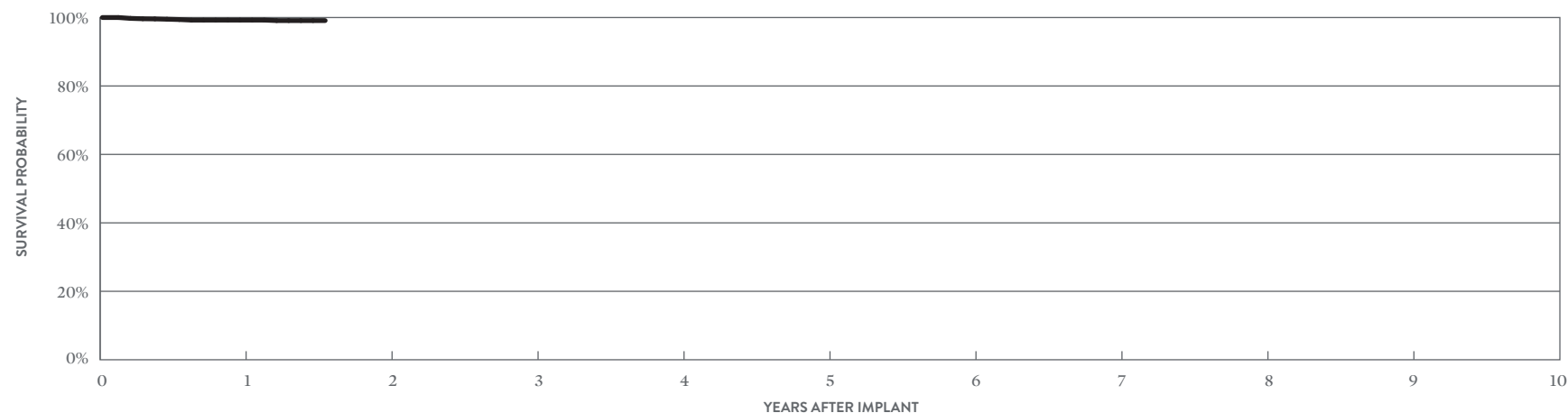
Quartet™

MODEL 1456Q

US Regulatory Approval	October 2015
Registered US Implants	3,358
Estimated Active US Implants	2,995
Insulation	Optim™*
Type and/or Fixation	S-Curve
Polarity	Quadpolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	1	0.03%	0	0.00%
Conductor Fracture	2	0.06%	0	0.00%
Lead Dislodgement	7	0.21%	14	0.42%
Failure to Capture	0	0.00%	5	0.15%
Oversensing	0	0.00%	0	0.00%
Failure to Sense	0	0.00%	0	0.00%
Insulation Breach	0	0.00%	0	0.00%
Abnormal Pacing Impedance	0	0.00%	0	0.00%
Extracardiac Stimulation	8	0.24%	4	0.12%
Other	3	0.09%	1	0.03%
Total	21	0.63%	24	0.71%
Total Returned for Analysis	3		12	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	0	0.00%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	12	0.36%
Total	12	0.36%



YEAR	1	AT 19 MONTHS
SURVIVAL PROBABILITY	99.25%	99.11%
± 1 STANDARD ERROR	0.18%	0.22%
SAMPLE SIZE	2,230	220

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

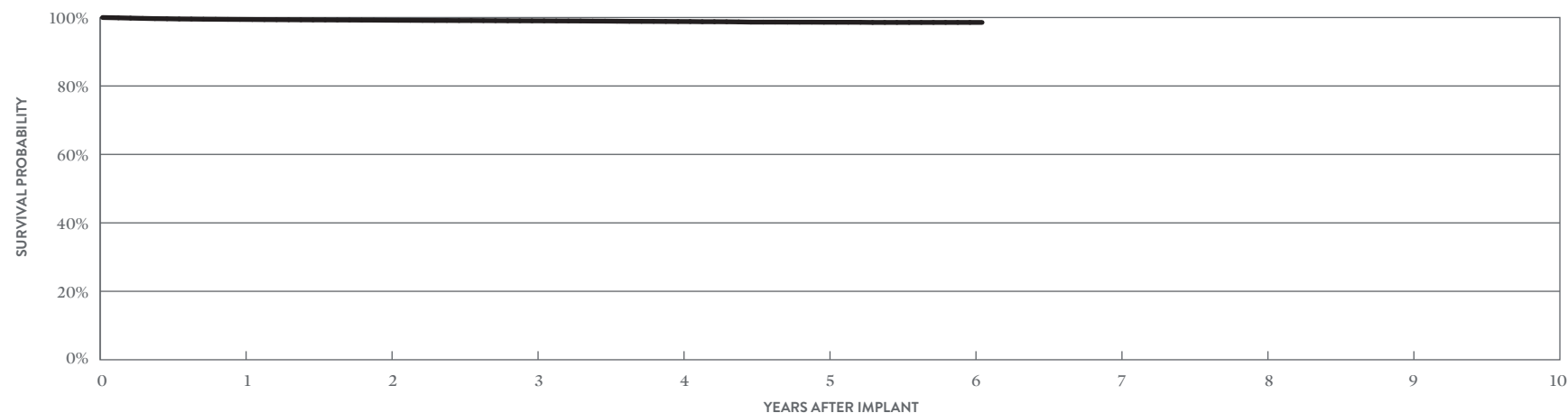
Quartet™

MODEL 1458Q

US Regulatory Approval	November 2011
Registered US Implants	122,300
Estimated Active US Implants	87,211
Insulation	Optim*
Type and/or Fixation	S-Curve
Polarity	Quadpolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	3	<0.01%	2	<0.01%
Conductor Fracture	0	0.00%	13	0.01%
Lead Dislodgement	170	0.14%	676	0.55%
Failure to Capture	72	0.06%	280	0.23%
Oversensing	2	<0.01%	8	<0.01%
Failure to Sense	0	0.00%	0	0.00%
Insulation Breach	1	<0.01%	6	<0.01%
Abnormal Pacing Impedance	4	<0.01%	46	0.04%
Extracardiac Stimulation	88	0.07%	148	0.12%
Other	75	0.06%	28	0.02%
Total	415	0.34%	1207	0.99%
Total Returned for Analysis	153		496	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	7	<0.01%
Clavicular Crush	0	0.00%
In the Pocket	2	<0.01%
Intravascular	5	<0.01%
Insulation Breach	3	<0.01%
Lead-to-Can Contact	1	<0.01%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	2	<0.01%
Crimps, Welds & Bonds	0	0.00%
Other	9	<0.01%
Extrinsic Factors	475	0.39%
Total	495	0.40%



YEAR	1	2	3	4	5	6	AT 73 MONTHS
SURVIVAL PROBABILITY	99.44%	99.21%	99.04%	98.83%	98.62%	98.57%	98.57%
± 1 STANDARD ERROR	0.02%	0.03%	0.03%	0.04%	0.06%	0.06%	0.06%
SAMPLE SIZE	106,870	78,200	51,730	28,950	13,660	4,130	320

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Left-Heart Leads

ACTIVELY MONITORED STUDY DATA

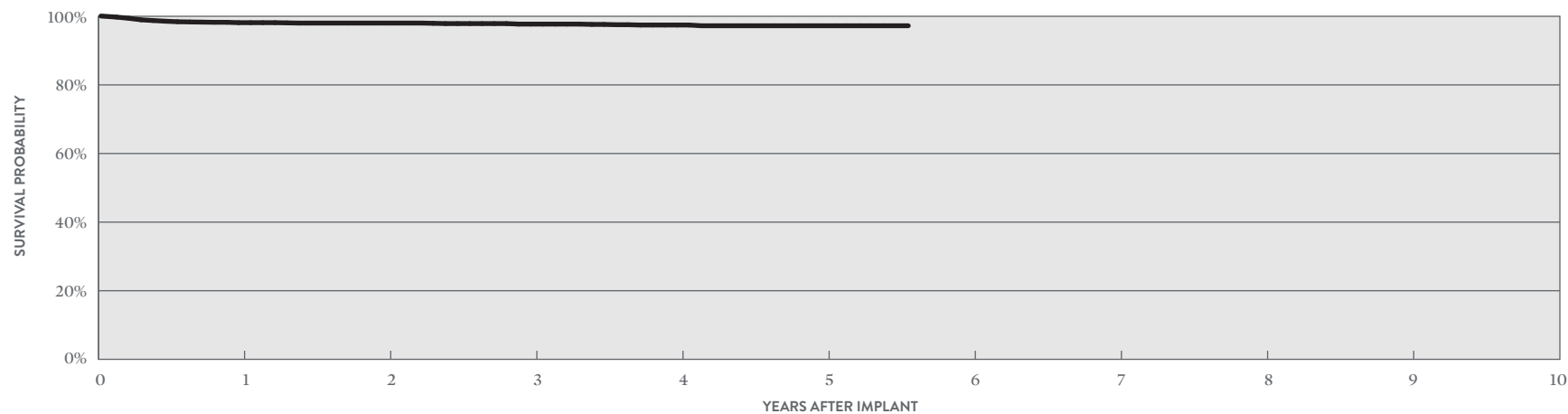
Quartet™

MODEL 1458Q

US Regulatory Approval	November 2011
Number of Devices Enrolled in Study	2,108
Active Devices Enrolled in Study	1,044
Cumulative Months of Follow-up	79,263
Insulation	Optim™*
Type and/or Fixation	S-Curve
Polarity	Quadpolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Pacing Impedance	1	0.05%
Extracardiac Stimulation	3	0.14%
Failure to Capture	5	0.24%
Lead Dislodgement	38	1.80%
Oversensing	1	0.05%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	0	0.00%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	21	1.00%
Total	21	1.00%



YEAR	1	2	3	4	5	AT 67 MONTHS
SURVIVAL PROBABILITY	98.14%	98.02%	97.72%	97.45%	97.24%	97.24%
± 1 STANDARD ERROR	0.29%	0.31%	0.35%	0.38%	0.41%	0.41%
SAMPLE SIZE	1,950	1,620	1,330	1,120	650	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

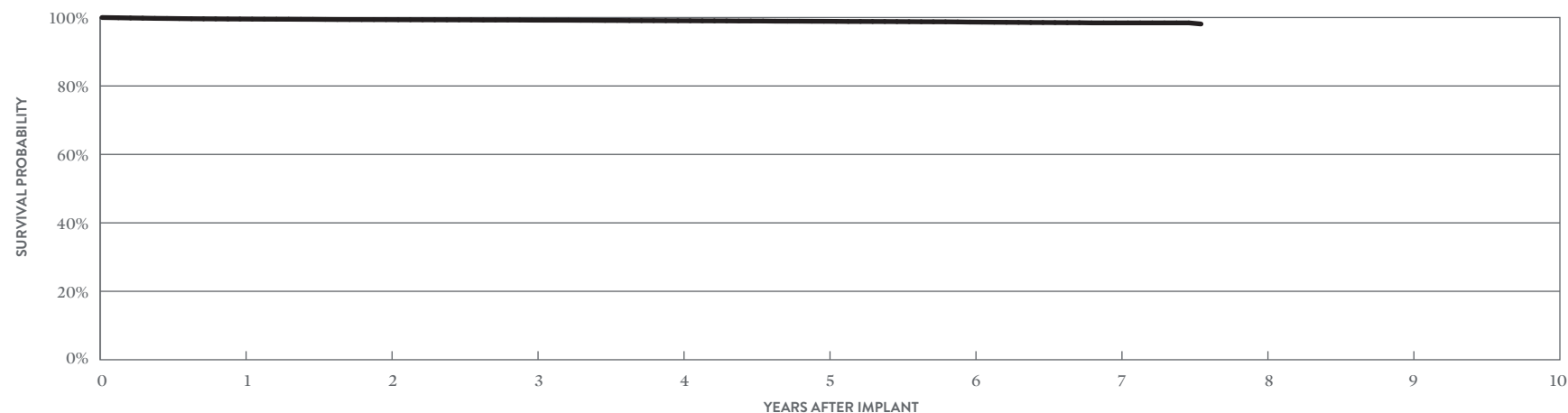
QuickFlex™ μ

MODEL 1258T

US Regulatory Approval	May 2010
Registered US Implants	46,740
Estimated Active US Implants	26,698
Insulation	Optim™*
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	1	<0.01%
Conductor Fracture	0	0.00%	20	0.04%
Lead Dislodgement	45	0.10%	191	0.41%
Failure to Capture	17	0.04%	158	0.34%
Oversensing	0	0.00%	15	0.03%
Failure to Sense	1	<0.01%	2	<0.01%
Insulation Breach	0	0.00%	7	0.01%
Abnormal Pacing Impedance	5	0.01%	41	0.09%
Extracardiac Stimulation	19	0.04%	68	0.15%
Other	12	0.03%	9	0.02%
Total	99	0.21%	512	1.10%
Total Returned for Analysis	53		197	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	5	0.01%
Clavicular Crush	1	<0.01%
In the Pocket	1	<0.01%
Intravascular	3	<0.01%
Insulation Breach	3	<0.01%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	2	<0.01%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	1	<0.01%
Crimps, Welds & Bonds	0	0.00%
Other	1	<0.01%
Extrinsic Factors	213	0.46%
Total	222	0.47%



YEAR	1	2	3	4	5	6	7	AT 91 MONTHS
SURVIVAL PROBABILITY	99.56%	99.38%	99.23%	99.07%	98.90%	98.67%	98.44%	98.11%
± 1 STANDARD ERROR	0.03%	0.04%	0.04%	0.05%	0.06%	0.07%	0.09%	0.09%
SAMPLE SIZE	42,980	36,640	31,610	26,020	19,440	13,210	6,700	440

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Left-Heart Leads

ACTIVELY MONITORED STUDY DATA

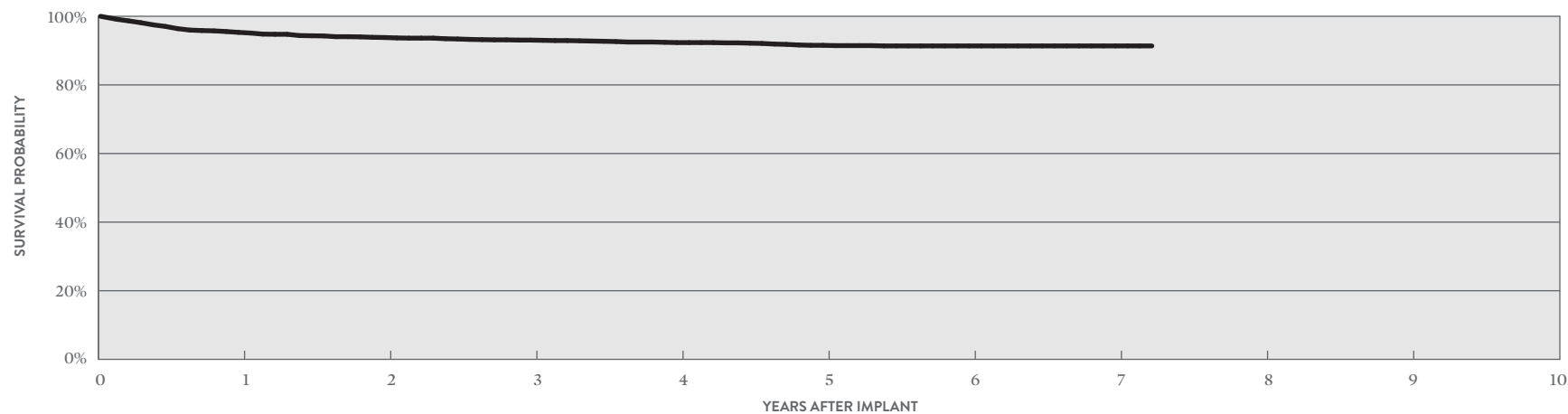
QuickFlex™ μ

MODEL 1258T

US Regulatory Approval	May 2010
Number of Devices Enrolled in Study	2,367
Active Devices Enrolled in Study	1,015
Cumulative Months of Follow-up	107,780
Insulation	Optim™*
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Pacing Impedance	6	0.25%
Conductor Fracture	2	0.08%
Extracardiac Stimulation	56	2.37%
Failure to Capture	49	2.07%
Lead Dislodgement	52	2.20%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	1	0.04%
Clavicular Crush	1	0.04%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	0	0.00%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	35	1.48%
Total	36	1.52%



YEAR	1	2	3	4	5	6	7	AT 87 MONTHS
SURVIVAL PROBABILITY	95.31%	93.81%	93.06%	92.33%	91.54%	91.36%	91.36%	91.36%
± 1 STANDARD ERROR	0.44%	0.52%	0.56%	0.60%	0.65%	0.67%	0.67%	0.67%
SAMPLE SIZE	2,150	1,760	1,480	1,270	1,080	880	440	70

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

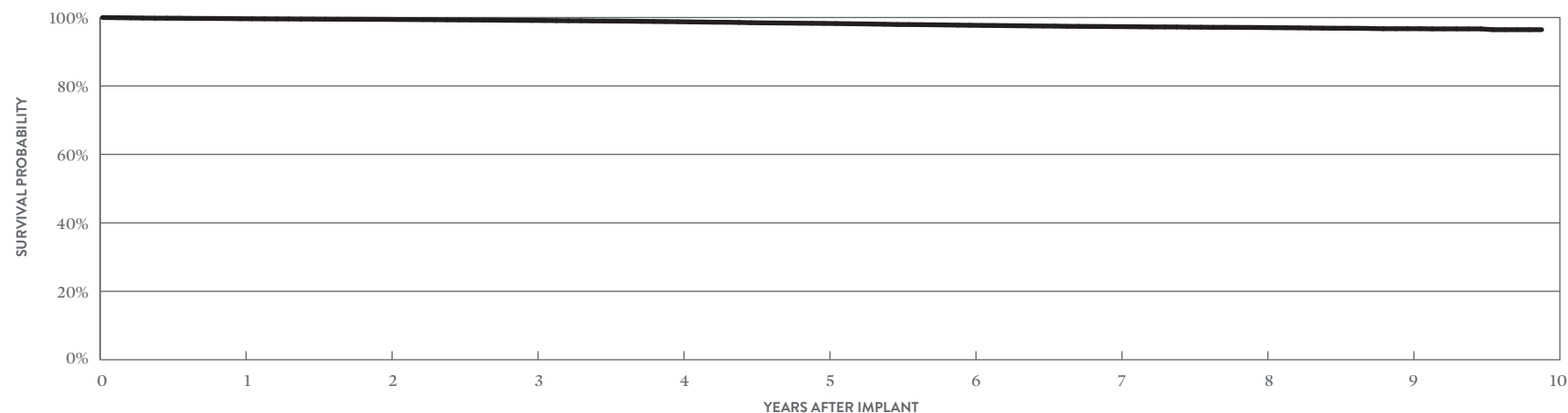
QuickFlex™

MODEL 1156T

US Regulatory Approval	July 2007
Registered US Implants	27,662
Estimated Active US Implants	11,431
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes
Number of US Advisories (see pg. 341)	One

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	1	<0.01%
Conductor Fracture	0	0.00%	5	0.02%
Lead Dislodgement	11	0.04%	133	0.48%
Failure to Capture	4	0.01%	183	0.66%
Oversensing	0	0.00%	14	0.05%
Failure to Sense	0	0.00%	0	0.00%
Insulation Breach	0	0.00%	46	0.17%
Abnormal Pacing Impedance	0	0.00%	61	0.22%
Extracardiac Stimulation	13	0.05%	85	0.31%
Other	9	0.03%	8	0.03%
Total	37	0.13%	536	1.94%
Total Returned for Analysis	14		156	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	6	0.02%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	6	0.02%
Insulation Breach	81	0.29%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	4	0.01%
Clavicular Crush	0	0.00%
Externalized Conductors	13	0.05%
Other	64	0.23%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	128	0.46%
Total	215	0.78%



YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.65%	99.45%	99.15%	98.77%	98.28%	97.74%	97.34%	97.08%	96.73%	96.45%
± 1 STANDARD ERROR	0.03%	0.05%	0.06%	0.08%	0.09%	0.11%	0.13%	0.14%	0.16%	0.23%
SAMPLE SIZE	25,310	21,570	19,100	17,110	15,400	13,780	11,580	8,180	4,230	260

Left-Heart Leads

ACTIVELY MONITORED STUDY DATA

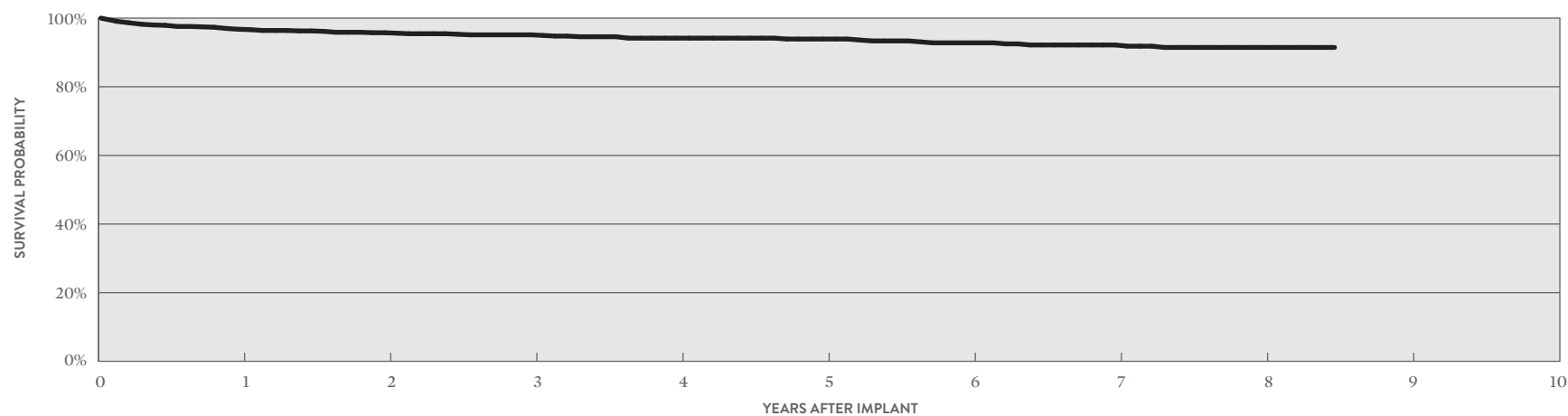
QuickFlex™

MODEL 1156T

US Regulatory Approval	July 2007
Number of Devices Enrolled in Study	985
Active Devices Enrolled in Study	264
Cumulative Months of Follow-up	47,545
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Pacing Impedance	1	0.10%
Extracardiac Stimulation	17	1.73%
Failure to Capture	9	0.91%
Insulation Breach	1	0.10%
Lead Dislodgement	27	2.74%
Skin Erosion	1	0.10%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	3	0.30%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	3	0.30%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	19	1.93%
Total	22	2.23%



YEAR	1	2	3	4	5	6	7	8	AT 102 MONTHS
SURVIVAL PROBABILITY	96.75%	95.74%	95.13%	94.16%	93.90%	92.79%	92.17%	91.45%	91.45%
± 1 STANDARD ERROR	0.56%	0.68%	0.74%	0.85%	0.89%	1.04%	1.12%	1.22%	1.22%
SAMPLE SIZE	900	750	610	480	380	340	290	200	60

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

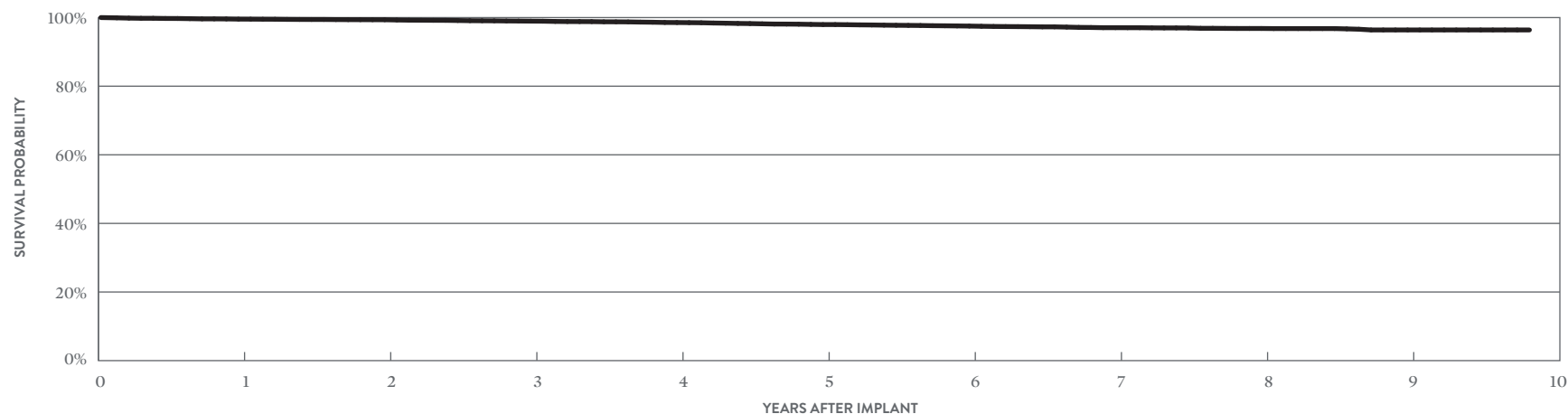
QuickFlex™ XL

MODEL 1158T

US Regulatory Approval	July 2007
Registered US Implants	15,337
Estimated Active US Implants	6,405
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes
Number of US Advisories (see pg. 341)	One

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	1	<0.01%
Conductor Fracture	0	0.00%	4	0.03%
Lead Dislodgement	9	0.06%	95	0.62%
Failure to Capture	2	0.01%	126	0.82%
Oversensing	0	0.00%	3	0.02%
Failure to Sense	0	0.00%	1	<0.01%
Insulation Breach	0	0.00%	35	0.23%
Abnormal Pacing Impedance	2	0.01%	23	0.15%
Extracardiac Stimulation	6	0.04%	32	0.21%
Other	6	0.04%	8	0.05%
Total	25	0.16%	328	2.14%
Total Returned for Analysis	13		112	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	5	0.03%
Clavicular Crush	0	0.00%
In the Pocket	1	<0.01%
Intravascular	4	0.03%
Insulation Breach	53	0.35%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	2	0.01%
Clavicular Crush	0	0.00%
Externalized Conductors	7	0.05%
Other	44	0.29%
Crimps, Welds & Bonds	1	<0.01%
Other	0	0.00%
Extrinsic Factors	86	0.56%
Total	145	0.95%



YEAR	1	2	3	4	5	6	7	8	9	AT 118 MONTHS
SURVIVAL PROBABILITY	99.55%	99.36%	98.96%	98.55%	97.96%	97.52%	97.05%	96.82%	96.39%	96.39%
± 1 STANDARD ERROR	0.05%	0.07%	0.09%	0.11%	0.14%	0.16%	0.18%	0.19%	0.24%	0.24%
SAMPLE SIZE	14,050	12,030	10,670	9,520	8,550	7,640	6,250	4,270	2,280	270

Left-Heart Leads

ACTIVELY MONITORED STUDY DATA

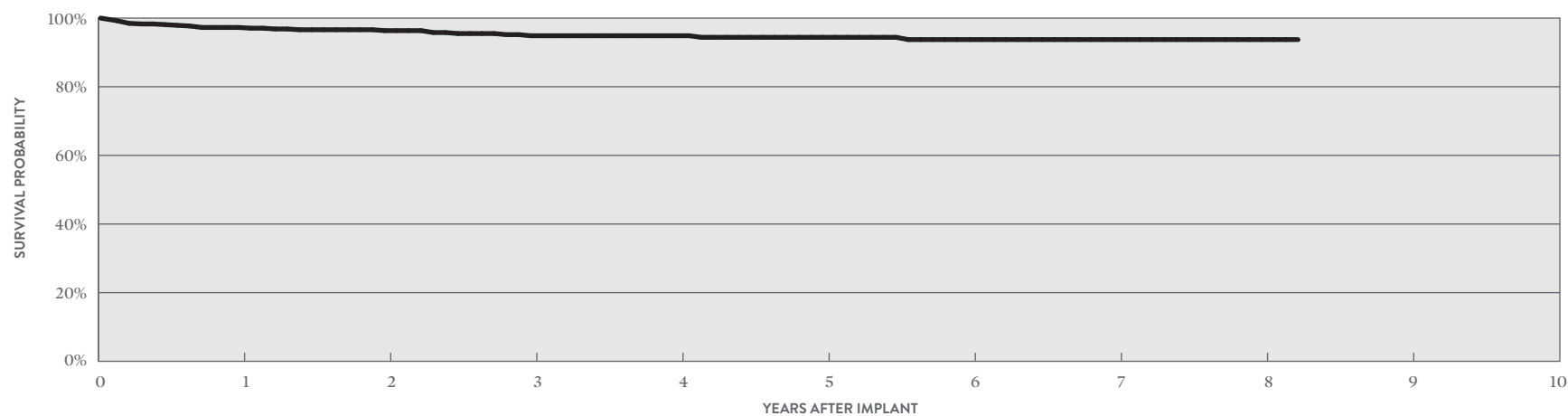
QuickFlex™ XL

MODEL 1158T

US Regulatory Approval	July 2007
Number of Devices Enrolled in Study	553
Active Devices Enrolled in Study	120
Cumulative Months of Follow-up	24,555
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Extracardiac Stimulation	9	1.63%
Failure to Capture	8	1.45%
Insulation Breach	1	0.18%
Lead Dislodgement	6	1.08%
Skin Erosion	1	0.18%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	1	0.18%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	1	0.18%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	7	1.27%
Total	8	1.45%



YEAR	1	2	3	4	5	6	7	8	AT 99 MONTHS
SURVIVAL PROBABILITY	97.27%	96.35%	94.86%	94.86%	94.40%	93.74%	93.74%	93.74%	93.74%
± 1 STANDARD ERROR	0.72%	0.81%	1.02%	1.07%	1.15%	1.32%	1.32%	1.32%	1.32%
SAMPLE SIZE	500	410	330	250	190	150	120	80	50

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

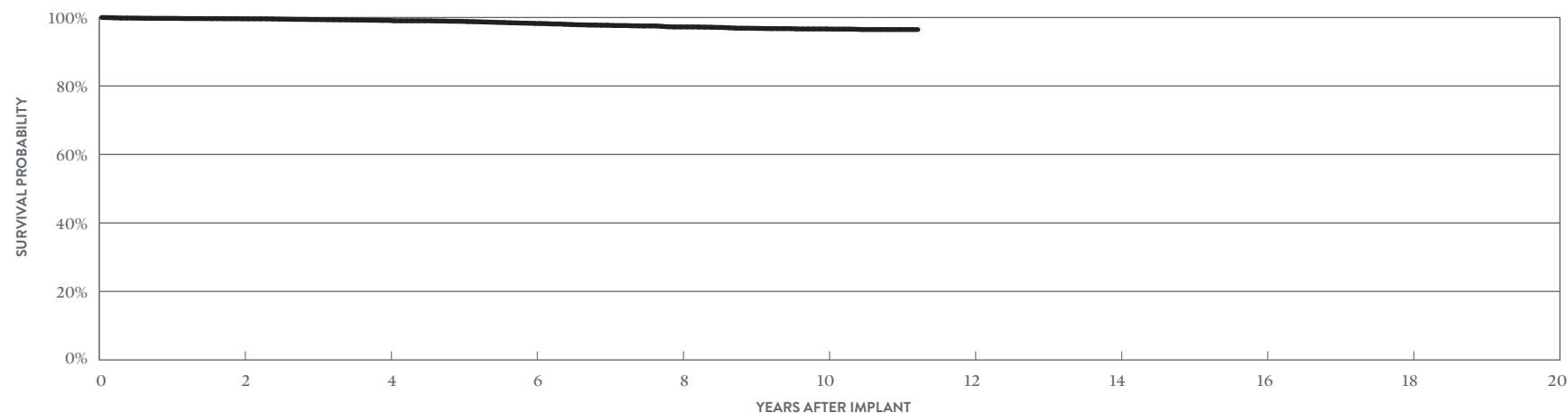
QuickSite™ XL

MODEL 1058T

US Regulatory Approval	February 2006
Registered US Implants	9,951
Estimated Active US Implants	3,399
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes
Number of US Advisories (see pg. 341)	One

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	0	0.00%
Conductor Fracture	0	0.00%	5	0.05%
Lead Dislodgement	10	0.10%	29	0.29%
Failure to Capture	3	0.03%	81	0.81%
Oversensing	1	0.01%	2	0.02%
Failure to Sense	0	0.00%	2	0.02%
Insulation Breach	0	0.00%	32	0.32%
Abnormal Pacing Impedance	2	0.02%	19	0.19%
Extracardiac Stimulation	9	0.09%	23	0.23%
Other	1	0.01%	2	0.02%
Total	26	0.26%	195	1.96%
Total Returned for Analysis	11		36	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	2	0.02%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	2	0.02%
Insulation Breach	24	0.24%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	1	0.01%
Clavicular Crush	0	0.00%
Externalized Conductors	6	0.06%
Other	17	0.17%
Crimps, Welds & Bonds	0	0.00%
Other	1	0.01%
Extrinsic Factors	29	0.29%
Total	56	0.56%



YEAR	2	4	6	8	10	AT 135 MONTHS
SURVIVAL PROBABILITY	99.62%	99.15%	98.27%	97.24%	96.65%	96.48%
± 1 STANDARD ERROR	0.06%	0.11%	0.17%	0.23%	0.26%	0.28%
SAMPLE SIZE	7,810	5,980	4,750	3,890	2,790	280

Left-Heart Leads

ACTIVELY MONITORED STUDY DATA

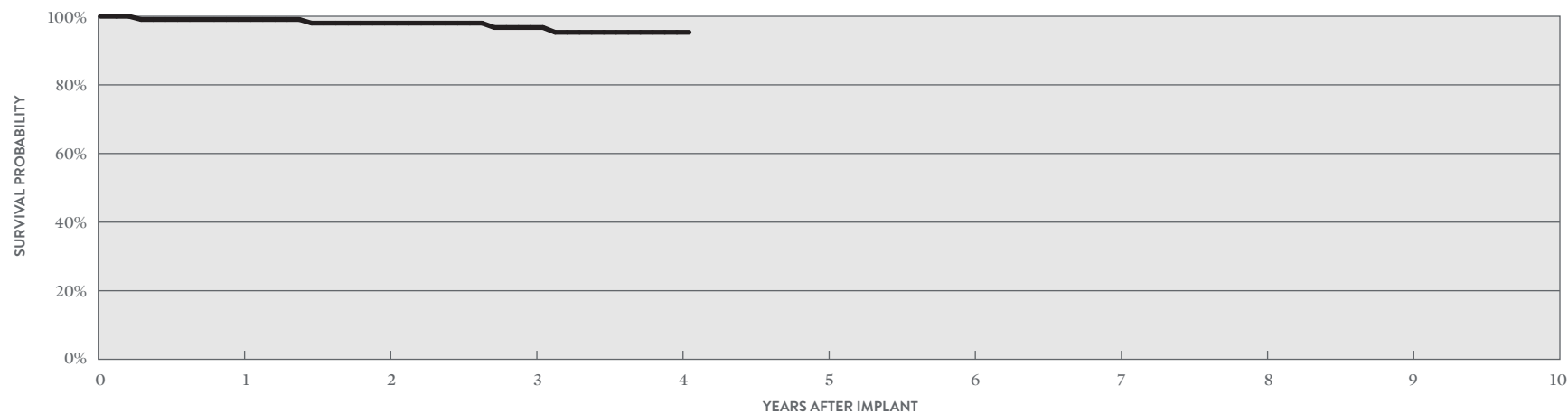
QuickSite™ XL

MODEL 1058T

US Regulatory Approval	February 2006
Number of Devices Enrolled in Study	111
Active Devices Enrolled in Study	19
Cumulative Months of Follow-up	5,486
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Failure to Capture	4	3.60%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	0	0.00%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	0	0.00%
Total	0	0.00%



YEAR	1	2	3	4	AT 49 MONTHS
SURVIVAL PROBABILITY	99.07%	98.01%	96.74%	95.33%	95.33%
± 1 STANDARD ERROR	0.92%	1.39%	1.87%	2.31%	2.31%
SAMPLE SIZE	100	90	80	60	50

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

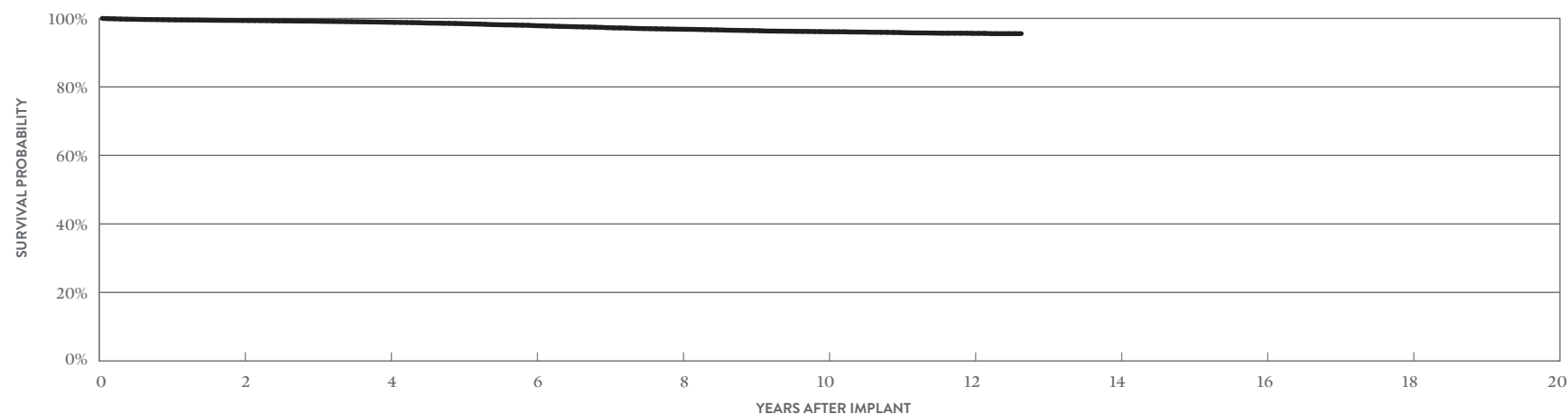
QuickSite™

MODEL 1056T

US Regulatory Approval	April 2005
Registered US Implants	32,332
Estimated Active US Implants	9,782
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes
Number of US Advisories (see pg. 341)	One

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	0	0.00%
Conductor Fracture	0	0.00%	9	0.03%
Lead Dislodgement	32	0.10%	166	0.51%
Failure to Capture	15	0.05%	270	0.84%
Oversensing	2	<0.01%	21	0.06%
Failure to Sense	0	0.00%	1	<0.01%
Insulation Breach	1	<0.01%	105	0.32%
Abnormal Pacing Impedance	3	<0.01%	59	0.18%
Extracardiac Stimulation	22	0.07%	103	0.32%
Other	9	0.03%	20	0.06%
Total	84	0.26%	754	2.33%
Total Returned for Analysis	28		196	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	6	0.02%
Clavicular Crush	0	0.00%
In the Pocket	2	<0.01%
Intravascular	4	0.01%
Insulation Breach	88	0.27%
Lead-to-Can Contact	1	<0.01%
Lead-to-Lead Contact	11	0.03%
Clavicular Crush	0	0.00%
Externalized Conductors	31	0.10%
Other	45	0.14%
Crimps, Welds & Bonds	0	0.00%
Other	1	<0.01%
Extrinsic Factors	158	0.49%
Total	253	0.78%



YEAR	2	4	6	8	10	12	AT 152 MONTHS
SURVIVAL PROBABILITY	99.39%	98.88%	97.89%	96.84%	96.11%	95.63%	95.55%
± 1 STANDARD ERROR	0.05%	0.07%	0.10%	0.14%	0.16%	0.18%	0.21%
SAMPLE SIZE	25,430	19,400	14,870	11,860	8,900	3,290	200

Left-Heart Leads

ACTIVELY MONITORED STUDY DATA

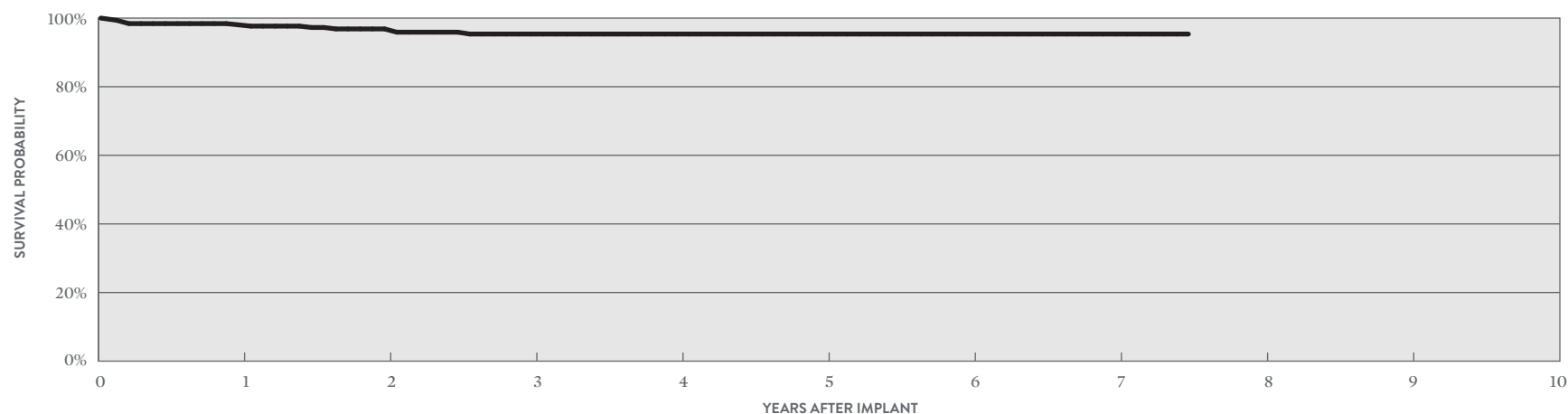
QuickSite™

MODEL 1056T

US Regulatory Approval	April 2005
Number of Devices Enrolled in Study	319
Active Devices Enrolled in Study	66
Cumulative Months of Follow-up	14,237
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Pacing Impedance	1	0.31%
Extracardiac Stimulation	2	0.63%
Failure to Capture	4	1.25%
Lead Dislodgement	5	1.57%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	1	0.31%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	1	0.31%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	4	1.25%
Total	5	1.57%



YEAR	1	2	3	4	5	6	7	AT 90 MONTHS
SURVIVAL PROBABILITY	98.03%	96.84%	95.35%	95.35%	95.35%	95.35%	95.35%	95.35%
± 1 STANDARD ERROR	0.71%	1.04%	1.34%	1.34%	1.34%	1.34%	1.34%	1.34%
SAMPLE SIZE	290	240	180	140	110	90	70	50

Left-Heart Leads

CUSTOMER REPORTED PERFORMANCE DATA

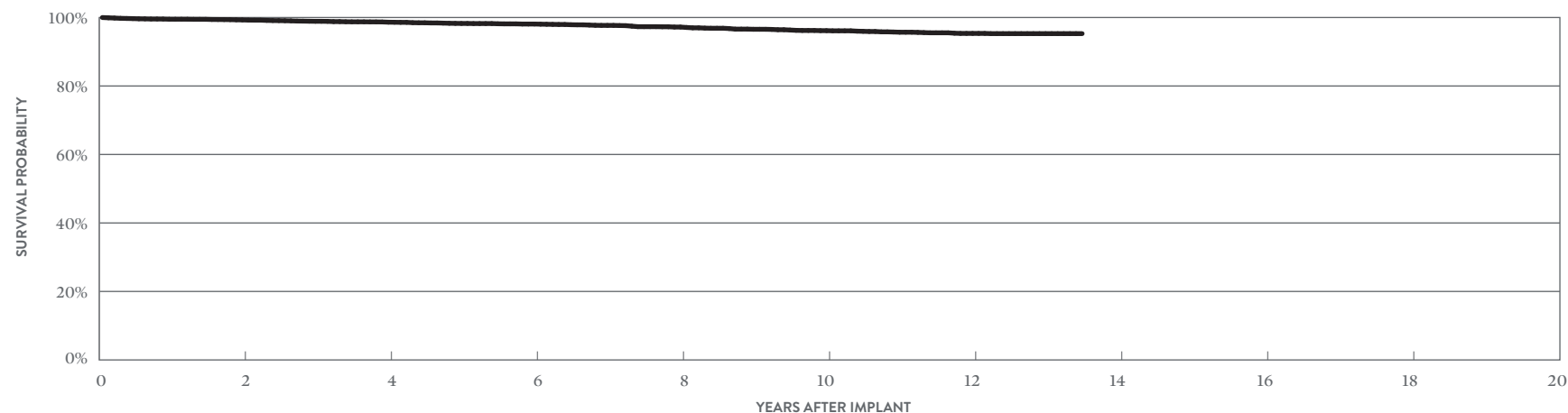
QuickSite™

MODEL 1056K

US Regulatory Approval	June 2004
Registered US Implants	7,874
Estimated Active US Implants	1,990
Insulation	Polyurethane/Silicone
Type and/or Fixation	S-Curve
Polarity	Unipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	0	0.00%
Conductor Fracture	0	0.00%	8	0.10%
Lead Dislodgement	10	0.13%	36	0.46%
Failure to Capture	3	0.04%	74	0.94%
Oversensing	0	0.00%	2	0.03%
Failure to Sense	0	0.00%	0	0.00%
Insulation Breach	0	0.00%	5	0.06%
Abnormal Pacing Impedance	0	0.00%	7	0.09%
Extracardiac Stimulation	10	0.13%	32	0.41%
Other	2	0.03%	10	0.13%
Total	25	0.32%	174	2.21%
Total Returned for Analysis	13		49	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	3	0.04%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	3	0.04%
Insulation Breach	2	0.03%
Lead-to-Can Contact	1	0.01%
Lead-to-Lead Contact	1	0.01%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	51	0.65%
Total	56	0.71%



YEAR	2	4	6	8	10	12	AT 162 MONTHS
SURVIVAL PROBABILITY	99.29%	98.63%	98.09%	97.21%	96.16%	95.35%	95.29%
± 1 STANDARD ERROR	0.10%	0.15%	0.19%	0.26%	0.33%	0.39%	0.39%
SAMPLE SIZE	6,220	4,660	3,440	2,650	2,140	1,650	260

Left-Heart Leads

Survival Probability Summary

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
1458QL	Quartet™	99.53%									
1457Q	QuickFlex™ μ	96.69%									
1456Q	QuickFlex™ μ	99.25%									
1458Q	Quartet™	99.44%	99.21%	99.04%	98.83%	98.62%	98.57%				
1258T	QuickFlex™ μ	99.56%	99.38%	99.23%	99.07%	98.90%	98.67%	98.44%			
1156T	QuickFlex™	99.65%	99.45%	99.15%	98.77%	98.28%	97.74%	97.34%	97.08%	96.73%	
1158T	QuickFlex™ XL	99.55%	99.36%	98.96%	98.55%	97.96%	97.52%	97.05%	96.82%	96.39%	
1058T	QuickSite™ XL	99.75%	99.62%	99.40%	99.15%	98.89%	98.27%	97.74%	97.24%	96.84%	96.65%
1056T	QuickSite™	99.59%	99.39%	99.18%	98.88%	98.47%	97.89%	97.30%	96.84%	96.44%	96.11%
1056K	QuickSite™	99.50%	99.29%	98.89%	98.63%	98.26%	98.09%	97.69%	97.21%	96.56%	96.16%

Left-Heart Leads

Acute Observation Summary

POST IMPLANT ≤30 DAYS

MODELS	US REGULATORY APPROVAL	REGISTERED US IMPLANTS	ESTIMATED ACTIVE US IMPLANTS	CARDIAC PERFORATION		CONDUCTOR FRACTURE		LEAD DISLODGEMENT		FAILURE TO CAPTURE		OVERSENSING		FAILURE TO SENSE		INSULATION BREACH		ABNORMAL PACING IMPEDANCE		EXTRACARDIAC STIMULATION		OTHER		TOTAL		TOTAL RETURNED FOR ANALYSIS
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	
1458QL	Oct-15	4,706	4,224	0	0.00%	0	0.00%	8	0.17%	8	0.17%	0	0.00%	0	0.00%	0	0.00%	1	0.02%	9	0.19%	2	0.04%	28	0.59%	4
1457Q	Oct-15	1,430	1,270	0	0.00%	0	0.00%	8	0.56%	0	0.00%	1	0.07%	0	0.00%	0	0.00%	0	0.00%	11	0.77%	2	0.14%	22	1.54%	2
1456Q	Oct-15	3,358	2,995	1	0.03%	2	0.06%	7	0.21%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	8	0.24%	3	0.09%	21	0.63%	3
1458Q	Nov-11	122,300	87,211	3	<0.01%	0	0.00%	170	0.14%	72	0.06%	2	<0.01%	0	0.00%	1	<0.01%	4	<0.01%	88	0.07%	75	0.06%	415	0.34%	153
1258T	May-10	46,740	26,698	0	0.00%	0	0.00%	45	0.10%	17	0.04%	0	0.00%	1	<0.01%	0	0.00%	5	0.01%	19	0.04%	12	0.03%	99	0.21%	53
1156T	Jul-07	27,662	11,431	0	0.00%	0	0.00%	11	0.04%	4	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	13	0.05%	9	0.03%	37	0.13%	14
1158T	Jul-07	15,337	6,405	0	0.00%	0	0.00%	9	0.06%	2	0.01%	0	0.00%	0	0.00%	0	0.00%	2	0.01%	6	0.04%	6	0.04%	25	0.16%	13
1058T	Feb-06	9,951	3,399	0	0.00%	0	0.00%	10	0.10%	3	0.03%	1	0.01%	0	0.00%	0	0.00%	2	0.02%	9	0.09%	1	0.01%	26	0.26%	11
1056T	Apr-05	32,332	9,782	0	0.00%	0	0.00%	32	0.10%	15	0.05%	2	<0.01%	0	0.00%	1	<0.01%	3	<0.01%	22	0.07%	9	0.03%	84	0.26%	28
1056K	Jun-04	7,874	1,990	0	0.00%	0	0.00%	10	0.13%	3	0.04%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	10	0.13%	2	0.03%	25	0.32%	13

Chronic Complication Summary

>30 DAYS

MODELS	US REGULATORY APPROVAL	REGISTERED US IMPLANTS	ESTIMATED ACTIVE US IMPLANTS	CARDIAC PERFORATION		CONDUCTOR FRACTURE		LEAD DISLODGEMENT		FAILURE TO CAPTURE		OVERSENSING		FAILURE TO SENSE		INSULATION BREACH		ABNORMAL PACING IMPEDANCE		EXTRACARDIAC STIMULATION		OTHER		TOTAL		TOTAL RETURNED FOR ANALYSIS
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	
1458QL	Oct-15	4,706	4,224	0	0.00%	0	0.00%	13	0.28%	5	0.11%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	11	0.23%	0	0.00%	29	0.62%	11
1457Q	Oct-15	1,430	1,270	0	0.00%	0	0.00%	12	0.84%	2	0.14%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	5	0.35%	2	0.14%	21	1.47%	8
1456Q	Oct-15	3,358	2,995	0	0.00%	0	0.00%	14	0.42%	5	0.15%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	0.12%	1	0.03%	24	0.71%	12
1458Q	Nov-11	122,300	87,211	2	<0.01%	13	0.01%	676	0.55%	280	0.23%	8	<0.01%	0	0.00%	6	<0.01%	46	0.04%	148	0.12%	28	0.02%	1207	0.99%	496
1258T	May-10	46,740	26,698	1	<0.01%	20	0.04%	191	0.41%	158	0.34%	15	0.03%	2	<0.01%	7	0.01%	41	0.09%	68	0.15%	9	0.02%	512	1.10%	197
1156T	Jul-07	27,662	11,431	1	<0.01%	5	0.02%	133	0.48%	183	0.66%	14	0.05%	0	0.00%	46	0.17%	61	0.22%	85	0.31%	8	0.03%	536	1.94%	156
1158T	Jul-07	15,337	6,405	1	<0.01%	4	0.03%	95	0.62%	126	0.82%	3	0.02%	1	<0.01%	35	0.23%	23	0.15%	32	0.21%	8	0.05%	328	2.14%	112
1058T	Feb-06	9,951	3,399	0	0.00%	5	0.05%	29	0.29%	81	0.81%	2	0.02%	2	0.02%	32	0.32%	19	0.19%	23	0.23%	2	0.02%	195	1.96%	36
1056T	Apr-05	32,332	9,782	0	0.00%	9	0.03%	166	0.51%	270	0.84%	21	0.06%	1	<0.01%	105	0.32%	59	0.18%	103	0.32%	20	0.06%	754	2.33%	196
1056K	Jun-04	7,874	1,990	0	0.00%	8	0.10%	36	0.46%	74	0.94%	2	0.03%	0	0.00%	5	0.06%	7	0.09%	32	0.41%	10	0.13%	174	2.21%	49

Definitions of observations and complications can be found on [page 7](#).

Left-Heart Leads

US Malfunction Summary

MODELS	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
1458QL	4,706	4.70%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	10	0.21%	10	0.21%
1457Q	1,430	6.80%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	13	0.91%	13	0.91%
1456Q	3,358	7.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	12	0.36%	12	0.36%
1458Q	122,300	6.00%	7	<0.01%	3	<0.01%	0	0.00%	9	<0.01%	475	0.39%	494	0.40%
1258T	46,740	10.50%	5	0.01%	3	<0.01%	0	0.00%	1	<0.01%	213	0.46%	222	0.47%
1156T	27,662	8.80%	6	0.02%	81	0.29%	0	0.00%	0	0.00%	128	0.46%	215	0.78%
1158T	15,337	9.90%	5	0.03%	53	0.35%	1	<0.01%	0	0.00%	86	0.56%	145	0.95%
1058T	9,951	9.70%	2	0.02%	24	0.24%	0	0.00%	1	0.01%	29	0.29%	56	0.56%
1056T	32,332	9.50%	6	0.02%	88	0.27%	0	0.00%	1	<0.01%	158	0.49%	253	0.78%
1056K	7,874	15.20%	3	0.04%	2	0.03%	0	0.00%	0	0.00%	51	0.65%	56	0.71%

Definitions of malfunction categories can be found on [pages 8-9](#).

Left-Heart Leads

Worldwide Malfunction Summary

MODELS	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
1458QL	9,038	2.4%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	10	0.11%	10	0.11%
1457Q	5,731	1.7%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	13	0.23%	13	0.23%
1456Q	8,409	3.0%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	17	0.20%	17	0.20%
1458Q	254,252	3.3%	21	0.01%	7	<0.01%	0	0.00%	18	0.01%	721	0.28%	767	0.30%
1258T	161,854	3.8%	38	0.02%	9	0.01%	0	0.00%	5	<0.01%	372	0.23%	424	0.26%

Definitions of malfunction categories can be found on [pages 8-9](#).

Left-Heart Leads

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	ABNORMAL PACING IMPEDANCE		CARDIAC PERFORATION		CONDUCTOR FRACTURE		EXTRACARDIAC STIMULATION		FAILURE TO CAPTURE		FAILURE TO SENSE		INSULATION BREACH		LEAD DISLODGE MENT		OVERSENSING		PERICARDIAL EFFUSION		SKIN EROSION		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
1458Q	2,108	1,044	79,263	1	0.05%	0	0.00%	0	0.00%	3	0.14%	5	0.24%	0	0.00%	0	0.00%	38	1.80%	1	0.05%	0	0.00%	0	0.00%	0	0.00%	48	2.28%
1258T	2,367	1,015	107,780	6	0.25%	0	0.00%	2	0.08%	56	2.37%	49	2.07%	0	0.00%	0	0.00%	52	2.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	165	6.97%
1156T	985	264	47,545	1	0.10%	0	0.00%	0	0.00%	17	1.73%	9	0.91%	0	0.00%	1	0.10%	27	2.74%	0	0.00%	0	0.00%	1	0.10%	56	5.69%		
1158T	553	120	24,555	0	0.00%	0	0.00%	0	0.00%	9	1.63%	8	1.45%	0	0.00%	1	0.18%	6	1.08%	0	0.00%	0	0.00%	1	0.18%	25	4.52%		
1058T	111	19	5,486	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	3.60%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	3.60%		
1056T	319	66	14,237	1	0.31%	0	0.00%	0	0.00%	2	0.63%	4	1.25%	0	0.00%	0	0.00%	5	1.57%	0	0.00%	0	0.00%	0	0.00%	12	3.76%		

MALFUNCTIONS

MODELS	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
1458Q	2,108	4.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	21	1.00%	21	1.00%
1258T	2,367	5.80%	1	0.04%	0	0.00%	0	0.00%	0	0.00%	35	1.48%	36	1.52%
1156T	985	8.50%	0	0.00%	3	0.30%	0	0.00%	0	0.00%	19	1.93%	22	2.23%
1158T	553	5.20%	0	0.00%	1	0.18%	0	0.00%	0	0.00%	7	1.27%	8	1.45%
1058T	111	6.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
1056T	319	7.20%	0	0.00%	1	0.31%	0	0.00%	0	0.00%	4	1.25%	5	1.57%

A list of complications can be found on [page 12](#).

Definitions of malfunction categories can be found on [pages 8-9](#).

DUAL-CHAMBER
Implantable Cardioverter
Defibrillator (ICD) Devices

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

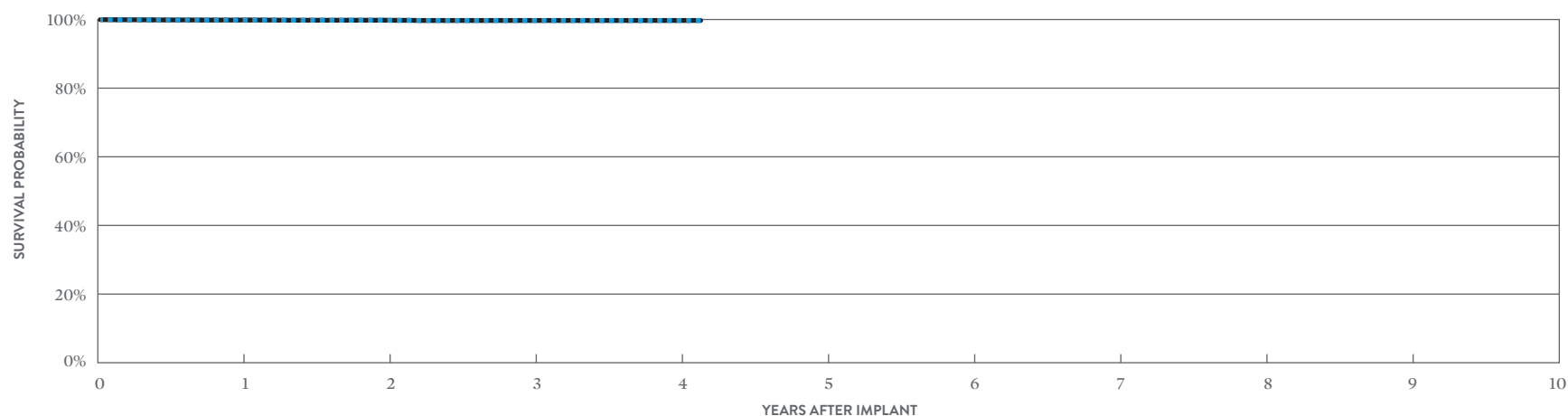
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ DR

MODEL CD2411-36Q*

US Regulatory Approval	June 2013
Registered US Implants	15,960
Estimated Active US Implants	12,580
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	0
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	2	0.01%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	2	0.01%	1	<0.01%
Software/Firmware	1	<0.01%	0	0.00%
Mechanical	1	<0.01%	2	0.01%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	1	<0.01%	2	0.01%
Total	6	0.04%	8	0.05%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.84%	99.79%	99.72%	99.72%	99.72%
± 1 STANDARD ERROR	0.03%	0.04%	0.05%	0.05%	0.05%
SAMPLE SIZE	12,890	7,900	4,330	1,500	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.84%	99.79%	99.72%	99.72%	99.72%
± 1 STANDARD ERROR	0.03%	0.04%	0.05%	0.05%	0.05%

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

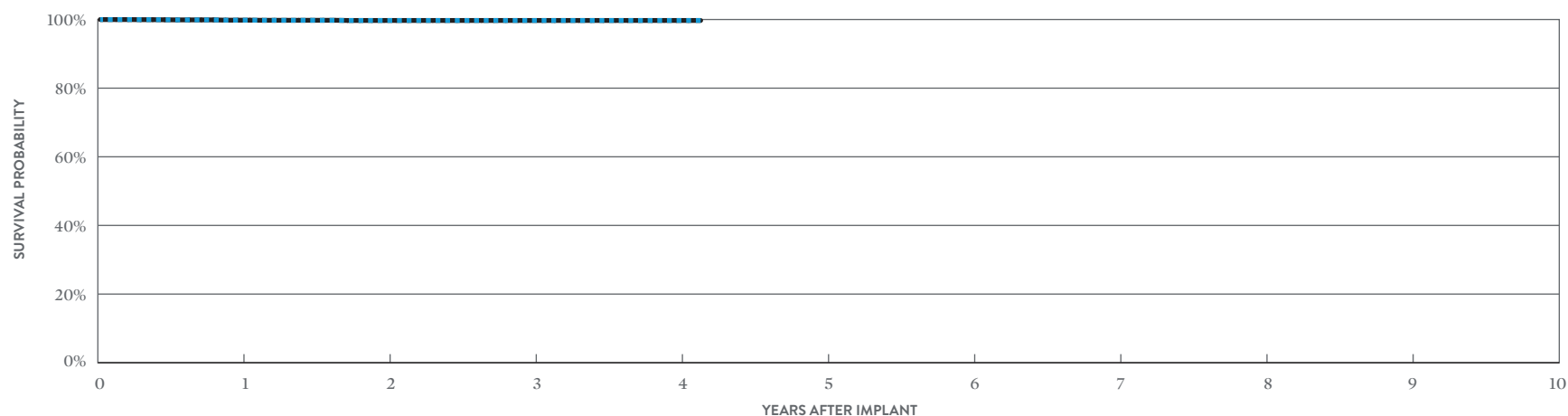
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ DR

MODEL CD2411-36C*

US Regulatory Approval	June 2013
Registered US Implants	8,338
Estimated Active US Implants	6,431
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	0
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.02%	1	0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	4	0.05%	1	0.01%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	6	0.07%	2	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.81%	99.72%	99.72%	99.72%	99.72%
± 1 STANDARD ERROR	0.05%	0.07%	0.07%	0.07%	0.07%
SAMPLE SIZE	7,050	4,620	2,630	1,030	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.81%	99.72%	99.72%	99.72%	99.72%
± 1 STANDARD ERROR	0.05%	0.07%	0.07%	0.07%	0.07%

*Parylene coating.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

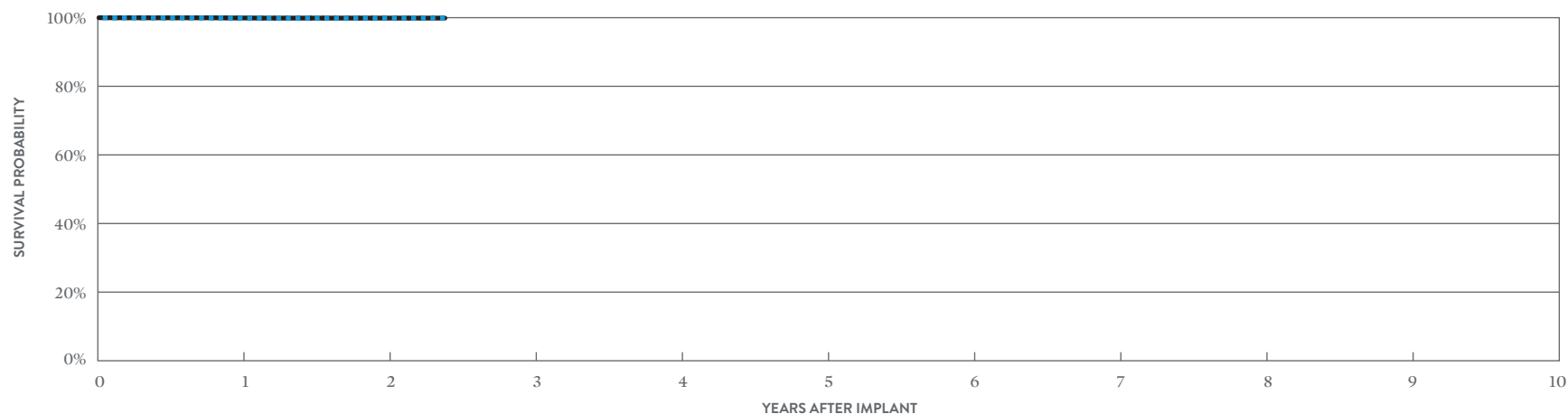
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ DR

MODEL CD2357-40Q* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	14,890
Estimated Active US Implants	12,814
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	2
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	<0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	<0.01%	0	0.00%
High Voltage Capacitor	2	0.01%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	2	0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	3	0.02%	3	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	99.86%	99.83%	99.83%
± 1 STANDARD ERROR	0.03%	0.04%	0.04%
SAMPLE SIZE	11,160	4,540	360

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	99.91%	99.88%	99.88%
± 1 STANDARD ERROR	0.02%	0.04%	0.04%

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

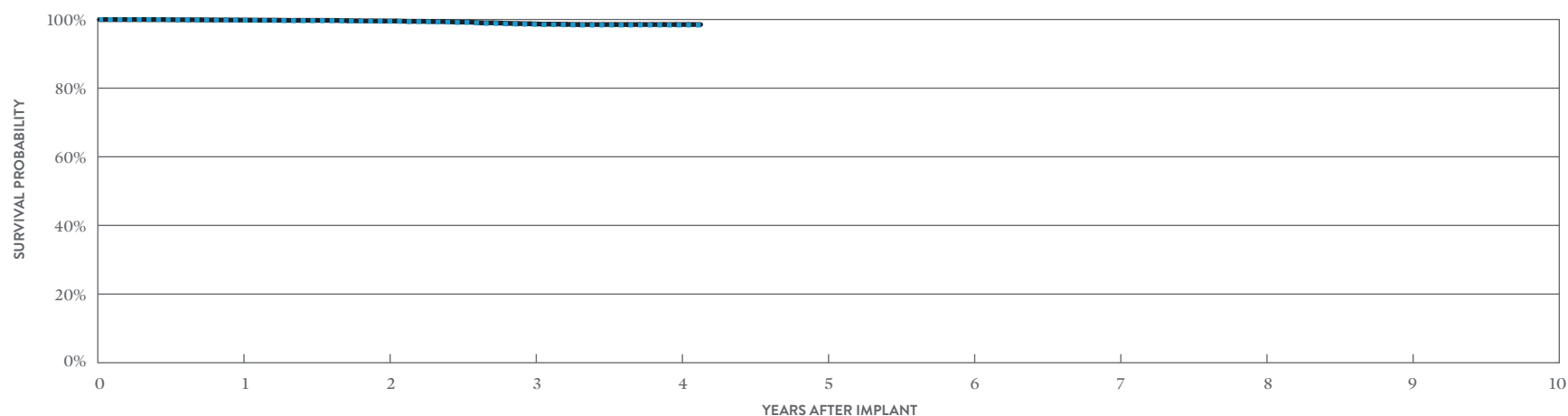
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ DR

MODEL CD2357-40Q* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	12,264
Estimated Active US Implants	8,223
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	5
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.02%	4	0.03%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	0	0.00%	1	<0.01%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	24	0.20%	8	0.07%
Other	0	0.00%	3	0.02%
Total	28	0.23%	18	0.15%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.79%	99.47%	98.60%	98.37%	98.37%
± 1 STANDARD ERROR	0.04%	0.07%	0.13%	0.15%	0.15%
SAMPLE SIZE	11,500	9,920	6,820	2,500	300

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.84%	99.56%	98.76%	98.53%	98.53%
± 1 STANDARD ERROR	0.04%	0.06%	0.12%	0.15%	0.15%

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

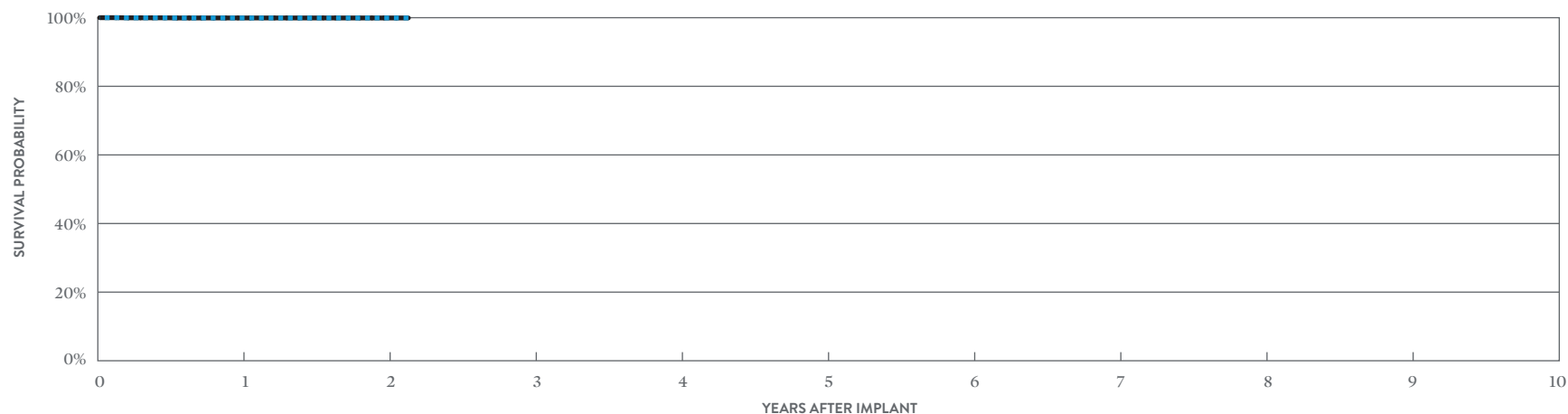
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ DR

MODEL CD2357-40C* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	6,180
Estimated Active US Implants	5,357
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	0
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	0.02%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.02%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	1	0.02%	1	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 26 MONTHS
SURVIVAL PROBABILITY	99.92%	99.92%	99.92%
± 1 STANDARD ERROR	0.04%	0.04%	0.04%
SAMPLE SIZE	4,690	1,800	300

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 26 MONTHS
SURVIVAL PROBABILITY	99.92%	99.92%	99.92%
± 1 STANDARD ERROR	0.04%	0.04%	0.04%

*Parylene coating.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

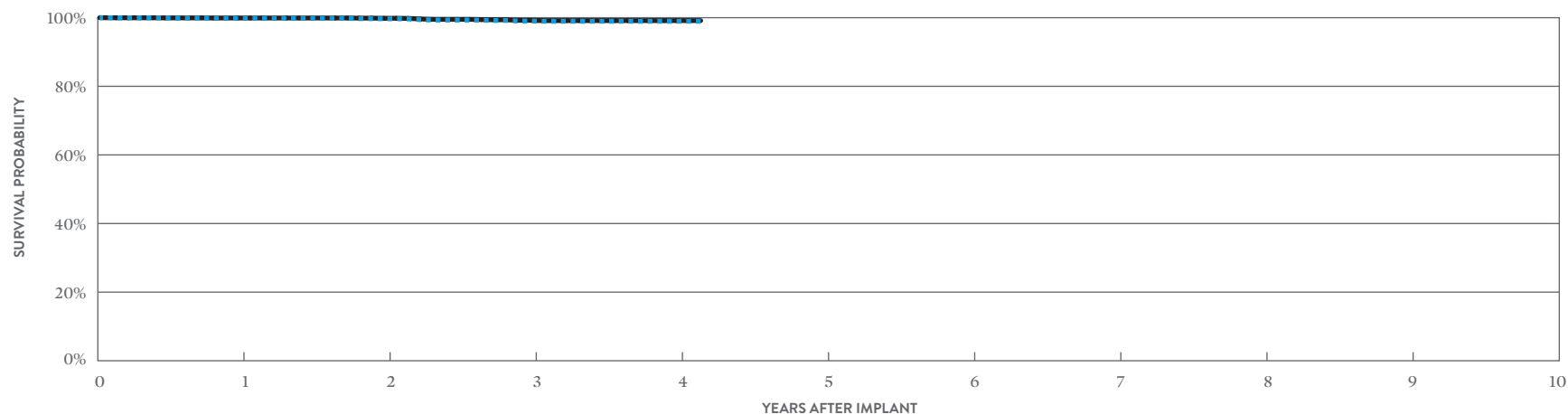
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ DR

MODEL CD2357-40C* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	6,956
Estimated Active US Implants	4,615
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	5
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.04%	0	0.00%
Electrical Interconnect	0	0.00%	1	0.01%
Battery	0	0.00%	2	0.03%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	9	0.13%	1	0.01%
Other	0	0.00%	0	0.00%
Total	12	0.17%	4	0.06%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.84%	99.67%	99.03%	98.95%	98.95%
± 1 STANDARD ERROR	0.05%	0.07%	0.15%	0.16%	0.16%
SAMPLE SIZE	6,520	5,550	3,820	1,490	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.91%	99.83%	99.23%	99.15%	99.15%
± 1 STANDARD ERROR	0.04%	0.05%	0.14%	0.15%	0.15%

*Parylene coating.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

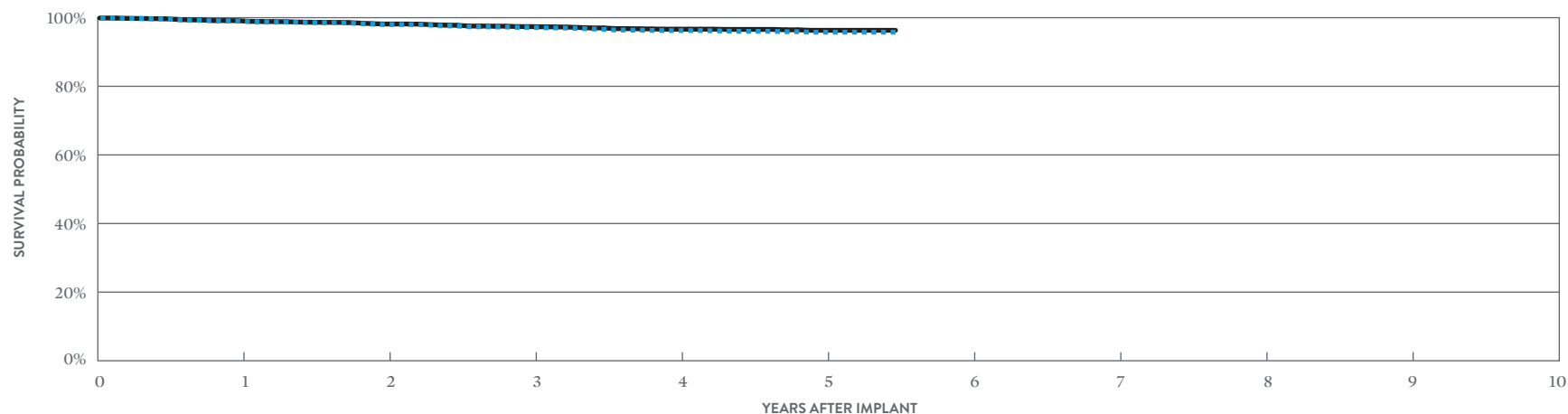
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ DR

MODEL CD2311-36Q*

US Regulatory Approval	May 2012
Registered US Implants	5,897
Estimated Active US Implants	3,525
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	9
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.05%	4	0.07%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	48	0.81%	5	0.08%
Software/Firmware	1	0.02%	0	0.00%
Mechanical	2	0.03%	3	0.05%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	2	0.03%	2	0.03%
Total	56	0.95%	14	0.24%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 66 MONTHS
SURVIVAL PROBABILITY	99.04%	98.00%	97.19%	96.25%	95.77%	95.77%
± 1 STANDARD ERROR	0.13%	0.19%	0.23%	0.28%	0.32%	0.32%
SAMPLE SIZE	5,530	4,900	4,390	3,710	2,140	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 66 MONTHS
SURVIVAL PROBABILITY	99.13%	98.16%	97.40%	96.65%	96.32%	96.32%
± 1 STANDARD ERROR	0.12%	0.18%	0.23%	0.26%	0.30%	0.30%

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

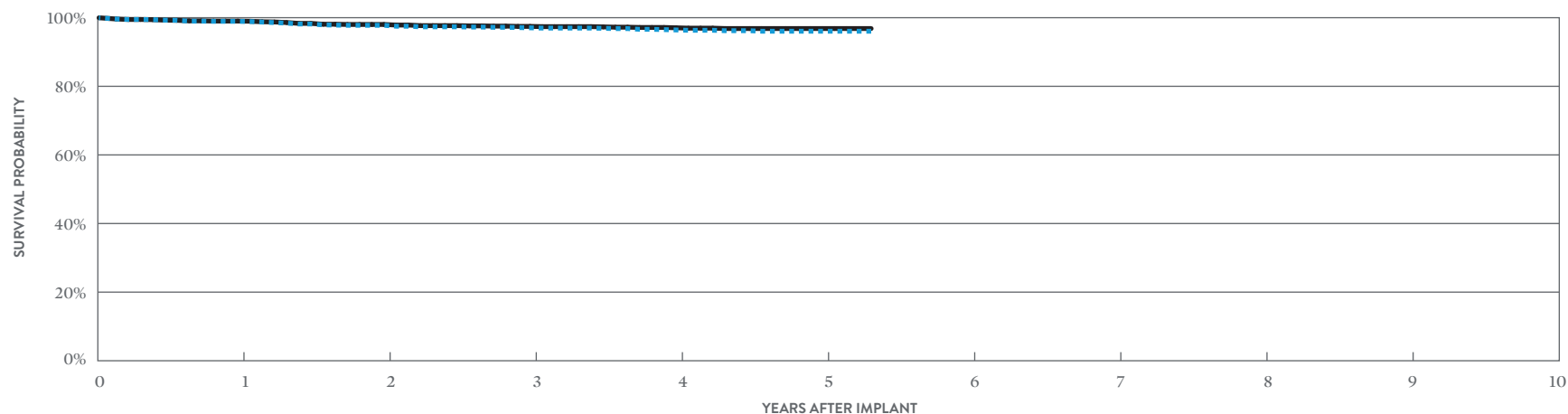
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ DR

MODEL CD2311-36

US Regulatory Approval	May 2012
Registered US Implants	3,746
Estimated Active US Implants	2,248
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	8
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	5	0.13%	2	0.05%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	20	0.53%	5	0.13%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	4	0.11%	3	0.08%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	3	0.08%	0	0.00%
Total	32	0.85%	10	0.27%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 64 MONTHS
SURVIVAL PROBABILITY	98.94%	97.78%	97.05%	96.42%	96.05%	96.05%
± 1 STANDARD ERROR	0.17%	0.25%	0.29%	0.33%	0.37%	0.37%
SAMPLE SIZE	3,520	3,100	2,770	2,310	1,290	280

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 64 MONTHS
SURVIVAL PROBABILITY	99.02%	98.02%	97.46%	97.05%	96.84%	96.84%
± 1 STANDARD ERROR	0.16%	0.24%	0.28%	0.30%	0.32%	0.32%

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

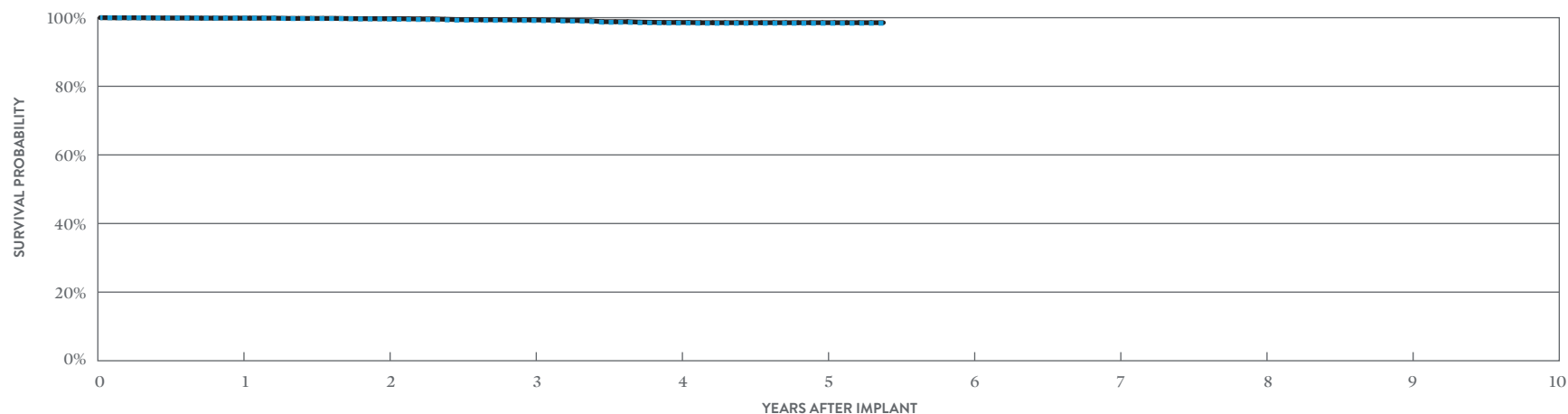
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ DR

MODEL CD2257-40Q* (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	6,798
Estimated Active US Implants	3,935
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	4
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	5	0.07%	1	0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	0.01%	1	0.01%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	0.01%
Mechanical	0	0.00%	1	0.01%
Possible Early Battery Depletion	12	0.18%	6	0.09%
Other	3	0.04%	0	0.00%
Total	21	0.31%	10	0.15%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 65 MONTHS
SURVIVAL PROBABILITY	99.87%	99.63%	99.18%	98.43%	98.37%	98.37%
± 1 STANDARD ERROR	0.04%	0.08%	0.12%	0.18%	0.18%	0.18%
SAMPLE SIZE	6,370	5,630	5,040	4,190	2,180	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 65 MONTHS
SURVIVAL PROBABILITY	99.87%	99.72%	99.36%	98.61%	98.55%	98.55%
± 1 STANDARD ERROR	0.04%	0.07%	0.11%	0.17%	0.17%	0.17%

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

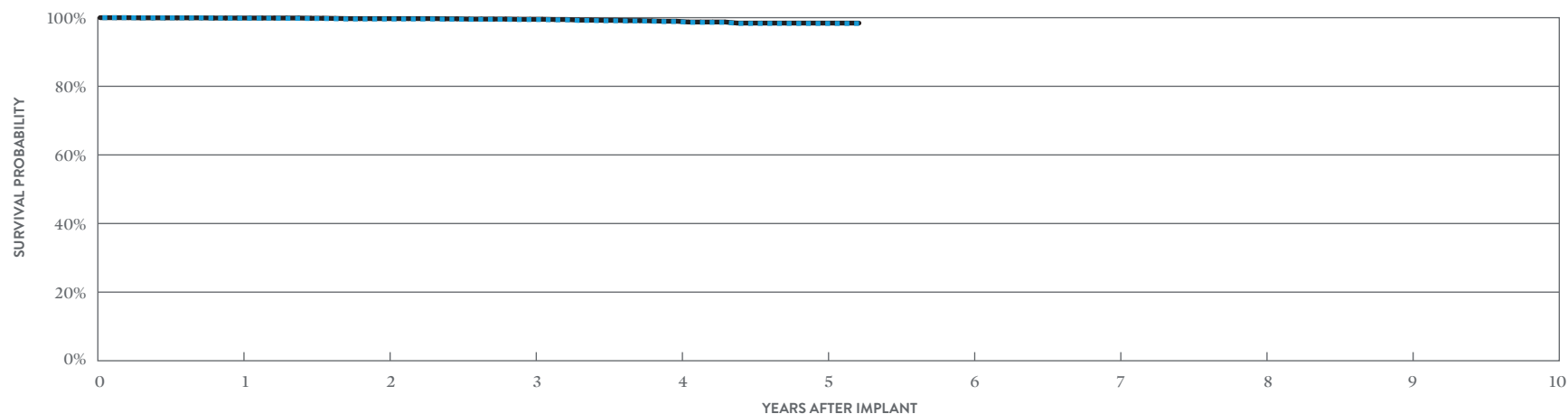
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ DR

MODEL CD2257-40 (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	4,235
Estimated Active US Implants	2,382
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	2
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	0.02%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	2	0.05%
High Voltage Capacitor	1	0.02%	0	0.00%
Software/Firmware	0	0.00%	1	0.02%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	9	0.21%	4	0.09%
Other	0	0.00%	1	0.02%
Total	11	0.26%	8	0.19%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 63 MONTHS
SURVIVAL PROBABILITY	99.85%	99.62%	99.43%	98.87%	98.32%	98.32%
± 1 STANDARD ERROR	0.06%	0.10%	0.13%	0.19%	0.26%	0.26%
SAMPLE SIZE	3,990	3,550	3,140	2,570	1,330	270

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 63 MONTHS
SURVIVAL PROBABILITY	99.90%	99.73%	99.54%	98.97%	98.43%	98.43%
± 1 STANDARD ERROR	0.05%	0.09%	0.12%	0.19%	0.25%	0.25%

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

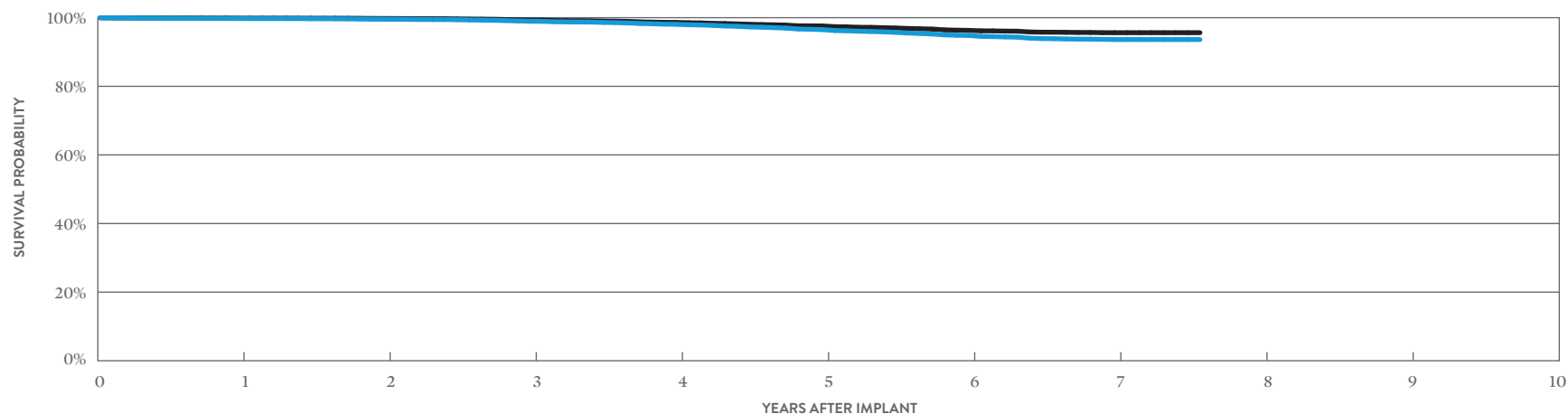
CUSTOMER REPORTED PERFORMANCE DATA

Fortify™ DR

MODEL CD2231-40Q* (ADVISORY POPULATION)

US Regulatory Approval	May 2010
Registered US Implants	26,868
Estimated Active US Implants	12,588
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	114
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	7	0.03%	8	0.03%
Electrical Interconnect	2	<0.01%	2	<0.01%
Battery	27	0.10%	27	0.10%
High Voltage Capacitor	4	0.01%	2	<0.01%
Software/Firmware	1	<0.01%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	120	0.45%	45	0.17%
Other	9	0.03%	5	0.02%
Total	170	0.63%	89	0.33%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 91 MONTHS
SURVIVAL PROBABILITY	99.75%	99.55%	98.97%	98.07%	96.52%	94.81%	93.63%	93.63%
± 1 STANDARD ERROR	0.03%	0.04%	0.07%	0.10%	0.14%	0.18%	0.22%	0.22%
SAMPLE SIZE	25,130	22,180	19,900	17,840	15,580	11,670	5,950	360

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 91 MONTHS
SURVIVAL PROBABILITY	99.87%	99.76%	99.31%	98.62%	97.59%	96.30%	95.63%	95.63%
± 1 STANDARD ERROR	0.02%	0.03%	0.05%	0.08%	0.11%	0.15%	0.18%	0.18%

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

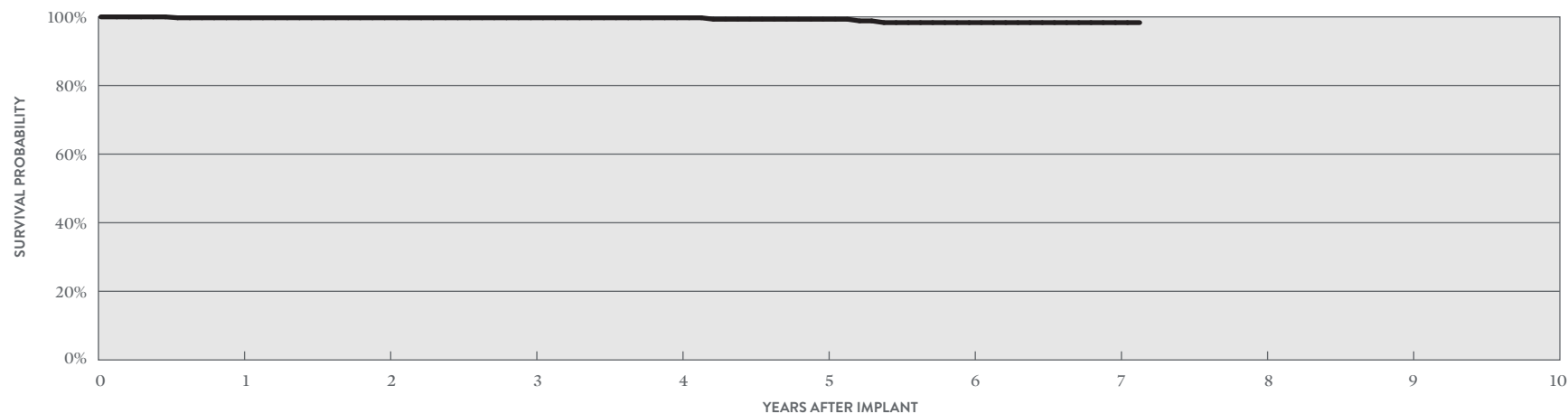
Fortify™ DR

MODEL CD2231-40Q*

US Regulatory Approval	May 2010
Number of Devices Enrolled in Study	390
Active Devices Enrolled in Study	159
Cumulative Months of Follow-up	21,754
Estimated Longevity	(see table on page 120)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Premature Battery Depletion	4	1.03%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	0.26%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	2	0.51%	2	0.51%
Other	1	0.26%	0	0.00%
Total	4	1.03%	2	0.51%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	AT 86 MONTHS
SURVIVAL PROBABILITY	99.74%	99.74%	99.74%	99.74%	99.32%	98.33%	98.33%	98.33%
± 1 STANDARD ERROR	0.26%	0.26%	0.26%	0.26%	0.49%	0.85%	0.85%	0.85%
SAMPLE SIZE	380	340	300	260	230	190	120	50

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

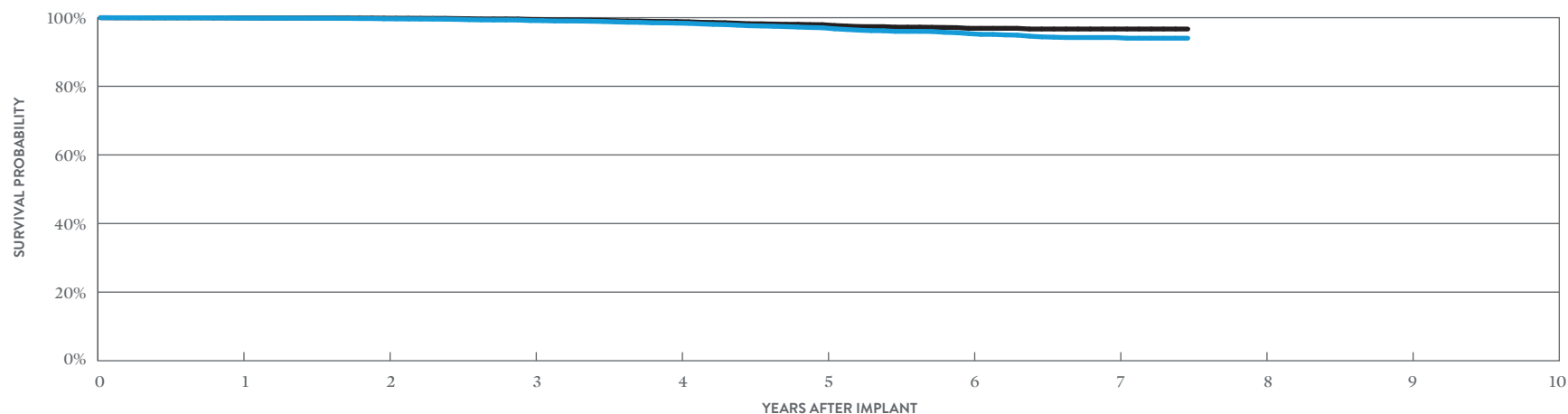
CUSTOMER REPORTED PERFORMANCE DATA

Fortify™ DR

MODEL CD2231-40 (ADVISORY POPULATION)

US Regulatory Approval	May 2010
Registered US Implants	12,094
Estimated Active US Implants	5,540
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	55
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	6	0.05%	2	0.02%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	5	0.04%	5	0.04%
High Voltage Capacitor	6	0.05%	1	<0.01%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	43	0.36%	15	0.12%
Other	4	0.03%	3	0.02%
Total	65	0.54%	27	0.22%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 90 MONTHS
SURVIVAL PROBABILITY	99.88%	99.67%	99.16%	98.42%	97.07%	95.36%	94.22%	94.02%
± 1 STANDARD ERROR	0.02%	0.05%	0.09%	0.13%	0.19%	0.25%	0.33%	0.36%
SAMPLE SIZE	11,320	9,970	8,850	7,860	6,800	4,880	2,320	270

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 90 MONTHS
SURVIVAL PROBABILITY	99.95%	99.86%	99.49%	98.87%	97.96%	96.90%	96.70%	96.70%
± 1 STANDARD ERROR	0.02%	0.03%	0.07%	0.11%	0.16%	0.20%	0.23%	0.23%

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

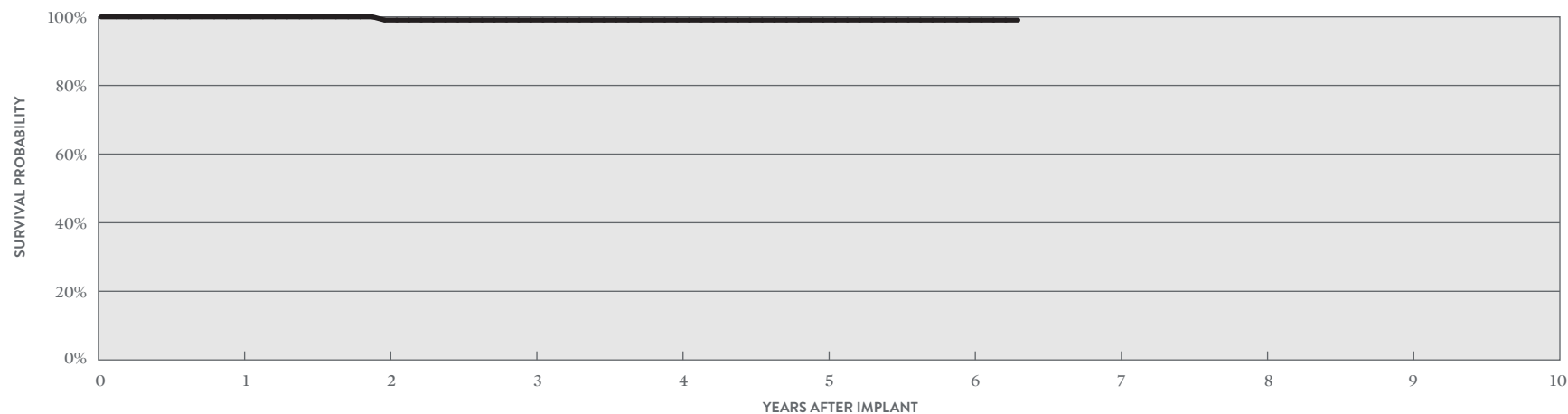
Fortify™ DR

MODEL CD2231-40

US Regulatory Approval	May 2010
Number of Devices Enrolled in Study	175
Active Devices Enrolled in Study	58
Cumulative Months of Follow-up	7,769
Estimated Longevity	(see table on page 120)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Premature Battery Depletion	1	0.57%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	AT 76 MONTHS
SURVIVAL PROBABILITY	100.00%	99.08%	99.08%	99.08%	99.08%	99.08%	99.08%
± 1 STANDARD ERROR	0.00%	0.00%	0.91%	0.91%	0.91%	0.91%	0.91%
SAMPLE SIZE	160	130	100	80	70	70	50

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

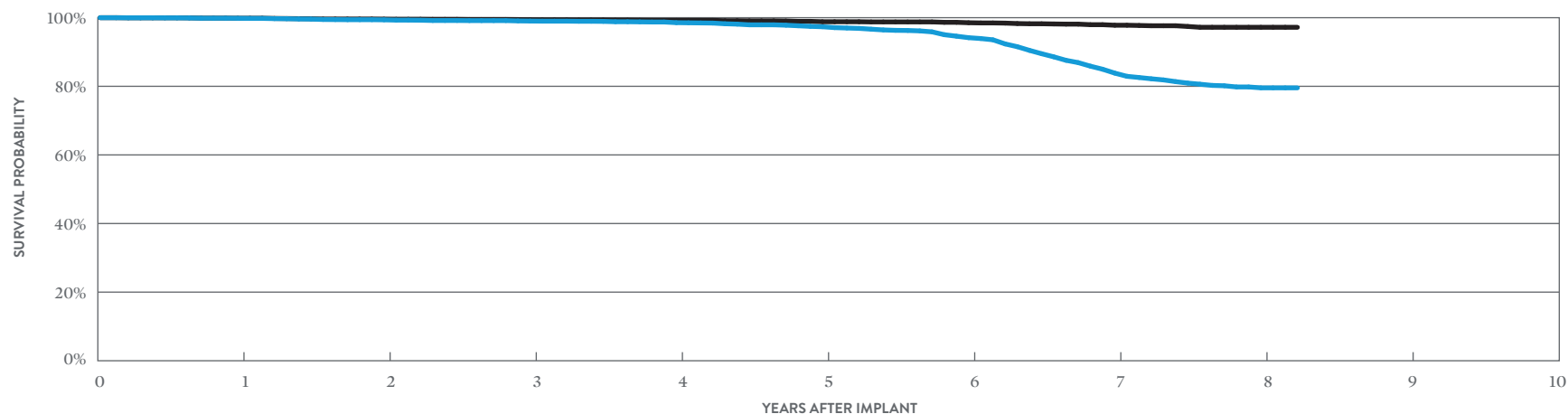
CUSTOMER REPORTED PERFORMANCE DATA

Current™ + DR

MODEL CD2211-36Q*

US Regulatory Approval	February 2009
Registered US Implants	8,144
Estimated Active US Implants	2,132
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	269
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	6	0.07%	4	0.05%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	6	0.07%	6	0.07%
High Voltage Capacitor	2	0.02%	0	0.00%
Software/Firmware	1	0.01%	11	0.14%
Mechanical	0	0.00%	2	0.02%
Possible Early Battery Depletion	4	0.05%	3	0.04%
Other	5	0.06%	3	0.04%
Total	24	0.29%	29	0.36%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 99 MONTHS
SURVIVAL PROBABILITY	99.82%	99.37%	99.02%	98.55%	97.36%	94.17%	83.84%	79.55%	79.55%
± 1 STANDARD ERROR	0.04%	0.09%	0.12%	0.14%	0.21%	0.33%	0.56%	0.67%	0.69%
SAMPLE SIZE	7,570	6,620	5,940	5,310	4,740	4,180	3,400	1,730	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 99 MONTHS
SURVIVAL PROBABILITY	99.85%	99.58%	99.41%	99.22%	98.82%	98.53%	97.80%	97.20%	97.20%
± 1 STANDARD ERROR	0.04%	0.07%	0.09%	0.11%	0.14%	0.16%	0.21%	0.27%	0.27%

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

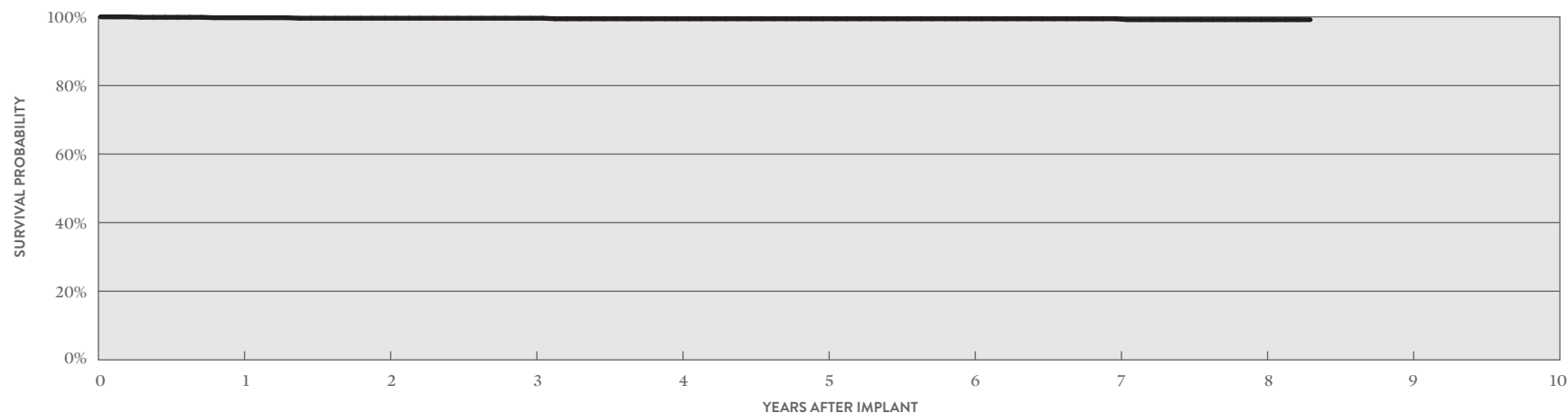
Current™ + DR

MODEL CD2211-36Q*

US Regulatory Approval	February 2009
Number of Devices Enrolled in Study	835
Active Devices Enrolled in Study	360
Cumulative Months of Follow-up	52,193
Estimated Longevity	(see table on page 120)
Max. Delivered Energy	36 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Premature Battery Depletion	4	0.48%
Skin Erosion	1	0.12%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.12%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	0.12%	2	0.24%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	2	0.24%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	1	0.12%
Other	0	0.00%	1	0.12%
Total	1	0.12%	7	0.84%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	8	AT 100 MONTHS
SURVIVAL PROBABILITY	99.75%	99.61%	99.61%	99.44%	99.44%	99.44%	99.44%	99.18%	99.18%
± 1 STANDARD ERROR	0.18%	0.23%	0.23%	0.28%	0.28%	0.28%	0.28%	0.39%	0.39%
SAMPLE SIZE	790	710	640	570	500	440	390	260	60

*DF4-LLHH connector type.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

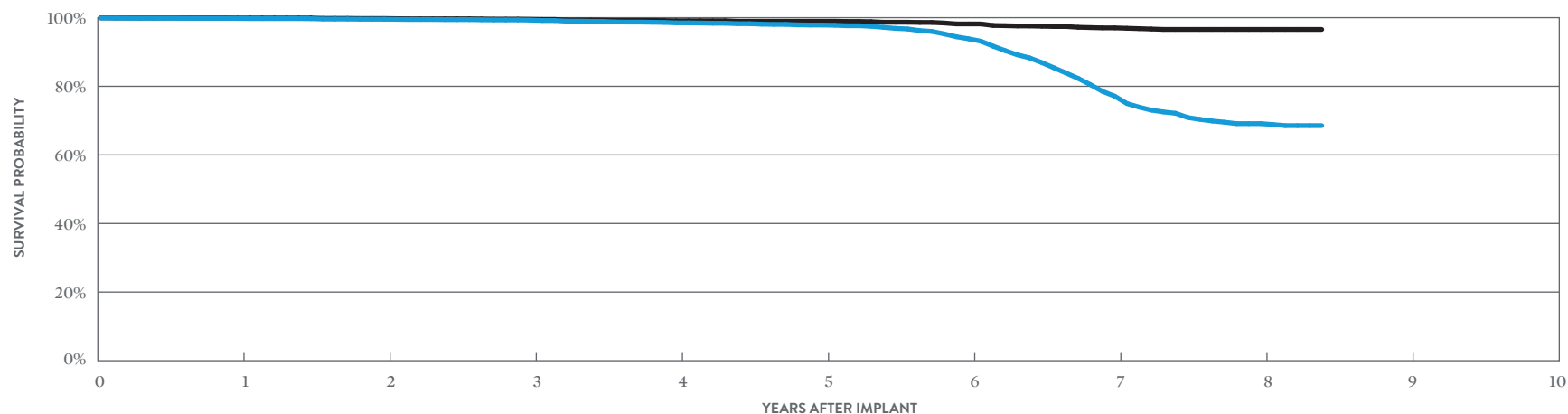
CUSTOMER REPORTED PERFORMANCE DATA

Current™ + DR

MODEL CD2211-36

US Regulatory Approval	February 2009
Registered US Implants	6,271
Estimated Active US Implants	1,516
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	296
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.03%	1	0.02%
Electrical Interconnect	2	0.03%	0	0.00%
Battery	7	0.11%	4	0.06%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	1	0.02%	13	0.21%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	9	0.14%	4	0.06%
Other	5	0.08%	0	0.00%
Total	26	0.41%	22	0.35%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	99.78%	99.57%	99.30%	98.49%	97.86%	93.84%	77.12%	69.12%	68.54%
± 1 STANDARD ERROR	0.05%	0.09%	0.11%	0.17%	0.22%	0.39%	0.77%	0.92%	0.96%
SAMPLE SIZE	5,840	5,090	4,510	4,020	3,560	3,070	2,350	1,230	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	8
SURVIVAL PROBABILITY	99.90%	99.76%	99.54%	99.10%	98.96%	98.18%	97.03%	96.58%	96.58%
± 1 STANDARD ERROR	0.03%	0.07%	0.09%	0.14%	0.15%	0.22%	0.31%	0.34%	0.34%

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

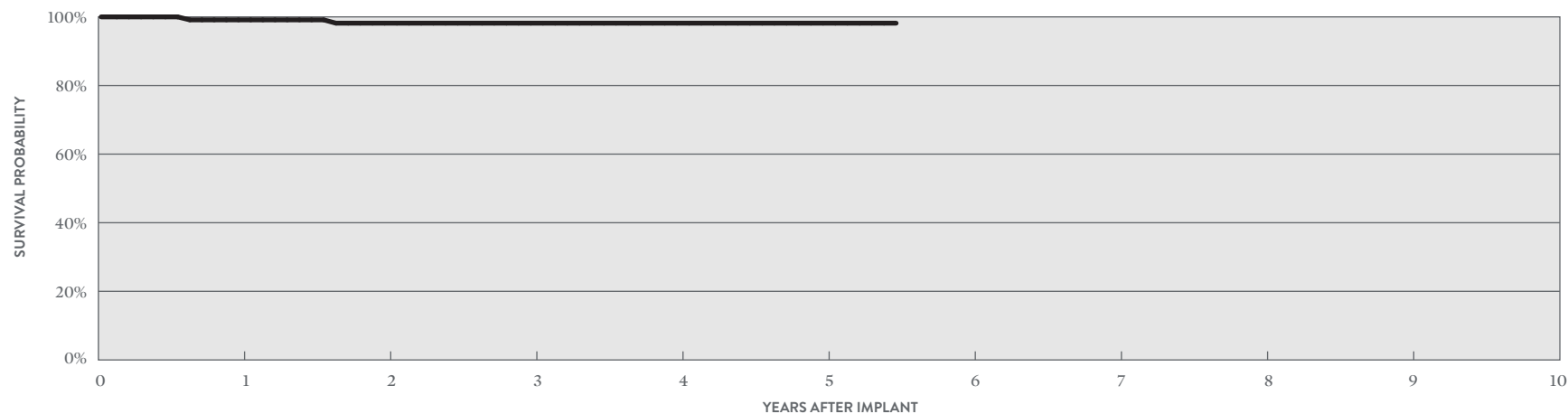
Current™ + DR

MODEL CD2211-36

US Regulatory Approval	February 2009
Number of Devices Enrolled in Study	123
Active Devices Enrolled in Study	26
Cumulative Months of Follow-up	6,344
Estimated Longevity	(see table on page 120)
Max. Delivered Energy	36 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Inappropriate Shock	1	0.81%
Premature Battery Depletion	1	0.81%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	1	0.81%	1	0.81%
Total	1	0.81%	1	0.81%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	5	4	AT 66 MONTHS
SURVIVAL PROBABILITY	99.13%	98.17%	98.17%	98.17%	98.17%	98.17%
± 1 STANDARD ERROR	0.86%	1.29%	1.29%	1.29%	1.29%	1.29%
SAMPLE SIZE	120	100	80	60	60	50

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

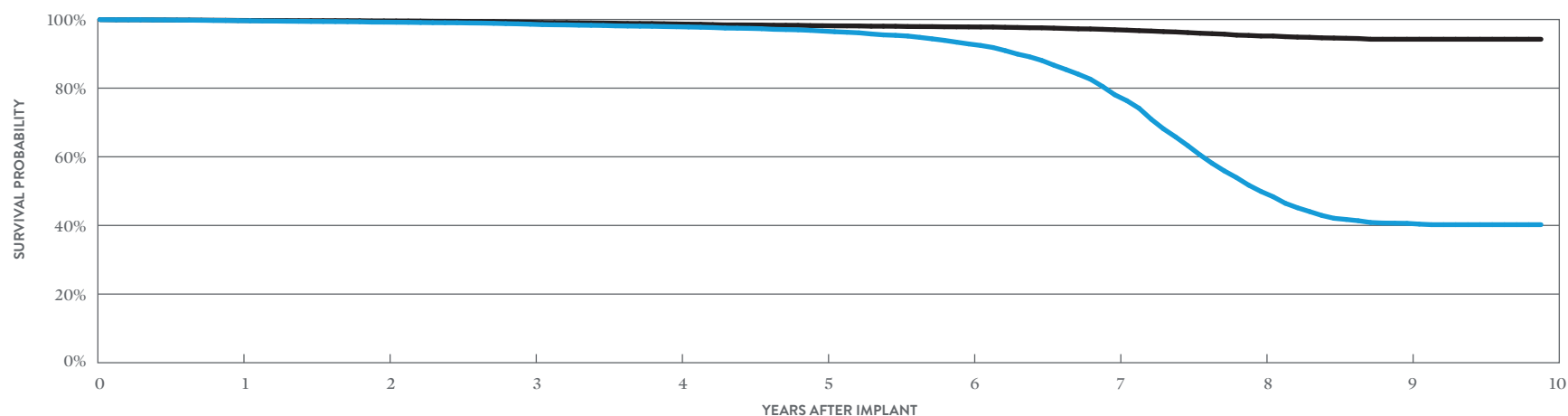
CUSTOMER REPORTED PERFORMANCE DATA

Current™ DR RF

MODEL 2207-36

US Regulatory Approval	September 2007
Registered US Implants	22,386
Estimated Active US Implants	3,307
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	2,169
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	10	0.04%	12	0.05%
Electrical Interconnect	6	0.03%	2	<0.01%
Battery	20	0.09%	9	0.04%
High Voltage Capacitor	1	<0.01%	0	0.00%
Software/Firmware	2	<0.01%	41	0.18%
Mechanical	1	<0.01%	20	0.09%
Possible Early Battery Depletion	40	0.18%	19	0.08%
Other	36	0.16%	6	0.03%
Total	116	0.52%	109	0.49%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.66%	99.23%	98.63%	97.89%	96.64%	92.88%	78.07%	49.91%	40.61%	40.22%
± 1 STANDARD ERROR	0.04%	0.06%	0.08%	0.11%	0.14%	0.22%	0.37%	0.50%	0.52%	0.52%
SAMPLE SIZE	20,830	18,100	15,960	14,200	12,670	11,150	9,220	6,240	2,960	260

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.73%	99.58%	99.20%	98.70%	98.19%	97.84%	97.02%	95.19%	94.25%	94.25%
± 1 STANDARD ERROR	0.03%	0.05%	0.06%	0.09%	0.11%	0.12%	0.15%	0.22%	0.28%	0.28%

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

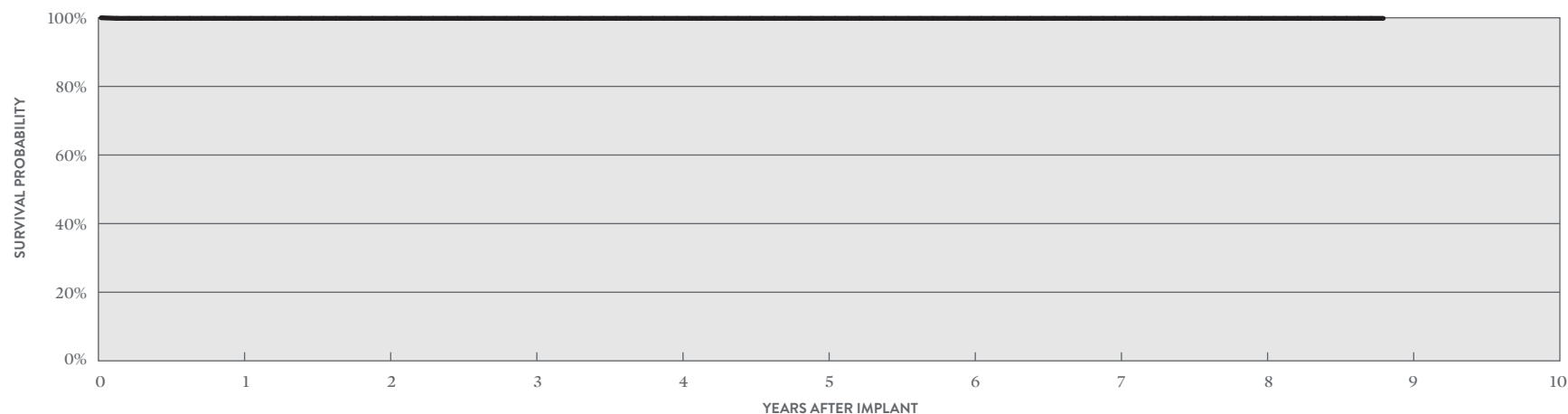
Current™ DR RF

MODEL 2207-36

US Regulatory Approval	September 2007
Number of Devices Enrolled in Study	631
Active Devices Enrolled in Study	73
Cumulative Months of Follow-up	32,853
Estimated Longevity	(see table on page 120)
Max. Delivered Energy	36 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Inappropriate Shock	1	0.16%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	2	0.32%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	1	0.16%	1	0.16%
Other	0	0.00%	0	0.00%
Total	1	0.16%	3	0.48%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	8	AT 106 MONTHS
SURVIVAL PROBABILITY	99.84%	99.84%	99.84%	99.84%	99.84%	99.84%	99.84%	99.84%	99.84%
± 1 STANDARD ERROR	0.16%	0.16%	0.16%	0.16%	0.16%	0.16%	0.16%	0.16%	0.16%
SAMPLE SIZE	600	520	430	340	280	230	180	120	50

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

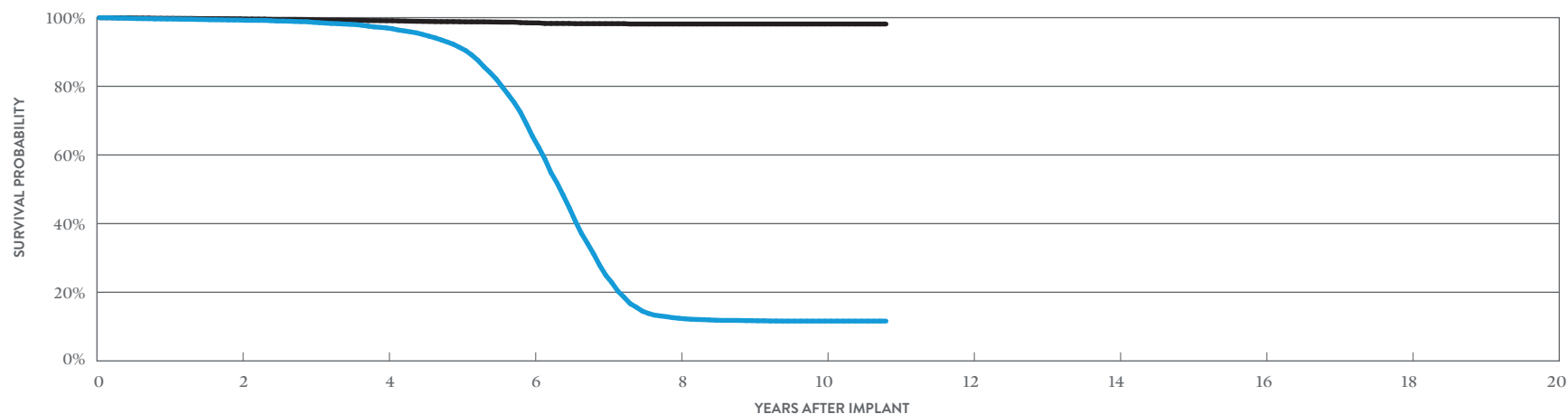
CUSTOMER REPORTED PERFORMANCE DATA

Atlas™ II + DR

MODEL V-268

US Regulatory Approval	July 2006
Registered US Implants	14,809
Estimated Active US Implants	1,283
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	2,920
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 330)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	6	0.04%	4	0.03%
Electrical Interconnect	4	0.03%	0	0.00%
Battery	9	0.06%	3	0.02%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	19	0.13%	6	0.04%
Other	10	0.07%	5	0.03%
Total	48	0.32%	19	0.13%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 130 MONTHS
SURVIVAL PROBABILITY	99.30%	97.01%	65.36%	12.43%	11.61%	11.61%
± 1 STANDARD ERROR	0.07%	0.16%	0.52%	0.34%	0.33%	0.33%
SAMPLE SIZE	11,930	9,120	6,120	1,930	900	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 130 MONTHS
SURVIVAL PROBABILITY	99.67%	99.11%	98.47%	98.15%	98.15%	98.15%
± 1 STANDARD ERROR	0.05%	0.09%	0.13%	0.17%	0.17%	0.17%

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

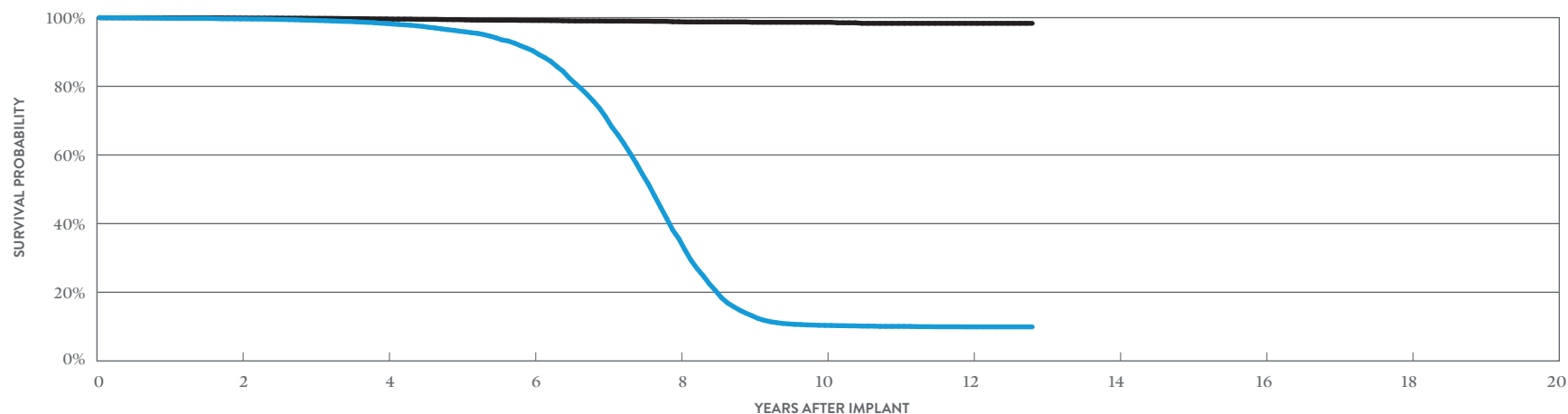
CUSTOMER REPORTED PERFORMANCE DATA

Atlas™ + DR

MODEL V-243

US Regulatory Approval	October 2003
Registered US Implants	21,082
Estimated Active US Implants	1,314
Estimated Longevity	(see table on page 120)
Normal Battery Depletion	3,623
Max. Delivered Energy	36 joules
Number of US Advisories (see pgs. 330, 331, 332)	Three

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	5	0.02%	3	0.01%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	12	0.06%	4	0.02%
High Voltage Capacitor	1	<0.01%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	4	0.02%
Possible Early Battery Depletion	6	0.03%	4	0.02%
Other	17	0.08%	2	<0.01%
Total	42	0.20%	17	0.08%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 154 MONTHS
SURVIVAL PROBABILITY	99.63%	98.31%	90.40%	35.61%	10.35%	9.92%	9.92%
± 1 STANDARD ERROR	0.04%	0.10%	0.27%	0.50%	0.28%	0.28%	0.28%
SAMPLE SIZE	17,170	13,150	9,500	5,090	1,630	780	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 154 MONTHS
SURVIVAL PROBABILITY	99.90%	99.63%	99.18%	98.81%	98.65%	98.35%	98.35%
± 1 STANDARD ERROR	0.02%	0.05%	0.08%	0.12%	0.14%	0.21%	0.21%

BATTERY LONGEVITY SUMMARY

**Dual-Chamber
Implantable Cardioverter
Defibrillator (ICD) Devices**

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Battery Longevity (years)

MODELS	FAMILY	NO PACING	25% PACING	50% PACING	100% PACING
CD2411-36Q	Ellipse™ DR*	10.4	9.6	8.9	7.7
CD2411-36C	Ellipse™ DR*	10.4	9.6	8.9	7.7
CD2357-40Q	Fortify Assura™ DR*	11.1	10.2	9.5	8.3
CD2357-40C	Fortify Assura™ DR*	11.1	10.2	9.5	8.3
CD2311-36Q	Ellipse™ DR*	10.4	9.6	8.9	7.7
CD2311-36	Ellipse™ DR*	10.4	9.6	8.9	7.7
CD2257-40Q	Fortify Assura™ DR*	11.1	10.2	9.5	8.3
CD2257-40	Fortify Assura™ DR*	11.1	10.2	9.5	8.3
CD2231-40Q	Fortify™ DR*	10.1	9.3	8.6	7.5
CD2231-40	Fortify™ DR*	10.1	9.3	8.6	7.5
CD2211-36Q	Current™ + DR**	8.2	7.5	7.0	6.1
CD2211-36	Current™ + DR**	8.2	7.5	7.0	6.1
2207-36	Current™ DR RF**	8.2	7.5	7.0	6.1
V-268	Atlas™ II + DR**	8.2	7.5	7.0	6.1
V-243	Atlas™ + DR**	7.9	7.3	6.9	6.1

Pacing parameters: DDD, 2.5V, 0.5 ms, 60 ppm, 500 ohms

*Battery voltage range: 3.20-2.59. Three maximum charges per year.

**Battery voltage range: 3.20-2.45. Four maximum charges per year as well as monthly charging during the battery's mid-life voltage range.

SUMMARY INFORMATION

**Dual-Chamber
Implantable Cardioverter
Defibrillator (ICD) Devices**

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Survival Probability Summary

INCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
CD2411-36Q	Ellipse™ DR	99.84%	99.79%	99.72%	99.72%						
CD2411-36C	Ellipse™ DR	99.81%	99.72%	99.72%	99.72%						
CD2357-40Q	Fortify Assura™ DR	99.86%	99.83%								
CD2357-40Q	Fortify Assura™ DR†	99.79%	99.47%	98.60%	98.37%						
CD2357-40C	Fortify Assura™ DR	99.92%	99.92%								
CD2357-40C	Fortify Assura™ DR†	99.84%	99.67%	99.03%	98.95%						
CD2311-36Q	Ellipse™ DR	99.04%	98.00%	97.19%	96.25%	95.77%					
CD2311-36	Ellipse™ DR	98.94%	97.78%	97.05%	96.42%	96.05%					
CD2257-40Q	Fortify Assura™ DR†	99.87%	99.63%	99.18%	98.43%	98.37%					
CD2257-40	Fortify Assura™ DR†	99.85%	99.62%	99.43%	98.87%	98.32%					
CD2231-40Q	Fortify™ DR†	99.75%	99.55%	98.97%	98.07%	96.52%	94.81%	93.63%			
CD2231-40	Fortify™ DR†	99.88%	99.67%	99.16%	98.42%	97.07%	95.36%	94.22%			
CD2211-36Q	Current™ + DR	99.82%	99.37%	99.02%	98.55%	97.36%	94.17%	83.84%	79.55%		
CD2211-36	Current™ + DR	99.78%	99.57%	99.30%	98.49%	97.86%	93.84%	77.12%	69.12%		
2207-36	Current™ DR RF	99.66%	99.23%	98.63%	97.89%	96.64%	92.88%	78.07%	49.91%	40.61%	
V-268	Atlas™ II + DR	99.60%	99.30%	98.64%	97.01%	91.35%	65.36%	24.98%	12.43%	11.74%	11.61%
V-243	Atlas™ + DR	99.82%	99.63%	99.23%	98.31%	96.07%	90.40%	71.06%	35.61%	13.16%	10.35%

†Premature battery depletion advisory population.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Survival Probability Summary

EXCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
CD2411-36Q	Ellipse™ DR	99.84%	99.79%	99.72%	99.72%						
CD2411-36C	Ellipse™ DR	99.81%	99.72%	99.72%	99.72%						
CD2357-40Q	Fortify Assura™ DR	99.91%	99.88%								
CD2357-40Q	Fortify Assura™ DR†	99.84%	99.56%	98.76%	98.53%						
CD2357-40C	Fortify Assura™ DR	99.92%	99.92%								
CD2357-40C	Fortify Assura™ DR†	99.91%	99.83%	99.23%	99.15%						
CD2311-36Q	Ellipse™ DR	99.13%	98.16%	97.40%	96.65%	96.32%					
CD2311-36	Ellipse™ DR	99.02%	98.02%	97.46%	97.05%	96.84%					
CD2257-40Q	Fortify Assura™ DR†	99.87%	99.72%	99.36%	98.61%	98.55%					
CD2257-40	Fortify Assura™ DR†	99.90%	99.73%	99.54%	98.97%	98.43%					
CD2231-40Q	Fortify™ DR†	99.87%	99.76%	99.31%	98.62%	97.59%	96.30%	95.63%			
CD2231-40	Fortify™ DR†	99.95%	99.86%	99.49%	98.87%	97.96%	96.90%	96.70%			
CD2211-36Q	Current™ + DR	99.85%	99.58%	99.41%	99.22%	98.82%	98.53%	97.80%	97.20%		
CD2211-36	Current™ + DR	99.90%	99.76%	99.54%	99.10%	98.96%	98.18%	97.03%	96.58%		
2207-36	Current™ DR RF	99.73%	99.58%	99.20%	98.70%	98.19%	97.84%	97.02%	95.19%	94.25%	
V-268	Atlas™ II + DR	99.80%	99.67%	99.40%	99.11%	98.81%	98.47%	98.25%	98.15%	98.15%	98.15%
V-243	Atlas™ + DR	99.97%	99.90%	99.80%	99.63%	99.42%	99.18%	99.01%	98.81%	98.65%	98.65%

†Premature battery depletion advisory population.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

US Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/ FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD2411-36Q	Ellipse™ DR	15,960	1.80%	0	0.00%	1	<0.01%	0	0.00%	2	0.01%	1	<0.01%	1	<0.01%	0	0.00%	1	<0.01%	6	0.04%
CD2411-36C	Ellipse™ DR	8,338	2.20%	2	0.02%	0	0.00%	0	0.00%	4	0.05%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	6	0.07%
CD2357-40Q	Fortify Assura™ DR	14,890	1.20%	0	0.00%	0	0.00%	1	<0.01%	2	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	3	0.02%
CD2357-40Q	Fortify Assura™ DR†	12,264	7.90%	3	0.02%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	24	0.20%	0	0.00%	28	0.23%
CD2357-40C	Fortify Assura™ DR	6,180	0.90%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.02%
CD2357-40C	Fortify Assura™ DR†	6,956	9.10%	3	0.04%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	9	0.13%	0	0.00%	12	0.17%
CD2311-36Q	Ellipse™ DR	5,897	6.50%	3	0.05%	0	0.00%	0	0.00%	48	0.81%	1	0.02%	2	0.03%	0	0.00%	2	0.03%	56	0.95%
CD2311-36	Ellipse™ DR	3,746	7.30%	5	0.13%	0	0.00%	0	0.00%	20	0.53%	0	0.00%	4	0.11%	0	0.00%	3	0.08%	32	0.85%
CD2257-40Q	Fortify Assura™ DR†	6,798	9.80%	5	0.07%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	12	0.18%	3	0.04%	21	0.31%
CD2257-40	Fortify Assura™ DR†	4,235	11.50%	1	0.02%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	9	0.21%	0	0.00%	11	0.26%
CD2231-40Q	Fortify™ DR†	26,868	10.50%	7	0.03%	2	<0.01%	27	0.10%	4	0.01%	1	<0.01%	0	0.00%	120	0.45%	9	0.03%	170	0.63%
CD2231-40	Fortify™ DR†	12,094	12.80%	6	0.05%	1	<0.01%	5	0.04%	6	0.05%	0	0.00%	0	0.00%	43	0.36%	4	0.03%	65	0.54%
CD2211-36Q	Current™ + DR	8,144	12.00%	6	0.07%	0	0.00%	6	0.07%	2	0.02%	1	0.01%	0	0.00%	4	0.05%	5	0.06%	24	0.29%
CD2211-36	Current™ + DR	6,271	15.60%	2	0.03%	2	0.03%	7	0.11%	0	0.00%	1	0.02%	0	0.00%	9	0.14%	5	0.08%	26	0.41%
2207-36	Current™ DR RF	22,386	21.50%	10	0.04%	6	0.03%	20	0.09%	1	<0.01%	2	<0.01%	1	<0.01%	40	0.18%	36	0.16%	116	0.52%
V-268	Atlas™ II + DR	14,809	29.40%	6	0.04%	4	0.03%	9	0.06%	0	0.00%	0	0.00%	0	0.00%	19	0.13%	10	0.07%	48	0.32%
V-243	Atlas™ + DR	21,082	26.90%	5	0.02%	1	<0.01%	12	0.06%	1	<0.01%	0	0.00%	0	0.00%	6	0.03%	17	0.08%	42	0.20%

Definitions of malfunction categories can be found on [pages 5-6](#).

†Premature battery depletion advisory population.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

US Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/ FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD2411-36Q	Ellipse™ DR	15,960	1.80%	2	0.01%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	2	0.01%	1	<0.01%	2	0.01%	8	0.05%
CD2411-36C	Ellipse™ DR	8,338	2.20%	1	0.01%	0	0.00%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.02%
CD2357-40Q	Fortify Assura™ DR	14,890	1.20%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.01%	0	0.00%	0	0.00%	3	0.02%
CD2357-40Q	Fortify Assura™ DR†	12,264	7.90%	4	0.03%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	1	<0.01%	8	0.07%	3	0.02%	18	0.15%
CD2357-40C	Fortify Assura™ DR	6,180	0.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	1	0.02%
CD2357-40C	Fortify Assura™ DR†	6,956	9.10%	0	0.00%	1	0.01%	2	0.03%	0	0.00%	0	0.00%	0	0.00%	1	0.01%	0	0.00%	4	0.06%
CD2311-36Q	Ellipse™ DR	5,897	6.50%	4	0.07%	0	0.00%	0	0.00%	5	0.08%	0	0.00%	3	0.05%	0	0.00%	2	0.03%	14	0.24%
CD2311-36	Ellipse™ DR	3,746	7.30%	2	0.05%	0	0.00%	0	0.00%	5	0.13%	0	0.00%	3	0.08%	0	0.00%	0	0.00%	10	0.27%
CD2257-40Q	Fortify Assura™ DR†	6,798	9.80%	1	0.01%	0	0.00%	1	0.01%	0	0.00%	1	0.01%	1	0.01%	6	0.09%	0	0.00%	10	0.15%
CD2257-40	Fortify Assura™ DR†	4,235	11.50%	0	0.00%	0	0.00%	2	0.05%	0	0.00%	1	0.02%	0	0.00%	4	0.09%	1	0.02%	8	0.19%
CD2231-40Q	Fortify™ DR†	26,868	10.50%	8	0.03%	2	<0.01%	27	0.10%	2	<0.01%	0	0.00%	0	0.00%	45	0.17%	5	0.02%	89	0.33%
CD2231-40	Fortify™ DR†	12,094	12.80%	2	0.02%	0	0.00%	5	0.04%	1	<0.01%	0	0.00%	1	<0.01%	15	0.12%	3	0.02%	27	0.22%
CD2211-36Q	Current™ + DR	8,144	12.00%	4	0.05%	0	0.00%	6	0.07%	0	0.00%	11	0.14%	2	0.02%	3	0.04%	3	0.04%	29	0.36%
CD2211-36	Current™ + DR	6,271	15.60%	1	0.02%	0	0.00%	4	0.06%	0	0.00%	13	0.21%	0	0.00%	4	0.06%	0	0.00%	22	0.35%
2207-36	Current™ DR RF	22,386	21.50%	12	0.05%	2	<0.01%	9	0.04%	0	0.00%	41	0.18%	20	0.09%	19	0.08%	6	0.03%	109	0.49%
V-268	Atlas™ II + DR	14,809	29.40%	4	0.03%	0	0.00%	3	0.02%	0	0.00%	0	0.00%	1	<0.01%	6	0.04%	5	0.03%	19	0.13%
V-243	Atlas™ + DR	21,082	26.90%	3	0.01%	0	0.00%	4	0.02%	0	0.00%	0	0.00%	4	0.02%	4	0.02%	2	<0.01%	17	0.08%

Definitions of malfunction categories can be found on [pages 5-6](#).

†Premature battery depletion advisory population.

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Worldwide Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD2411-36Q	Ellipse™ DR	16,363	2.06%	0	0.00%	1	<0.01%	0	0.00%	2	0.01%	1	<0.01%	1	<0.01%	0	0.00%	1	<0.01%	6	0.04%
CD2411-36C	Ellipse™ DR	8,476	2.65%	2	0.02%	0	0.00%	0	0.00%	4	0.05%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	6	0.07%
CD2357-40Q	Fortify Assura™ DR	27,558	4.34%	3	0.01%	1	<0.01%	1	<0.01%	2	<0.01%	0	0.00%	0	0.00%	24	0.09%	0	0.00%	31	0.11%
CD2357-40C	Fortify Assura™ DR	13,400	5.51%	4	0.03%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	9	0.07%	0	0.00%	13	0.10%
CD2311-36Q	Ellipse™ DR	5,906	7.94%	3	0.05%	0	0.00%	0	0.00%	48	0.81%	1	0.02%	2	0.03%	0	0.00%	2	0.03%	56	0.95%
CD2311-36	Ellipse™ DR	3,756	8.15%	5	0.13%	0	0.00%	0	0.00%	20	0.53%	0	0.00%	4	0.11%	0	0.00%	3	0.08%	32	0.85%
CD2257-40Q	Fortify Assura™ DR	6,781	10.16%	5	0.07%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	12	0.18%	3	0.04%	21	0.31%
CD2257-40	Fortify Assura™ DR	4,236	12.06%	1	0.02%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	9	0.21%	0	0.00%	11	0.26%
CD2231-40Q	Fortify™ DR	28,132	10.66%	7	0.02%	2	<0.01%	28	0.10%	4	0.01%	1	<0.01%	0	0.00%	123	0.44%	10	0.04%	175	0.62%
CD2231-40	Fortify™ DR	13,904	12.01%	6	0.04%	2	0.01%	5	0.04%	6	0.04%	0	0.00%	0	0.00%	44	0.32%	5	0.04%	68	0.49%
CD2211-36Q	Current™ + DR	15,192	7.83%	7	0.05%	1	<0.01%	8	0.05%	4	0.03%	1	<0.01%	0	0.00%	7	0.05%	10	0.07%	38	0.25%
CD2211-36	Current™ + DR	13,485	8.21%	4	0.03%	5	0.04%	8	0.06%	1	<0.01%	1	<0.01%	0	0.00%	11	0.08%	9	0.07%	39	0.29%
2207-36	Current™ DR RF	33,051	17.28%	17	0.05%	11	0.03%	28	0.08%	12	0.04%	3	<0.01%	2	<0.01%	59	0.18%	45	0.14%	177	0.54%
V-268	Atlas™ II + DR	25,779	19.22%	15	0.06%	5	0.02%	19	0.07%	1	<0.01%	0	0.00%	0	0.00%	32	0.12%	19	0.07%	91	0.35%
V-243	Atlas™ + DR	34,105	18.83%	5	0.01%	3	<0.01%	25	0.07%	1	<0.01%	0	0.00%	0	0.00%	14	0.04%	30	0.09%	78	0.23%

Definitions of malfunction categories can be found on [pages 5-6](#).

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Worldwide Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD2411-36Q	Ellipse™ DR	16,363	2.06%	2	0.01%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	2	0.01%	1	<0.01%	2	0.01%	8	0.05%
CD2411-36C	Ellipse™ DR	8,476	2.65%	1	0.01%	0	0.00%	0	0.00%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.02%
CD2357-40Q	Fortify Assura™ DR	27,558	4.34%	5	0.02%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	3	0.01%	8	0.03%	3	0.01%	21	0.08%
CD2357-40C	Fortify Assura™ DR	13,400	5.51%	0	0.00%	1	<0.01%	2	0.01%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%	0	0.00%	5	0.04%
CD2311-36Q	Ellipse™ DR	5,906	7.94%	4	0.07%	0	0.00%	0	0.00%	5	0.08%	0	0.00%	3	0.05%	0	0.00%	2	0.03%	14	0.24%
CD2311-36	Ellipse™ DR	3,756	8.15%	2	0.05%	0	0.00%	0	0.00%	5	0.13%	0	0.00%	3	0.08%	0	0.00%	0	0.00%	10	0.27%
CD2257-40Q	Fortify Assura™ DR	6,781	10.16%	1	0.01%	0	0.00%	1	0.01%	0	0.00%	1	0.01%	1	0.01%	6	0.09%	0	0.00%	10	0.15%
CD2257-40	Fortify Assura™ DR	4,236	12.06%	0	0.00%	0	0.00%	2	0.05%	0	0.00%	1	0.02%	0	0.00%	4	0.09%	1	0.02%	8	0.19%
CD2231-40Q	Fortify™ DR	28,132	10.66%	8	0.03%	2	<0.01%	28	0.10%	2	<0.01%	0	0.00%	0	0.00%	48	0.17%	5	0.02%	93	0.33%
CD2231-40	Fortify™ DR	13,904	12.01%	2	0.01%	0	0.00%	5	0.04%	1	<0.01%	0	0.00%	2	0.01%	16	0.12%	3	0.02%	29	0.21%
CD2211-36Q	Current™ + DR	15,192	7.83%	9	0.06%	0	0.00%	9	0.06%	0	0.00%	13	0.09%	3	0.02%	5	0.03%	4	0.03%	43	0.28%
CD2211-36	Current™ + DR	13,485	8.21%	1	<0.01%	0	0.00%	4	0.03%	1	<0.01%	13	0.10%	1	<0.01%	5	0.04%	1	<0.01%	26	0.19%
2207-36	Current™ DR RF	33,051	17.28%	19	0.06%	5	0.02%	14	0.04%	4	0.01%	81	0.25%	32	0.10%	26	0.08%	11	0.03%	192	0.58%
V-268	Atlas™ II + DR	25,779	19.22%	7	0.03%	0	0.00%	8	0.03%	1	<0.01%	0	0.00%	2	<0.01%	9	0.03%	6	0.02%	33	0.13%
V-243	Atlas™ + DR	34,105	18.83%	6	0.02%	0	0.00%	6	0.02%	0	0.00%	0	0.00%	8	0.02%	6	0.02%	4	0.01%	30	0.09%

Definitions of malfunction categories can be found on [pages 5-6](#).

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	INAPPROPRIATE SHOCK		LOSS OF TELEMETRY		PERICARDIAL EFFUSION		PREMATURE BATTERY DEPLETION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD2231-40Q	390	159	21,754	0	0.00%	0	0.00%	0	0.00%	4	1.03%	0	0.00%	4	1.03%
CD2231-40	175	58	7,769	0	0.00%	0	0.00%	0	0.00%	1	0.57%	0	0.00%	1	0.57%
CD2211-36Q	835	360	52,193	0	0.00%	0	0.00%	0	0.00%	4	0.48%	1	0.12%	5	0.60%
CD2211-36	123	26	6,344	1	0.81%	0	0.00%	0	0.00%	1	0.81%	0	0.00%	2	1.63%
2207-36	631	73	32,853	1	0.16%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.16%

A list of of complications can be found on [page 12](#).

Dual-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Actively Monitored Study Data Summary

MALFUNCTIONS WITH COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD2231-40Q	Fortify™ DR	390	14.10%	0	0.00%	0	0.00%	1	0.26%	0	0.00%	0	0.00%	0	0.00%	2	0.51%	1	0.26%	4	1.03%
CD2231-40	Fortify™ DR	175	15.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD2211-36Q	Current™ + DR	835	14.00%	0	0.00%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.12%
CD2211-36	Current™ + DR	123	18.70%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.81%	1	0.81%
2207-36	Current™ DR RF	631	30.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.16%	0	0.00%	1	0.16%

MALFUNCTIONS WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD2231-40Q	Fortify™ DR	390	14.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.51%	0	0.00%	2	0.51%
CD2231-40	Fortify™ DR	175	15.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD2211-36Q	Current™ + DR	835	14.00%	1	0.12%	0	0.00%	2	0.24%	0	0.00%	2	0.24%	0	0.00%	1	0.12%	1	0.12%	7	0.84%
CD2211-36	Current™ + DR	123	18.70%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.81%	1	0.81%
2207-36	Current™ DR RF	631	30.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.32%	0	0.00%	1	0.16%	0	0.00%	3	0.48%

Definitions of malfunction categories can be found on [pages 5-6](#).

SINGLE-CHAMBER
**Implantable Cardioverter
Defibrillator (ICD) Devices**

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

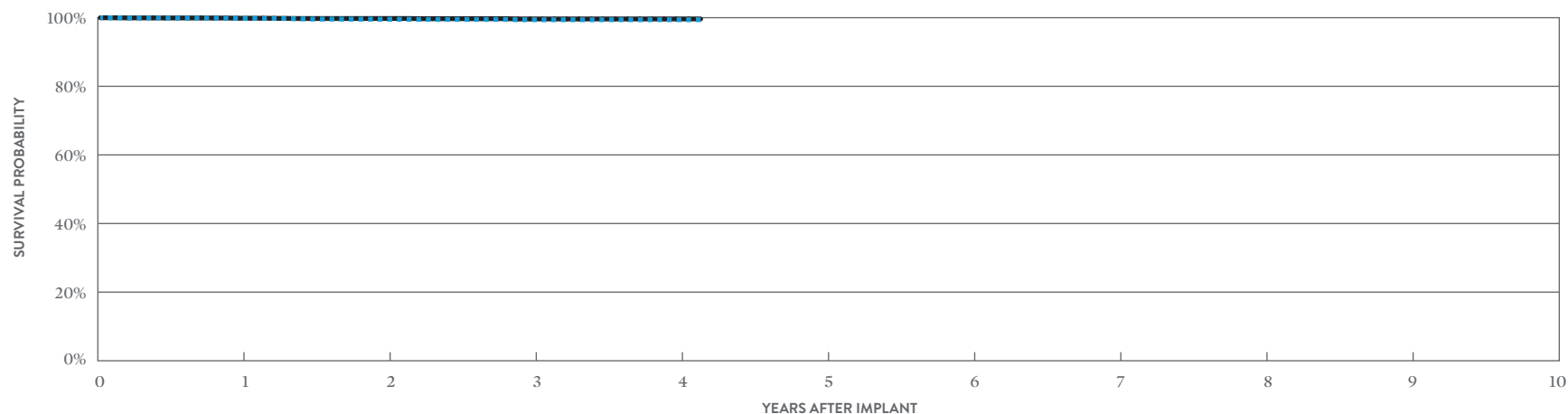
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ VR

MODEL CD1411-36Q*

US Regulatory Approval	June 2013
Registered US Implants	12,672
Estimated Active US Implants	9,901
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	7
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.02%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	6	0.05%	1	<0.01%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	2	0.02%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	8	0.06%	4	0.03%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.80%	99.53%	99.41%	99.41%	99.41%
± 1 STANDARD ERROR	0.04%	0.07%	0.10%	0.10%	0.10%
SAMPLE SIZE	10,510	6,720	3,620	1,220	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 50 MONTHS
SURVIVAL PROBABILITY	99.84%	99.76%	99.63%	99.63%	99.63%
± 1 STANDARD ERROR	0.03%	0.05%	0.08%	0.08%	0.08%

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

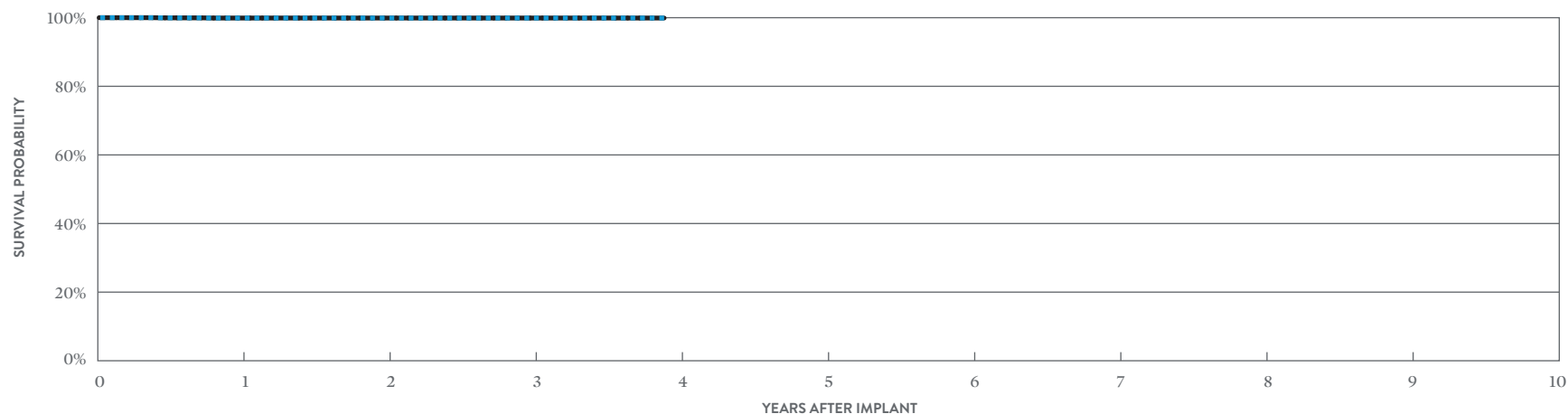
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ VR

MODEL CD1411-36C*

US Regulatory Approval	June 2013
Registered US Implants	5,095
Estimated Active US Implants	3,950
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	0
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.02%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	1	0.02%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	2	0.04%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 47 MONTHS
SURVIVAL PROBABILITY	99.90%	99.90%	99.90%	99.90%
± 1 STANDARD ERROR	0.05%	0.05%	0.05%	0.05%
SAMPLE SIZE	4,280	2,830	1,560	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 47 MONTHS
SURVIVAL PROBABILITY	99.90%	99.90%	99.90%	99.90%
± 1 STANDARD ERROR	0.05%	0.05%	0.05%	0.05%

*Parylene coating.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

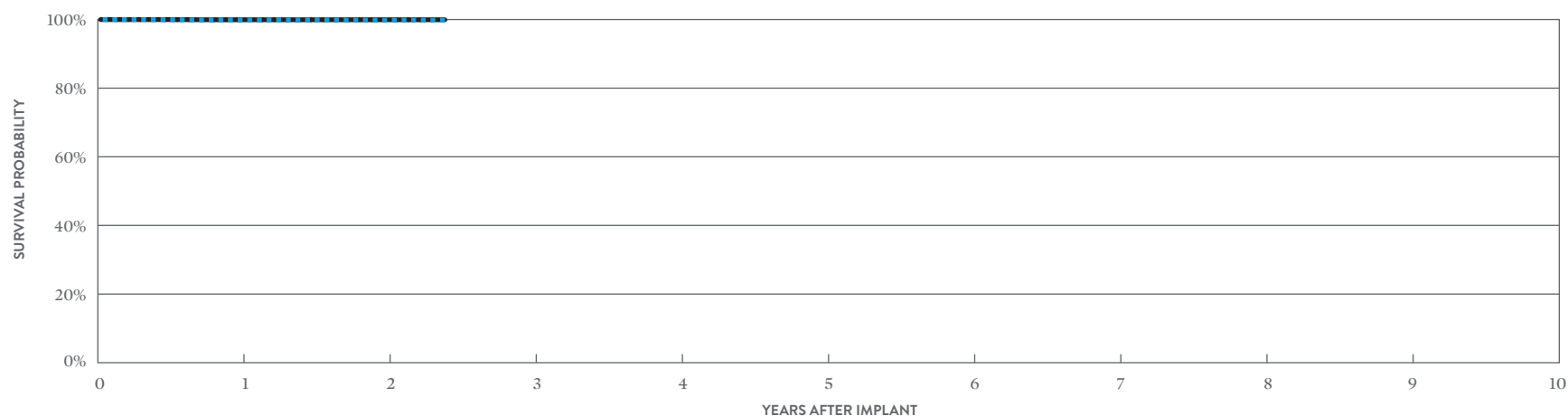
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ VR

MODEL CD1357-40Q* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	10,558
Estimated Active US Implants	9,063
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	3
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	1	<0.01%	0	0.00%
Total	1	<0.01%	2	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	99.88%	99.81%	99.81%
± 1 STANDARD ERROR	0.04%	0.06%	0.06%
SAMPLE SIZE	8,210	3,690	350

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	99.93%	99.93%	99.93%
± 1 STANDARD ERROR	0.03%	0.03%	0.03%

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

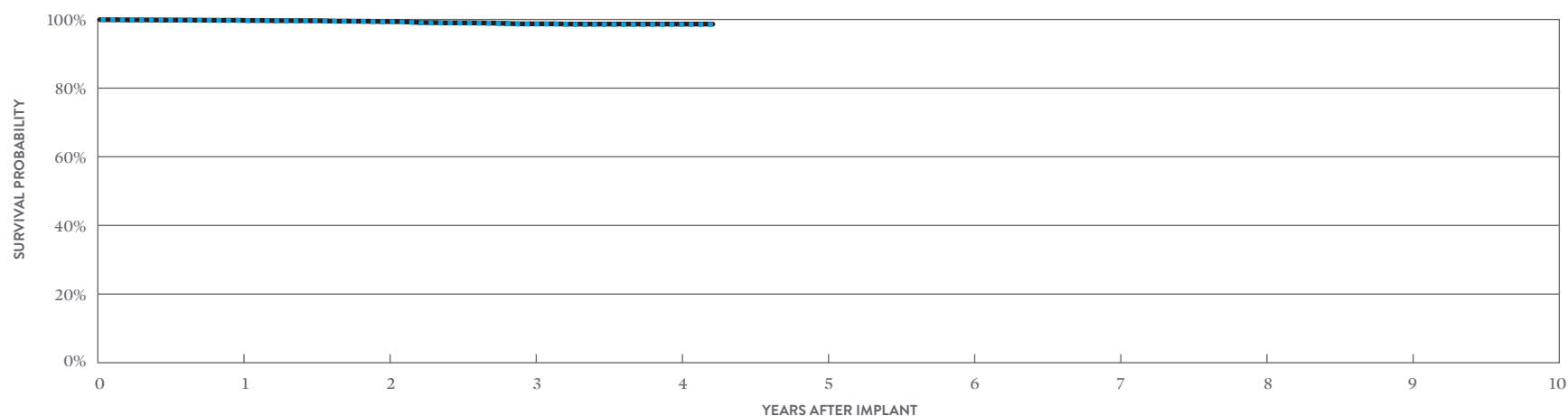
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ VR

MODEL CD1357-40Q* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	10,215
Estimated Active US Implants	6,875
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	3
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.03%	8	0.08%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	0	0.00%	1	<0.01%
High Voltage Capacitor	1	<0.01%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	15	0.15%	8	0.08%
Other	2	0.02%	2	0.02%
Total	22	0.22%	19	0.19%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 51 MONTHS
SURVIVAL PROBABILITY	99.74%	99.33%	98.67%	98.53%	98.53%
± 1 STANDARD ERROR	0.05%	0.08%	0.14%	0.15%	0.15%
SAMPLE SIZE	9,560	8,240	5,710	2,140	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	AT 51 MONTHS
SURVIVAL PROBABILITY	99.77%	99.40%	98.74%	98.67%	98.67%
± 1 STANDARD ERROR	0.05%	0.08%	0.13%	0.14%	0.14%

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

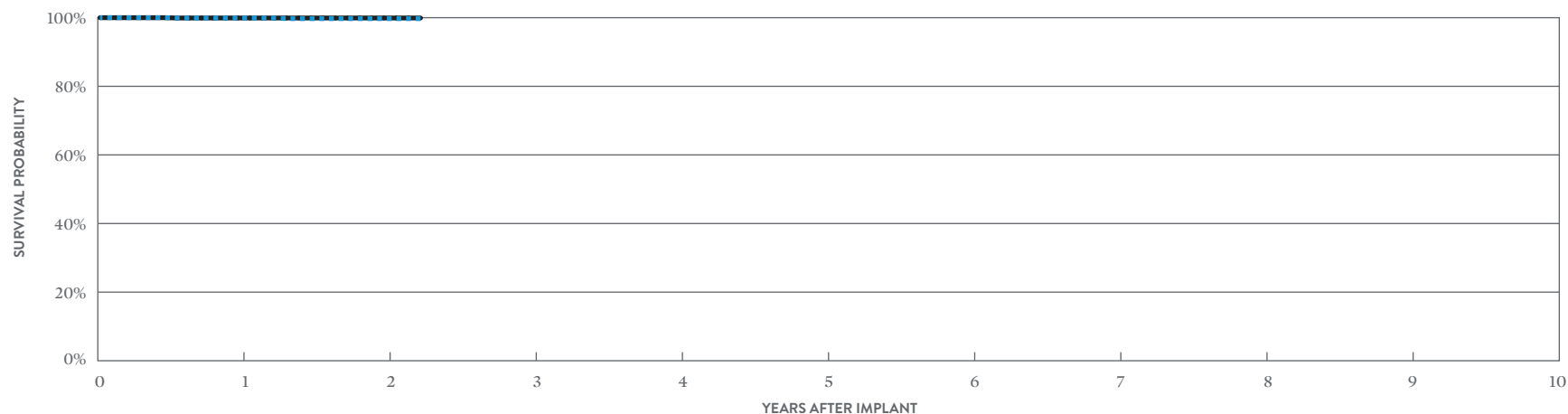
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ VR

MODEL CD1357-40C* (NON-ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	3,122
Estimated Active US Implants	2,711
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	1
Max. Delivered Energy	40 joules
Number of US Advisories	None for this population

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.03%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	1	0.03%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 27 MONTHS
SURVIVAL PROBABILITY	99.91%	99.72%	99.72%
± 1 STANDARD ERROR	0.07%	0.15%	0.15%
SAMPLE SIZE	2,290	910	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 27 MONTHS
SURVIVAL PROBABILITY	99.91%	99.91%	99.91%
± 1 STANDARD ERROR	0.07%	0.07%	0.07%

*Parylene coating.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

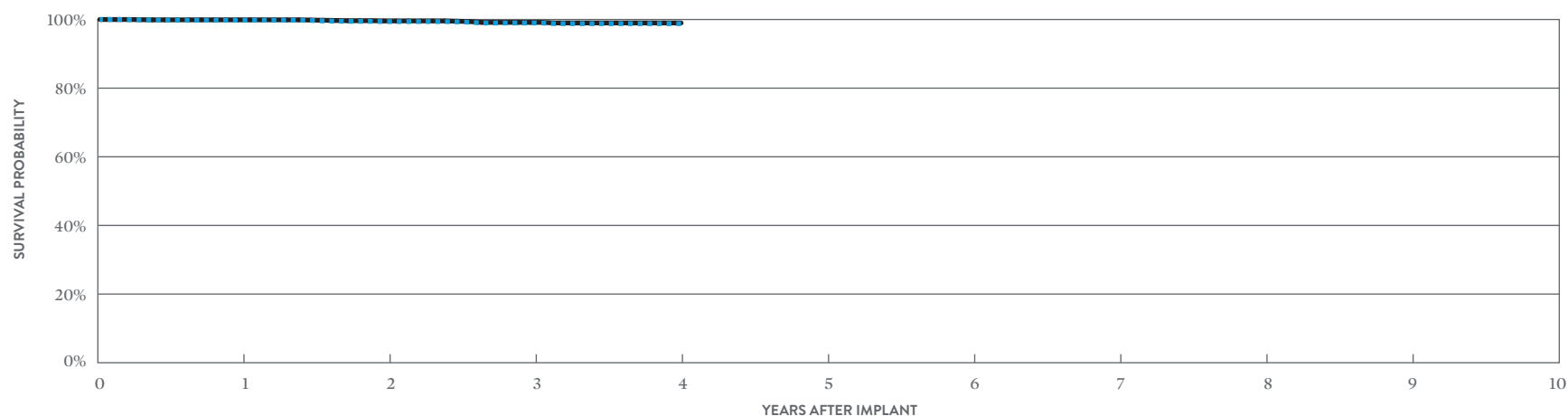
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ VR

MODEL CD1357-40C* (ADVISORY POPULATION)

US Regulatory Approval	June 2013
Registered US Implants	4,131
Estimated Active US Implants	2,792
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	3
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.07%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	2	0.05%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	1	0.02%	0	0.00%
Possible Early Battery Depletion	2	0.05%	4	0.10%
Other	0	0.00%	0	0.00%
Total	6	0.15%	6	0.15%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4
SURVIVAL PROBABILITY	99.80%	99.39%	98.97%	98.79%
± 1 STANDARD ERROR	0.07%	0.12%	0.19%	0.23%
SAMPLE SIZE	3,890	3,280	2,120	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4
SURVIVAL PROBABILITY	99.90%	99.59%	99.16%	98.99%
± 1 STANDARD ERROR	0.05%	0.10%	0.18%	0.22%

*Parylene coating.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

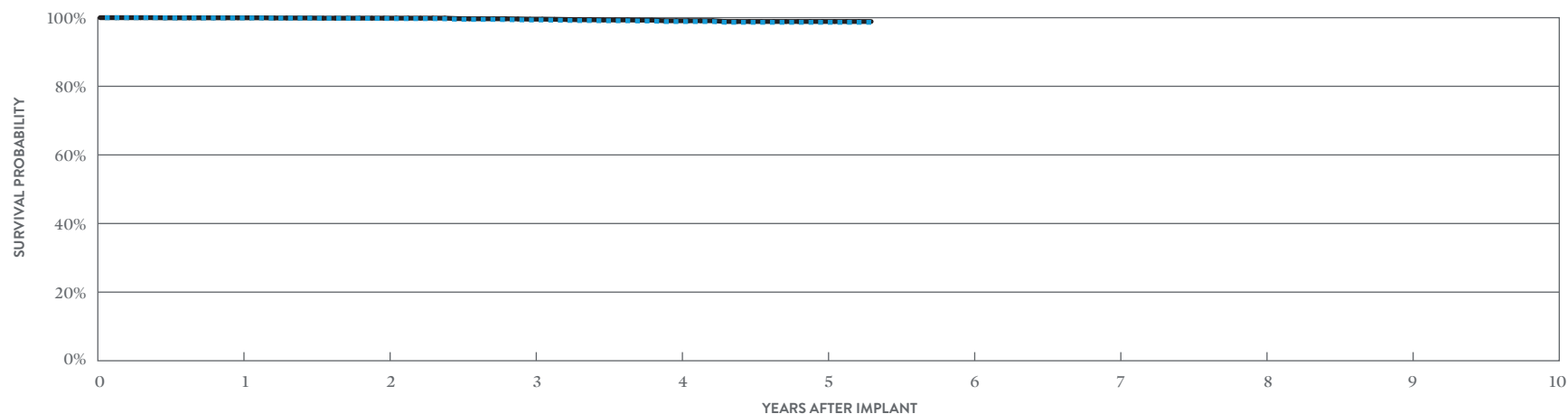
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ VR

MODEL CD1257-40Q* (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	5,077
Estimated Active US Implants	3,000
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	5
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	1	0.02%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	8	0.16%	7	0.14%
Other	1	0.02%	0	0.00%
Total	10	0.20%	7	0.14%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 64 MONTHS
SURVIVAL PROBABILITY	99.92%	99.77%	99.33%	98.87%	98.65%	98.65%
± 1 STANDARD ERROR	0.04%	0.07%	0.12%	0.18%	0.20%	0.20%
SAMPLE SIZE	4,800	4,270	3,810	3,190	1,650	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 64 MONTHS
SURVIVAL PROBABILITY	99.96%	99.87%	99.57%	99.11%	98.89%	98.89%
± 1 STANDARD ERROR	0.03%	0.05%	0.10%	0.16%	0.19%	0.19%

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

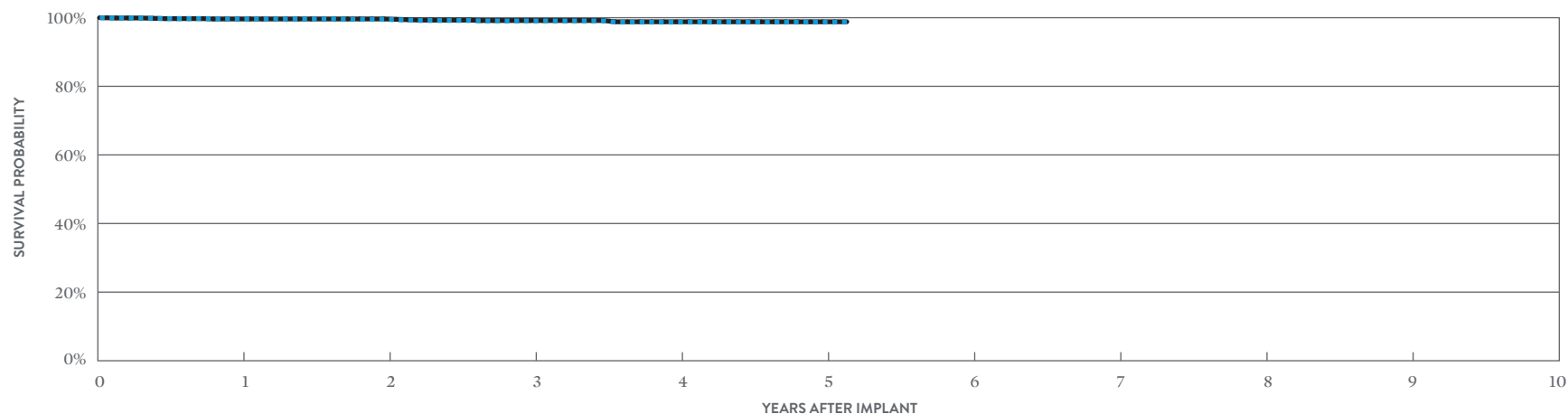
CUSTOMER REPORTED PERFORMANCE DATA

Fortify Assura™ VR

MODEL CD1257-40 (ADVISORY POPULATION)

US Regulatory Approval	May 2012
Registered US Implants	2,294
Estimated Active US Implants	1,338
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	1
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	0.04%	0	0.00%
Electrical Interconnect	2	0.09%	0	0.00%
Battery	1	0.04%	1	0.04%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	2	0.09%	1	0.04%
Other	1	0.04%	1	0.04%
Total	7	0.31%	3	0.13%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 62 MONTHS
SURVIVAL PROBABILITY	99.63%	99.52%	99.06%	98.70%	98.70%	98.70%
± 1 STANDARD ERROR	0.13%	0.15%	0.22%	0.27%	0.27%	0.27%
SAMPLE SIZE	2,150	1,900	1,680	1,330	700	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 62 MONTHS
SURVIVAL PROBABILITY	99.63%	99.63%	99.17%	98.81%	98.81%	98.81%
± 1 STANDARD ERROR	0.13%	0.13%	0.21%	0.26%	0.26%	0.26%

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

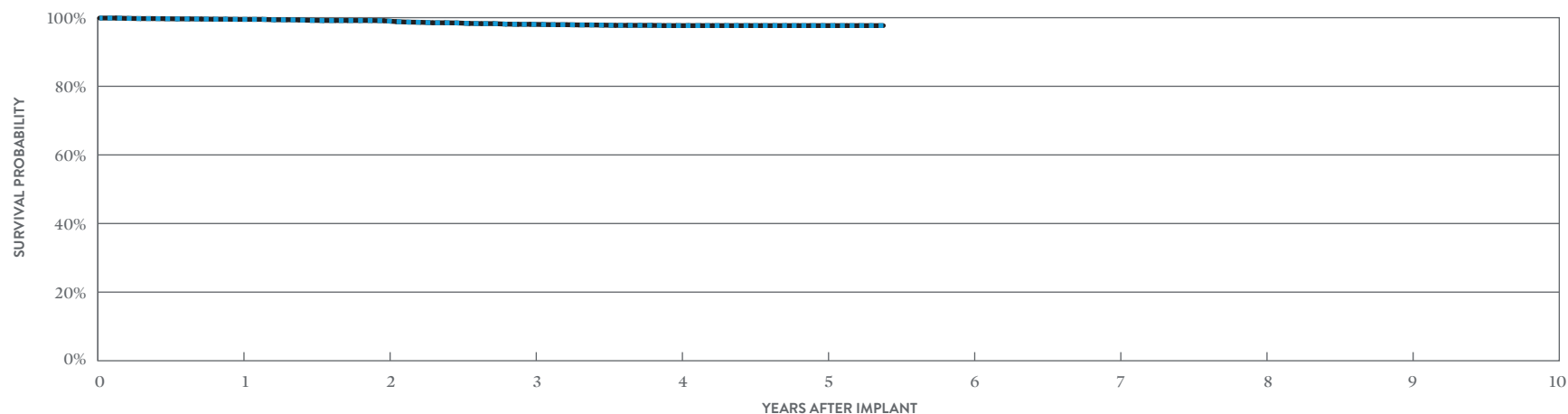
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ VR

MODEL CD1311-36Q*

US Regulatory Approval	May 2012
Registered US Implants	4,742
Estimated Active US Implants	2,854
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	0
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	0.02%	1	0.02%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	29	0.61%	3	0.06%
Software/Firmware	1	0.02%	0	0.00%
Mechanical	1	0.02%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	1	0.02%	2	0.04%
Total	33	0.70%	6	0.13%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 65 MONTHS
SURVIVAL PROBABILITY	99.51%	99.11%	98.07%	97.69%	97.69%	97.69%
± 1 STANDARD ERROR	0.10%	0.14%	0.22%	0.24%	0.24%	0.24%
SAMPLE SIZE	4,470	3,970	3,530	2,970	1,680	270

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 65 MONTHS
SURVIVAL PROBABILITY	99.51%	99.11%	98.07%	97.69%	97.69%	97.69%
± 1 STANDARD ERROR	0.10%	0.14%	0.22%	0.24%	0.24%	0.24%

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

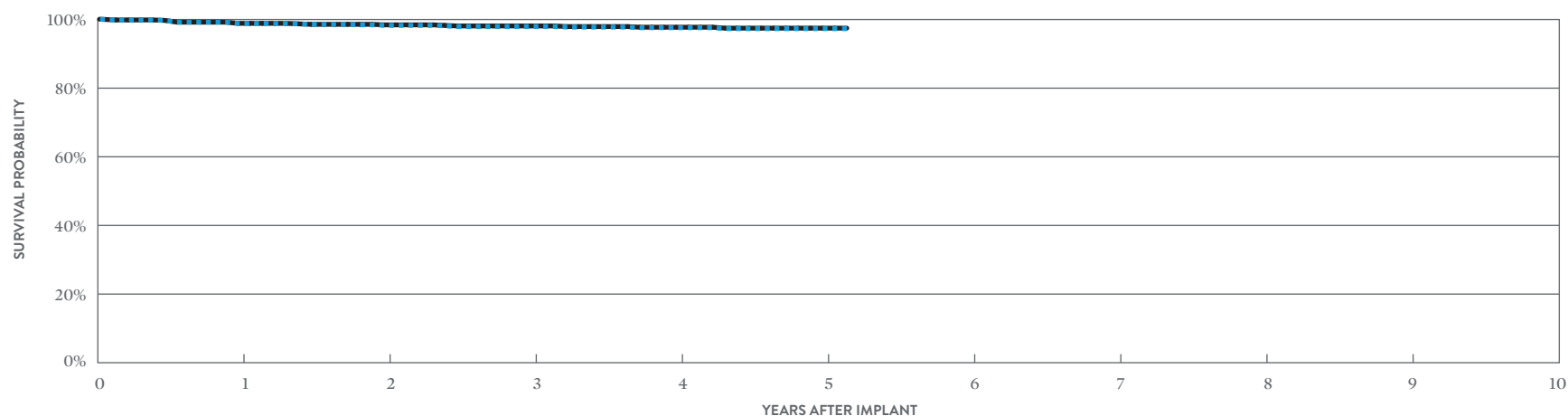
CUSTOMER REPORTED PERFORMANCE DATA

Ellipse™ VR

MODEL CD1311-36

US Regulatory Approval	May 2012
Registered US Implants	1,620
Estimated Active US Implants	992
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	1
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 327)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.12%	0	0.00%
Electrical Interconnect	1	0.06%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	6	0.37%	2	0.12%
Software/Firmware	0	0.00%	1	0.06%
Mechanical	2	0.12%	1	0.06%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	11	0.68%	4	0.25%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 62 MONTHS
SURVIVAL PROBABILITY	98.88%	98.29%	97.97%	97.61%	97.33%	97.33%
± 1 STANDARD ERROR	0.22%	0.32%	0.37%	0.42%	0.46%	0.46%
SAMPLE SIZE	1,530	1,360	1,220	1,020	570	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	AT 62 MONTHS
SURVIVAL PROBABILITY	98.88%	98.44%	98.12%	97.75%	97.48%	97.48%
± 1 STANDARD ERROR	0.22%	0.31%	0.36%	0.40%	0.45%	0.45%

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

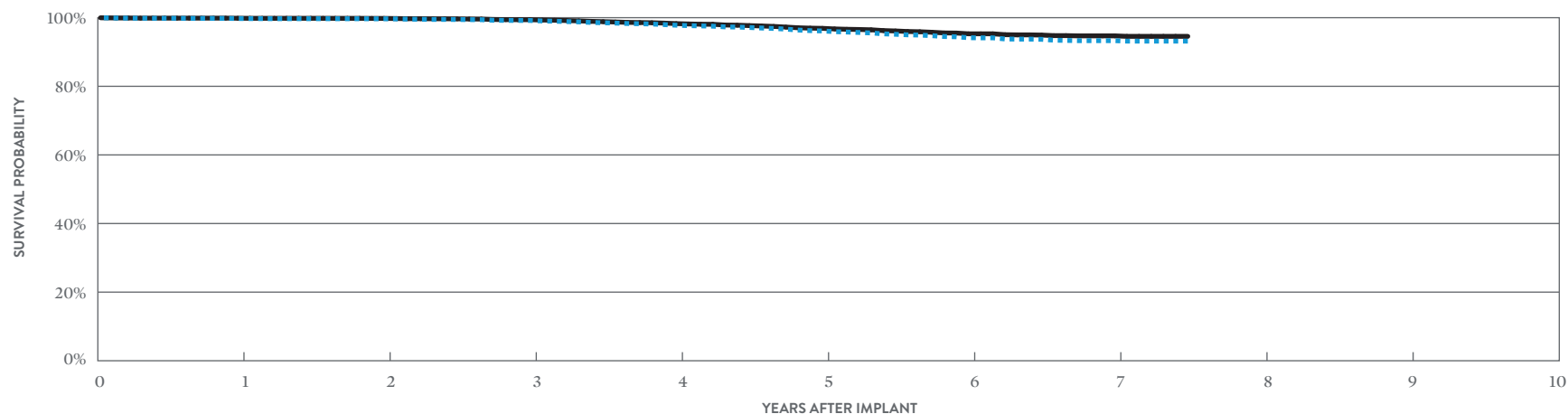
CUSTOMER REPORTED PERFORMANCE DATA

Fortify™ VR

MODEL CD1231-40Q* (ADVISORY POPULATION)

US Regulatory Approval	May 2010
Registered US Implants	16,186
Estimated Active US Implants	7,717
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	49
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	7	0.04%	4	0.02%
Electrical Interconnect	2	0.01%	0	0.00%
Battery	15	0.09%	21	0.13%
High Voltage Capacitor	1	<0.01%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	92	0.57%	40	0.25%
Other	6	0.04%	2	0.01%
Total	123	0.76%	67	0.41%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 90 MONTHS
SURVIVAL PROBABILITY	99.75%	99.67%	99.18%	97.80%	96.13%	94.16%	93.30%	93.16%
± 1 STANDARD ERROR	0.04%	0.05%	0.08%	0.13%	0.19%	0.24%	0.29%	0.30%
SAMPLE SIZE	15,130	13,300	11,870	10,570	9,210	6,790	3,290	350

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	at 90 months
SURVIVAL PROBABILITY	99.84%	99.79%	99.41%	98.27%	96.89%	95.30%	94.70%	94.56%
± 1 STANDARD ERROR	0.03%	0.04%	0.07%	0.12%	0.17%	0.22%	0.26%	0.27%

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

Fortify™ VR

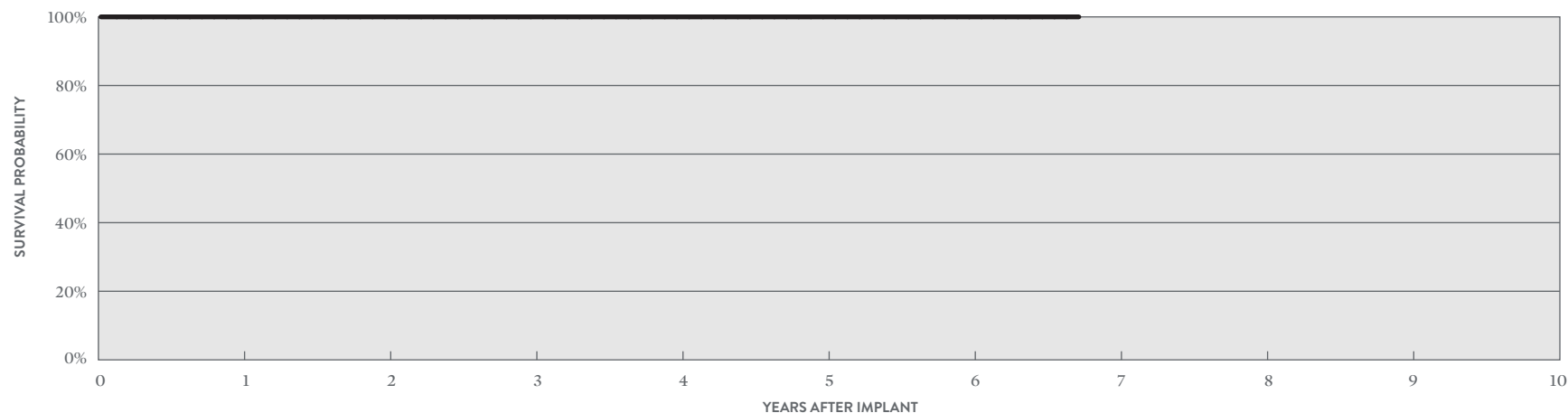
MODEL CD1231-40Q*

US Regulatory Approval	May 2010
Number of Devices Enrolled in Study	159
Active Devices Enrolled in Study	82
Cumulative Months of Follow-up	9,579
Estimated Longevity	(see table on page 152)
Max. Delivered Energy	40 joules

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	1	0.63%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	2	1.26%	1	0.63%
Other	0	0.00%	0	0.00%
Total	2	1.26%	2	1.26%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	AT 81 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
SAMPLE SIZE	160	150	130	110	100	90	50

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

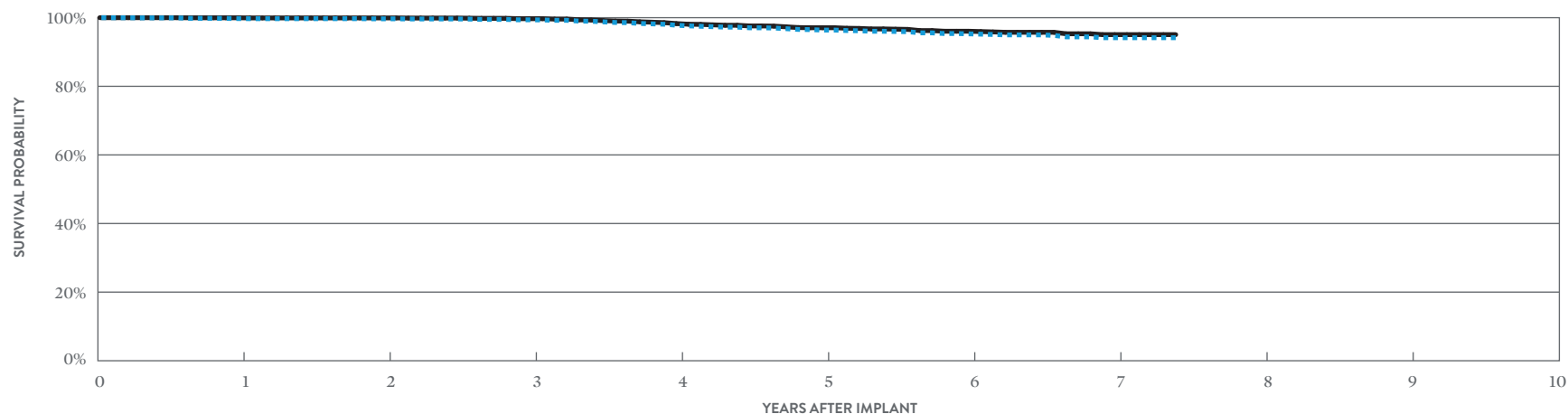
CUSTOMER REPORTED PERFORMANCE DATA

Fortify™ VR

MODEL CD1231-40 (ADVISORY POPULATION)

US Regulatory Approval	May 2010
Registered US Implants	6,782
Estimated Active US Implants	3,147
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	16
Max. Delivered Energy	40 joules
Number of US Advisories (see pg. 326)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	0.03%	4	0.06%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	4	0.06%	2	0.03%
High Voltage Capacitor	6	0.09%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.01%
Possible Early Battery Depletion	29	0.43%	16	0.24%
Other	3	0.04%	3	0.04%
Total	44	0.65%	26	0.38%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 89 MONTHS
SURVIVAL PROBABILITY	99.74%	99.67%	99.38%	97.80%	96.39%	95.29%	94.16%	94.16%
± 1 STANDARD ERROR	0.06%	0.07%	0.11%	0.20%	0.28%	0.33%	0.45%	0.45%
SAMPLE SIZE	6,330	5,550	4,920	4,370	3,830	2,810	1,350	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 89 MONTHS
SURVIVAL PROBABILITY	99.97%	99.93%	99.71%	98.28%	97.07%	96.05%	95.06%	95.06%
± 1 STANDARD ERROR	0.02%	0.03%	0.08%	0.18%	0.26%	0.32%	0.42%	0.42%

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

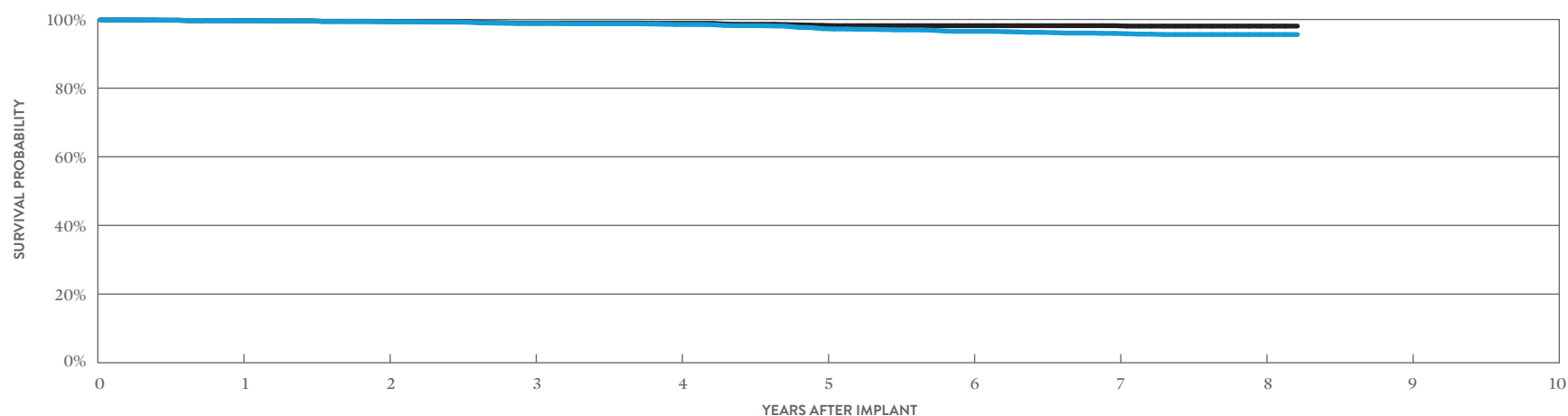
CUSTOMER REPORTED PERFORMANCE DATA

Current™ + VR

MODEL CD1211-36Q*

US Regulatory Approval	February 2009
Registered US Implants	4,431
Estimated Active US Implants	1,849
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	26
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	4	0.09%	3	0.07%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	5	0.11%	3	0.07%
High Voltage Capacitor	1	0.02%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	6	0.14%	1	0.02%
Other	2	0.05%	2	0.05%
Total	18	0.41%	9	0.20%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 99 MONTHS
SURVIVAL PROBABILITY	99.61%	99.36%	98.83%	98.55%	97.32%	96.56%	95.96%	95.62%	95.62%
± 1 STANDARD ERROR	0.09%	0.12%	0.18%	0.20%	0.28%	0.35%	0.39%	0.41%	0.41%
SAMPLE SIZE	4,120	3,610	3,220	2,850	2,520	2,200	1,900	1,120	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 99 MONTHS
SURVIVAL PROBABILITY	99.67%	99.41%	98.95%	98.87%	98.28%	98.20%	98.20%	98.08%	98.08%
± 1 STANDARD ERROR	0.09%	0.12%	0.17%	0.18%	0.23%	0.24%	0.24%	0.25%	0.25%

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

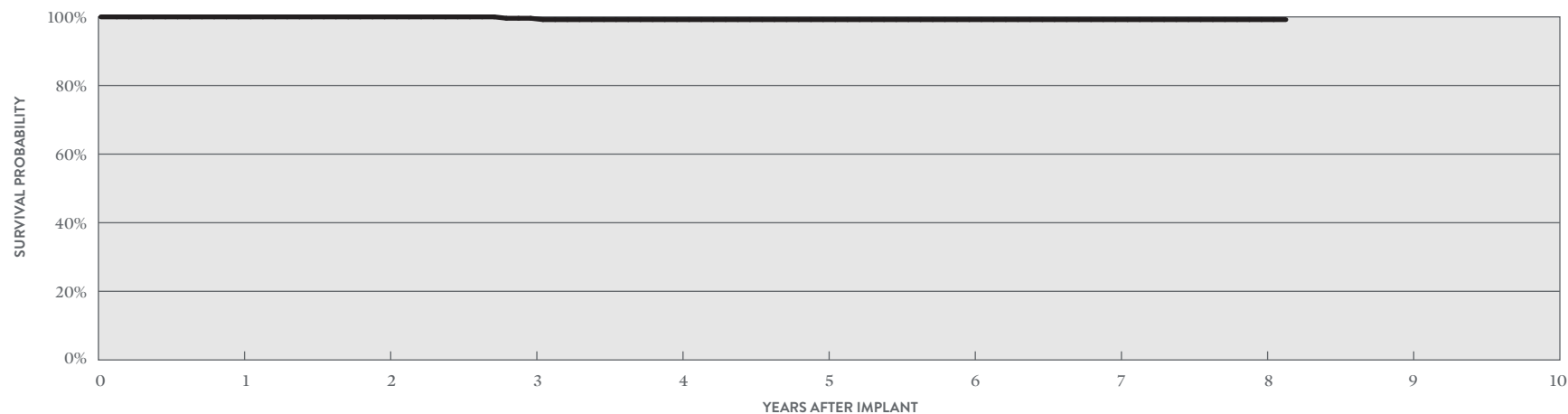
Current™ + VR

MODEL CD1211-36Q*

US Regulatory Approval	February 2009
Number of Devices Enrolled in Study	363
Active Devices Enrolled in Study	166
Cumulative Months of Follow-up	22,028
Estimated Longevity	(see table on page 152)
Max. Delivered Energy	36 joules

QUALIFYING COMPLICATIONS	QTY	RATE
Inappropriate Shock	1	0.28%
Premature Battery Depletion	1	0.28%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	1	0.28%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	1	0.28%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	8	AT 98 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	99.60%	99.20%	99.20%	99.20%	99.20%	99.20%	99.20%
± 1 STANDARD ERROR	0.00%	0.00%	0.40%	0.57%	0.57%	0.57%	0.57%	0.57%	0.57%
SAMPLE SIZE	350	310	260	230	200	180	170	120	50

*DF4-LLHH connector type.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

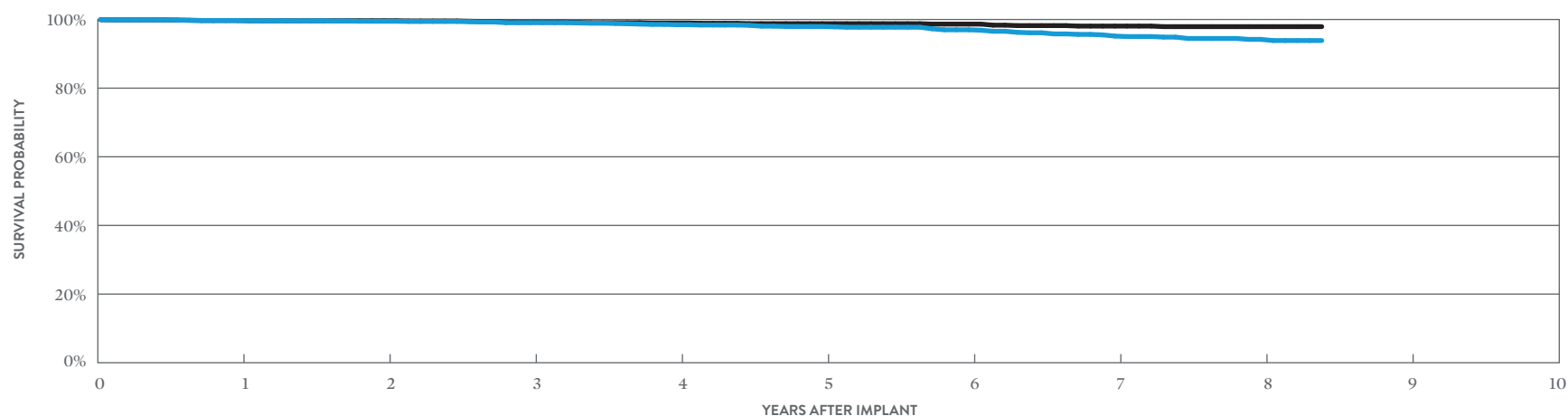
CUSTOMER REPORTED PERFORMANCE DATA

Current™ + VR

MODEL CD1211-36

US Regulatory Approval	February 2009
Registered US Implants	3,637
Estimated Active US Implants	1,392
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	28
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	0.08%	3	0.08%
Electrical Interconnect	2	0.05%	0	0.00%
Battery	4	0.11%	0	0.00%
High Voltage Capacitor	2	0.05%	0	0.00%
Software/Firmware	0	0.00%	1	0.03%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	4	0.11%	2	0.05%
Other	1	0.03%	0	0.00%
Total	16	0.44%	6	0.16%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	99.71%	99.50%	99.08%	98.49%	97.96%	97.00%	95.17%	94.21%	93.87%
± 1 STANDARD ERROR	0.09%	0.12%	0.18%	0.23%	0.28%	0.36%	0.47%	0.57%	0.62%
SAMPLE SIZE	3,380	2,960	2,640	2,340	2,060	1,760	1,470	970	260

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	99.71%	99.64%	99.22%	98.97%	98.78%	98.66%	98.08%	97.92%	97.92%
± 1 STANDARD ERROR	0.09%	0.10%	0.16%	0.19%	0.21%	0.23%	0.30%	0.32%	0.32%

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

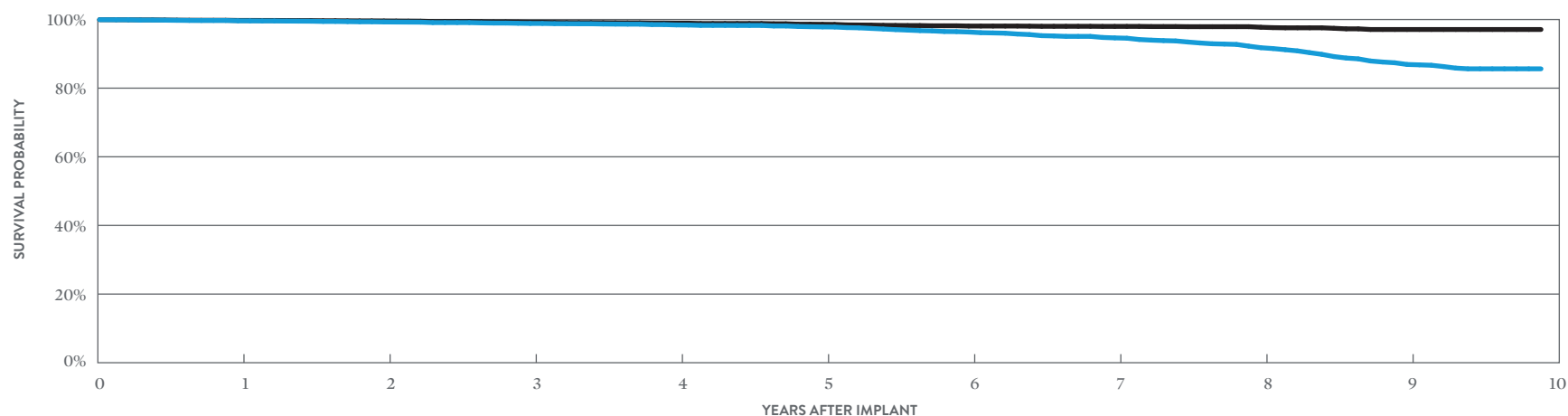
CUSTOMER REPORTED PERFORMANCE DATA

Current™ VR RF

MODEL 1207-36

US Regulatory Approval	September 2007
Registered US Implants	13,287
Estimated Active US Implants	3,437
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	216
Max. Delivered Energy	36 joules
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	6	0.05%	8	0.06%
Electrical Interconnect	10	0.08%	0	0.00%
Battery	10	0.08%	5	0.04%
High Voltage Capacitor	1	<0.01%	1	<0.01%
Software/Firmware	0	0.00%	4	0.03%
Mechanical	0	0.00%	4	0.03%
Possible Early Battery Depletion	13	0.10%	14	0.11%
Other	9	0.07%	6	0.05%
Total	49	0.37%	42	0.32%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.62%	99.28%	98.85%	98.42%	97.84%	96.35%	94.64%	91.78%	86.90%	85.63%
± 1 STANDARD ERROR	0.05%	0.08%	0.10%	0.12%	0.15%	0.21%	0.26%	0.33%	0.47%	0.59%
SAMPLE SIZE	12,350	10,720	9,450	8,410	7,510	6,680	5,910	4,970	3,030	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.73%	99.57%	99.19%	98.93%	98.60%	98.09%	97.99%	97.74%	97.09%	97.09%
± 1 STANDARD ERROR	0.04%	0.06%	0.08%	0.10%	0.12%	0.15%	0.15%	0.16%	0.22%	0.22%

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

ACTIVELY MONITORED STUDY DATA

Current™ VR RF

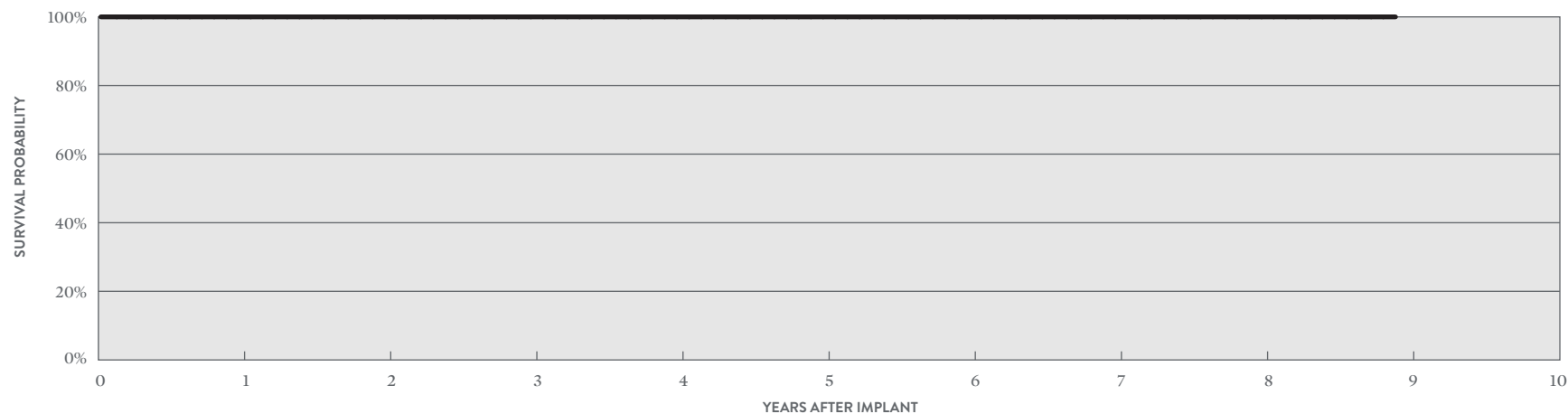
MODEL 1207-36

US Regulatory Approval	September 2007
Number of Devices Enrolled in Study	395
Active Devices Enrolled in Study	78
Cumulative Months of Follow-up	21,682
Estimated Longevity	(see table on page 152)
Max. Delivered Energy	36 joules

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.25%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
High Voltage Capacitor	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	1	0.25%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	8	AT 107 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
SAMPLE SIZE	380	340	280	220	170	140	120	100	50

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

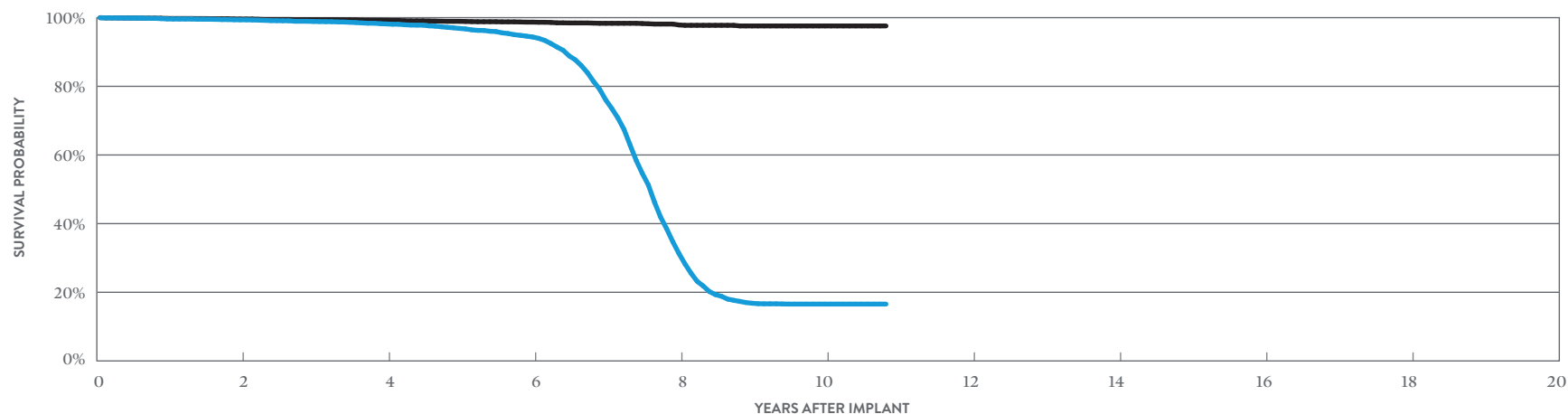
CUSTOMER REPORTED PERFORMANCE DATA

Atlas™ II VR

MODEL V-168

US Regulatory Approval	July 2006
Registered US Implants	10,605
Estimated Active US Implants	1,137
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	1,708
Max. Delivered Energy	36 joules
Number of US Advisories (see pg. 330)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	4	0.04%	3	0.03%
Electrical Interconnect	2	0.02%	0	0.00%
Battery	10	0.09%	2	0.02%
High Voltage Capacitor	1	<0.01%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	1	<0.01%	4	0.04%
Possible Early Battery Depletion	10	0.09%	5	0.05%
Other	10	0.09%	5	0.05%
Total	38	0.36%	19	0.18%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 130 MONTHS
SURVIVAL PROBABILITY	99.36%	98.17%	94.37%	31.34%	16.56%	16.56%
± 1 STANDARD ERROR	0.08%	0.15%	0.29%	0.67%	0.49%	0.49%
SAMPLE SIZE	8,640	6,620	5,100	2,750	800	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 130 MONTHS
SURVIVAL PROBABILITY	99.60%	99.22%	98.70%	97.88%	97.59%	97.59%
± 1 STANDARD ERROR	0.06%	0.10%	0.14%	0.18%	0.26%	0.26%

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

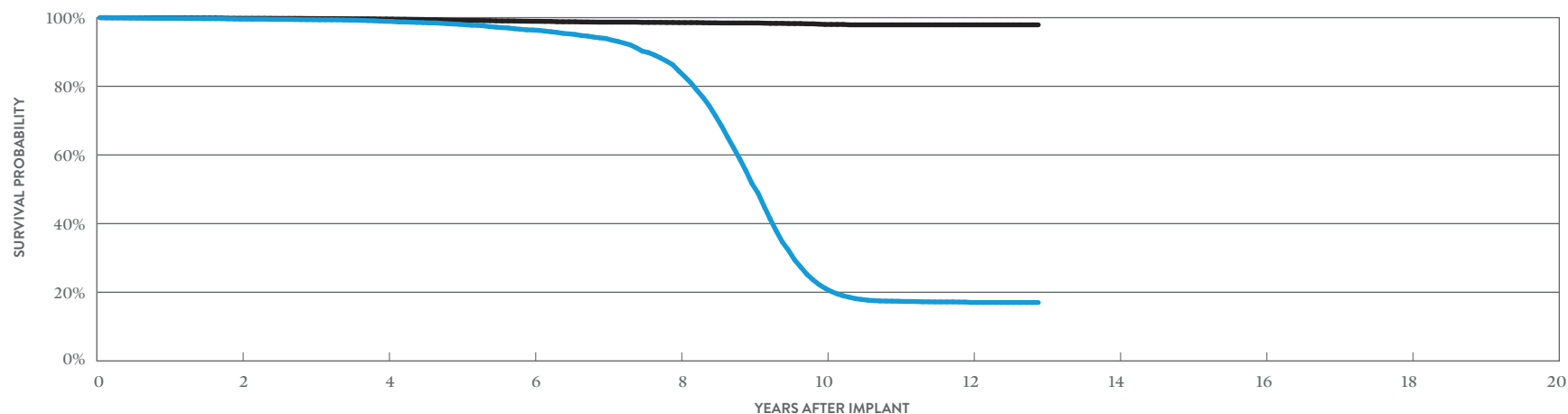
CUSTOMER REPORTED PERFORMANCE DATA

Atlas™ + VR

MODEL V-193

US Regulatory Approval	October 2003
Registered US Implants	20,793
Estimated Active US Implants	1,775
Estimated Longevity	(see table on page 152)
Normal Battery Depletion	2,659
Max. Delivered Energy	36 joules
Number of US Advisories (see pgs. 330, 331, 332)	Three

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	2	<0.01%
Electrical Interconnect	5	0.02%	1	<0.01%
Battery	9	0.04%	2	<0.01%
High Voltage Capacitor	2	<0.01%	1	<0.01%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	5	0.02%
Possible Early Battery Depletion	26	0.13%	5	0.02%
Other	13	0.06%	7	0.03%
Total	57	0.27%	24	0.12%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 155 MONTHS
SURVIVAL PROBABILITY	99.57%	98.94%	96.41%	84.45%	21.17%	17.02%	17.02%
± 1 STANDARD ERROR	0.05%	0.08%	0.17%	0.37%	0.45%	0.41%	0.41%
SAMPLE SIZE	16,890	12,910	9,670	6,990	3,110	1,010	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 155 MONTHS
SURVIVAL PROBABILITY	99.81%	99.60%	98.94%	98.56%	98.02%	97.91%	97.91%
± 1 STANDARD ERROR	0.03%	0.05%	0.09%	0.12%	0.17%	0.19%	0.19%

BATTERY LONGEVITY SUMMARY

**Single-Chamber
Implantable Cardioverter
Defibrillator (ICD) Devices**

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Battery Longevity (years)

MODELS	FAMILY	NO PACING	25% PACING	50% PACING	100% PACING
CD1411-36Q	Ellipse [™] VR*	11.1	10.6	10.1	9.4
CD1411-36C	Ellipse [™] VR*	11.1	10.6	10.1	9.4
CD1357-40Q	Fortify Assura [™] VR*	11.7	11.3	10.8	10.1
CD1357-40C	Fortify Assura [™] VR*	11.7	11.3	10.8	10.1
CD1257-40Q	Fortify Assura [™] VR*	11.7	11.3	10.8	10.1
CD1257-40	Fortify Assura [™] VR*	11.7	11.3	10.8	10.1
CD1311-36Q	Ellipse [™] VR*	11.1	10.6	10.1	9.4
CD1311-36	Ellipse [™] VR*	11.1	10.6	10.1	9.4
CD1231-40Q	Fortify [™] VR*	10.8	10.3	9.9	9.1
CD1231-40	Fortify [™] VR*	10.8	10.3	9.9	9.1
CD1211-36Q	Current [™] + VR**	8.4	8.0	7.6	7.0
CD1211-36	Current [™] + VR**	8.4	8.0	7.6	7.0
1207-36	Current [™] VR RF**	8.4	8.0	7.6	7.0
V-168	Atlas [™] II VR**	8.4	8.0	7.6	7.0
V-193	Atlas [™] + VR**	8.6	8.2	7.9	7.3

Pacing parameters: VVI, 2.5V, 0.5 ms, 60 ppm, 500 ohms

*Battery voltage range: 3.20-2.59. Three maximum charges per year.

**Battery voltage range: 3.20-2.45. Four maximum charges per year as well as monthly charging during the battery's mid-life voltage range.

SUMMARY INFORMATION

**Single-Chamber
Implantable Cardioverter
Defibrillator (ICD) Devices**

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices Survival Probability Summary

INCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
CDI411-36Q	Ellipse™ VR	99.80%	99.53%	99.41%	99.41%						
CDI411-36C	Ellipse™ VR	99.90%	99.90%	99.90%							
CDI357-40Q	Fortify Assura™ VR	99.88%	99.81%								
CDI357-40Q	Fortify Assura™ VR†	99.74%	99.33%	98.67%	98.53%						
CDI357-40C	Fortify Assura™ VR	99.91%	99.72%								
CDI357-40C	Fortify Assura™ VR†	99.80%	99.39%	98.97%	98.79%						
CDI257-40Q	Fortify Assura™ VR†	99.92%	99.77%	99.33%	98.87%	98.65%					
CDI257-40	Fortify Assura™ VR†	99.63%	99.52%	99.06%	98.70%	98.70%					
CDI311-36Q	Ellipse™ VR	99.51%	99.11%	98.07%	97.69%	97.69%					
CDI311-36	Ellipse™ VR	98.88%	98.29%	97.97%	97.61%	97.33%					
CDI231-40Q	Fortify™ VR†	99.75%	99.67%	99.18%	97.80%	96.13%	94.16%	93.30%			
CDI231-40	Fortify™ VR†	99.74%	99.67%	99.38%	97.80%	96.39%	95.29%	94.16%			
CDI211-36Q	Current™ + VR	99.61%	99.36%	98.83%	98.55%	97.32%	96.56%	95.96%	95.62%		
CDI211-36	Current™ + VR	99.71%	99.50%	99.08%	98.49%	97.96%	97.00%	95.17%	94.21%		
1207-36	Current™ VR RF	99.62%	99.28%	98.85%	98.42%	97.84%	96.35%	94.64%	91.78%	86.90%	
V-168	Atlas™ II VR	99.63%	99.36%	98.89%	98.17%	96.86%	94.37%	76.04%	31.34%	16.81%	16.56%
V-193	Atlas™ + VR	99.82%	99.57%	99.40%	98.94%	98.06%	96.41%	93.94%	84.45%	51.66%	21.17%

†Premature battery depletion advisory population.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices Survival Probability Summary

EXCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
CD1411-36Q	Ellipse™ VR	99.84%	99.76%	99.63%	99.63%						
CD1411-36C	Ellipse™ VR	99.90%	99.90%	99.90%							
CD1357-40Q	Fortify Assura™ VR	99.93%	99.93%								
CD1357-40Q	Fortify Assura™ VR†	99.77%	99.40%	98.74%	98.67%						
CD1357-40C	Fortify Assura™ VR	99.91%	99.91%								
CD1357-40C	Fortify Assura™ VR†	99.90%	99.59%	99.16%	98.99%						
CD1257-40Q	Fortify Assura™ VR†	99.96%	99.87%	99.57%	99.11%	98.89%					
CD1257-40	Fortify Assura™ VR†	99.63%	99.63%	99.17%	98.81%	98.81%					
CD1311-36Q	Ellipse™ VR	99.51%	99.11%	98.07%	97.69%	97.69%					
CD1311-36	Ellipse™ VR	98.88%	98.44%	98.12%	97.75%	97.48%					
CD1231-40Q	Fortify™ VR†	99.84%	99.79%	99.41%	98.27%	96.89%	95.30%	94.70%			
CD1231-40	Fortify™ VR†	99.97%	99.93%	99.71%	98.28%	97.07%	96.05%	95.06%			
CD1211-36Q	Current™ + VR	99.67%	99.41%	98.95%	98.87%	98.28%	98.20%	98.20%	98.08%		
CD1211-36	Current™ + VR	99.71%	99.64%	99.22%	98.97%	98.78%	98.66%	98.08%	97.92%		
1207-36	Current™ VR RF	99.73%	99.57%	99.19%	98.93%	98.60%	98.09%	97.99%	97.74%	97.09%	
V-168	Atlas™ II VR	99.77%	99.60%	99.44%	99.22%	98.93%	98.70%	98.36%	97.88%	97.59%	97.59%
V-193	Atlas™ + VR	99.95%	99.81%	99.74%	99.60%	99.19%	98.94%	98.69%	98.56%	98.45%	98.02%

†Premature battery depletion advisory population.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

US Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD1411-36Q	Ellipse [™] VR	12,672	1.80%	2	0.02%	0	0.00%	0	0.00%	6	0.05%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	8	0.06%
CD1411-36C	Ellipse [™] VR	5,095	2.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD1357-40Q	Fortify Assura [™] VR	10,558	1.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%
CD1357-40Q	Fortify Assura [™] VR†	10,215	5.90%	3	0.03%	1	<0.01%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	15	0.15%	2	0.02%	22	0.22%
CD1357-40C	Fortify Assura [™] VR	3,122	1.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD1357-40C	Fortify Assura [™] VR†	4,131	7.90%	3	0.07%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.02%	2	0.05%	0	0.00%	6	0.15%
CD1257-40Q	Fortify Assura [™] VR†	5,077	7.20%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	8	0.16%	1	0.02%	10	0.20%
CD1257-40	Fortify Assura [™] VR†	2,294	10.20%	1	0.04%	2	0.09%	1	0.04%	0	0.00%	0	0.00%	0	0.00%	2	0.09%	1	0.04%	7	0.31%
CD1311-36Q	Ellipse [™] VR	4,742	5.80%	1	0.02%	0	0.00%	0	0.00%	29	0.61%	1	0.02%	1	0.02%	0	0.00%	1	0.02%	33	0.70%
CD1311-36	Ellipse [™] VR	1,620	7.30%	2	0.12%	1	0.06%	0	0.00%	6	0.37%	0	0.00%	2	0.12%	0	0.00%	0	0.00%	11	0.68%
CD1231-40Q	Fortify [™] VR†	16,186	10.20%	7	0.04%	2	0.01%	15	0.09%	1	<0.01%	0	0.00%	0	0.00%	92	0.57%	6	0.04%	123	0.76%
CD1231-40	Fortify [™] VR†	6,782	11.50%	2	0.03%	0	0.00%	4	0.06%	6	0.09%	0	0.00%	0	0.00%	29	0.43%	3	0.04%	44	0.65%
CD1211-36Q	Current [™] + VR	4,431	9.00%	4	0.09%	0	0.00%	5	0.11%	1	0.02%	0	0.00%	0	0.00%	6	0.14%	2	0.05%	18	0.41%
CD1211-36	Current [™] + VR	3,637	9.20%	3	0.08%	2	0.05%	4	0.11%	2	0.05%	0	0.00%	0	0.00%	4	0.11%	1	0.03%	16	0.44%
1207-36	Current [™] VR RF	13,287	11.70%	6	0.05%	10	0.08%	10	0.08%	1	<0.01%	0	0.00%	0	0.00%	13	0.10%	9	0.07%	49	0.37%
V-168	Atlas [™] II VR	10,605	26.60%	4	0.04%	2	0.02%	10	0.09%	1	<0.01%	0	0.00%	1	<0.01%	10	0.09%	10	0.09%	38	0.36%
V-193	Atlas [™] + VR	20,793	24.00%	2	<0.01%	5	0.02%	9	0.04%	2	<0.01%	0	0.00%	0	0.00%	26	0.13%	13	0.06%	57	0.27%

Definitions of malfunction categories can be found on [pages 5-6](#).

†Premature battery depletion advisory population.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

US Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/ FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD1411-36Q	Ellipse™ VR	12,672	1.80%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%	2	0.02%	0	0.00%	0	0.00%	4	0.03%
CD1411-36C	Ellipse™ VR	5,095	2.10%	1	0.02%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.04%
CD1357-40Q	Fortify Assura™ VR	10,558	1.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%	0	0.00%	0	0.00%	2	0.02%
CD1357-40Q	Fortify Assura™ VR†	10,215	5.90%	8	0.08%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	8	0.08%	2	0.02%	19	0.19%
CD1357-40C	Fortify Assura™ VR	3,122	1.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.03%	0	0.00%	0	0.00%	1	0.03%
CD1357-40C	Fortify Assura™ VR†	4,131	7.90%	0	0.00%	0	0.00%	2	0.05%	0	0.00%	0	0.00%	0	0.00%	4	0.10%	0	0.00%	6	0.15%
CD1257-40Q	Fortify Assura™ VR†	5,077	7.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	7	0.14%	0	0.00%	7	0.14%
CD1257-40	Fortify Assura™ VR†	2,294	10.20%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	0	0.00%	0	0.00%	1	0.04%	1	0.04%	3	0.13%
CD1311-36Q	Ellipse™ VR	4,742	5.80%	1	0.02%	0	0.00%	0	0.00%	3	0.06%	0	0.00%	0	0.00%	0	0.00%	2	0.04%	6	0.13%
CD1311-36	Ellipse™ VR	1,620	7.30%	0	0.00%	0	0.00%	0	0.00%	2	0.12%	1	0.06%	1	0.06%	0	0.00%	0	0.00%	4	0.25%
CD1231-40Q	Fortify™ VR†	16,186	10.20%	4	0.02%	0	0.00%	21	0.13%	0	0.00%	0	0.00%	0	0.00%	40	0.25%	2	0.01%	67	0.41%
CD1231-40	Fortify™ VR†	6,782	11.50%	4	0.06%	0	0.00%	2	0.03%	0	0.00%	0	0.00%	1	0.01%	16	0.24%	3	0.04%	26	0.38%
CD1211-36Q	Current™ + VR	4,431	9.00%	3	0.07%	0	0.00%	3	0.07%	0	0.00%	0	0.00%	0	0.00%	1	0.02%	2	0.05%	9	0.20%
CD1211-36	Current™ + VR	3,637	9.20%	3	0.08%	0	0.00%	0	0.00%	0	0.00%	1	0.03%	0	0.00%	2	0.05%	0	0.00%	6	0.16%
1207-36	Current™ VR RF	13,287	11.70%	8	0.06%	0	0.00%	5	0.04%	1	<0.01%	4	0.03%	4	0.03%	14	0.11%	6	0.05%	42	0.32%
V-168	Atlas™ II VR	10,605	26.60%	3	0.03%	0	0.00%	2	0.02%	0	0.00%	0	0.00%	4	0.04%	5	0.05%	5	0.05%	19	0.18%
V-193	Atlas™ + VR	20,793	24.00%	2	<0.01%	1	<0.01%	2	<0.01%	1	<0.01%	1	<0.01%	5	0.02%	5	0.02%	7	0.03%	24	0.12%

Definitions of malfunction categories can be found on [pages 5-6](#).

†Premature battery depletion advisory population.

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Worldwide Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD1411-36Q	Ellipse™ VR	13,031	2.01%	2	0.02%	0	0.00%	0	0.00%	6	0.05%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	8	0.06%
CD1411-36C	Ellipse™ VR	5,229	2.62%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD1357-40Q	Fortify Assura™ VR	21,056	3.60%	3	0.01%	1	<0.01%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	15	0.07%	3	0.01%	23	0.11%
CD1357-40C	Fortify Assura™ VR	7,393	5.48%	3	0.04%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.01%	2	0.03%	0	0.00%	6	0.08%
CD1257-40Q	Fortify Assura™ VR	5,040	7.62%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	8	0.16%	1	0.02%	10	0.20%
CD1257-40	Fortify Assura™ VR	2,299	10.87%	1	0.04%	2	0.09%	1	0.04%	0	0.00%	0	0.00%	0	0.00%	2	0.09%	1	0.04%	7	0.30%
CD1311-36Q	Ellipse™ VR	4,893	6.23%	1	0.02%	0	0.00%	0	0.00%	29	0.59%	1	0.02%	1	0.02%	0	0.00%	1	0.02%	33	0.67%
CD1311-36	Ellipse™ VR	1,633	9.12%	2	0.12%	1	0.06%	0	0.00%	7	0.43%	0	0.00%	2	0.12%	0	0.00%	0	0.00%	12	0.73%
CD1231-40Q	Fortify™ VR†	17,561	9.92%	7	0.04%	2	0.01%	15	0.09%	1	<0.01%	0	0.00%	0	0.00%	101	0.58%	6	0.03%	132	0.75%
CD1231-40	Fortify™ VR†	8,054	10.52%	2	0.02%	0	0.00%	4	0.05%	6	0.07%	0	0.00%	0	0.00%	30	0.37%	3	0.04%	45	0.56%
CD1211-36Q	Current™ + VR	16,155	3.27%	8	0.05%	3	0.02%	8	0.05%	3	0.02%	0	0.00%	0	0.00%	8	0.05%	7	0.04%	37	0.23%
CD1211-36	Current™ + VR	14,877	2.89%	3	0.02%	4	0.03%	4	0.03%	5	0.03%	0	0.00%	0	0.00%	8	0.05%	6	0.04%	30	0.20%
1207-36	Current™ VR RF	24,846	8.22%	11	0.04%	30	0.12%	17	0.07%	1	<0.01%	0	0.00%	1	<0.01%	30	0.12%	11	0.04%	101	0.41%
V-168	Atlas™ II VR	23,946	14.75%	8	0.03%	5	0.02%	19	0.08%	1	<0.01%	0	0.00%	1	<0.01%	22	0.09%	21	0.09%	77	0.32%
V-193	Atlas™ + VR	39,596	15.47%	6	0.02%	9	0.02%	15	0.04%	5	0.01%	1	<0.01%	1	<0.01%	71	0.18%	32	0.08%	140	0.35%

Definitions of malfunction categories can be found on [pages 5-6](#).

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Worldwide Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD1411-36Q	Ellipse™ VR	13,031	2.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%	2	0.02%	0	0.00%	0	0.00%	4	0.03%
CD1411-36C	Ellipse™ VR	5,229	2.62%	1	0.02%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.04%
CD1357-40Q	Fortify Assura™ VR	21,056	3.60%	8	0.04%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	1	<0.01%	8	0.04%	2	<0.01%	21	0.10%
CD1357-40C	Fortify Assura™ VR	7,393	5.48%	1	0.01%	0	0.00%	2	0.03%	0	0.00%	0	0.00%	1	0.01%	4	0.05%	0	0.00%	8	0.11%
CD1257-40Q	Fortify Assura™ VR	5,040	7.62%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	7	0.14%	0	0.00%	7	0.14%
CD1257-40	Fortify Assura™ VR	2,299	10.87%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	0	0.00%	0	0.00%	1	0.04%	1	0.04%	3	0.13%
CD1311-36Q	Ellipse™ VR	4,893	6.23%	1	0.02%	0	0.00%	0	0.00%	3	0.06%	0	0.00%	0	0.00%	0	0.00%	2	0.04%	6	0.12%
CD1311-36	Ellipse™ VR	1,633	9.12%	0	0.00%	0	0.00%	0	0.00%	2	0.12%	1	0.06%	1	0.06%	0	0.00%	0	0.00%	4	0.24%
CD1231-40Q	Fortify™ VR†	17,561	9.92%	5	0.03%	1	<0.01%	22	0.13%	0	0.00%	0	0.00%	0	0.00%	42	0.24%	2	0.01%	72	0.41%
CD1231-40	Fortify™ VR†	8,054	10.52%	4	0.05%	0	0.00%	2	0.02%	0	0.00%	0	0.00%	1	0.01%	16	0.20%	3	0.04%	26	0.32%
CD1211-36Q	Current™ + VR	16,155	3.27%	5	0.03%	0	0.00%	5	0.03%	0	0.00%	0	0.00%	0	0.00%	3	0.02%	4	0.02%	17	0.11%
CD1211-36	Current™ + VR	14,877	2.89%	4	0.03%	0	0.00%	2	0.01%	0	0.00%	1	<0.01%	0	0.00%	3	0.02%	0	0.00%	10	0.07%
1207-36	Current™ VR RF	24,846	8.22%	14	0.06%	3	0.01%	12	0.05%	1	<0.01%	13	0.05%	6	0.02%	20	0.08%	10	0.04%	79	0.32%
V-168	Atlas™ II VR	23,946	14.75%	4	0.02%	0	0.00%	6	0.03%	0	0.00%	0	0.00%	12	0.05%	10	0.04%	9	0.04%	41	0.17%
V-193	Atlas™ + VR	39,596	15.47%	4	0.01%	3	<0.01%	8	0.02%	1	<0.01%	1	<0.01%	14	0.04%	11	0.03%	13	0.03%	55	0.14%

Definitions of malfunction categories can be found on [pages 5-6](#).

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	INAPPROPRIATE SHOCK		LOSS OF TELEMETRY		PERICARDIAL EFFUSION		PREMATURE BATTERY DEPLETION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD1231-40Q	159	82	9,579	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
CD1211-36Q	363	166	22,028	1	0.28%	0	0.00%	0	0.00%	1	0.28%	0	0.00%	2	0.55%
1207-36	395	78	21,682	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

A list of of complications can be found on [page 12](#).

Single-Chamber Implantable Cardioverter Defibrillator (ICD) Devices

Actively Monitored Study Data Summary

MALFUNCTIONS WITH COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD1231-40Q	Fortify [™] VR	159	13.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	1.26%	0	0.00%	2	1.26%
CD1211-36Q	Current [™] + VR	363	8.30%	0	0.00%	0	0.00%	1	0.28%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.28%
1207-36	Current [™] VR RF	395	15.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

MALFUNCTIONS WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		HIGH VOLTAGE CAPACITOR		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
CD1231-40Q	Fortify [™] VR	159	13.20%	0	0.00%	0	0.00%	1	0.63%	0	0.00%	0	0.00%	0	0.00%	1	0.63%	0	0.00%	2	1.26%
CD1211-36Q	Current [™] + VR	363	8.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
1207-36	Current [™] VR RF	395	15.40%	1	0.25%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.25%

Definitions of malfunction categories can be found on [pages 5-6](#).

Defibrillation Leads

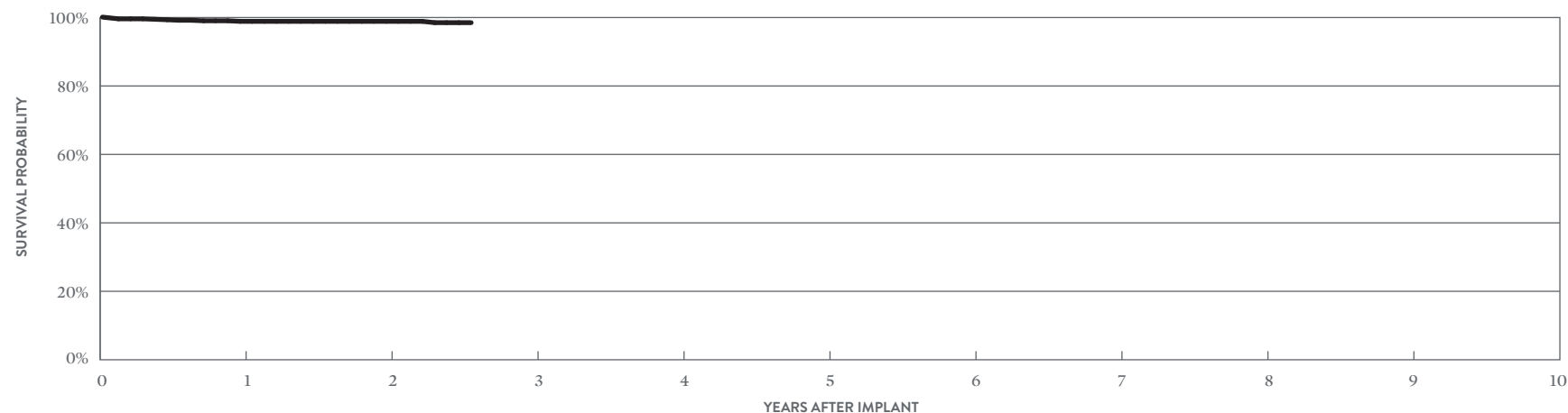
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Optisure™ DF4

MODEL LDA230Q

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	February 2014	Cardiac Perforation	0	0.00%	0	0.00%	Conductor Fracture	0	0.00%
Registered US Implants	857	Conductor Fracture	0	0.00%	0	0.00%	Clavicular Crush	0	0.00%
Estimated Active US Implants	674	Lead Dislodgement	1	0.12%	2	0.23%	In the Pocket	0	0.00%
Insulation	Optim™*	Failure to Capture	0	0.00%	3	0.35%	Intravascular	0	0.00%
Type and/or Fixation	Dual Coil, Active	Oversensing	0	0.00%	2	0.23%	Insulation Breach	1	0.12%
Polarity	Bipolar	Failure to Sense	0	0.00%	0	0.00%	Lead-to-Can Contact	0	0.00%
Steroid	Yes	Insulation Breach	0	0.00%	0	0.00%	Lead-to-Lead Contact	0	0.00%
Number of US Advisories (see pg. 342)	One	Abnormal Pacing Impedance	0	0.00%	0	0.00%	Clavicular Crush	1	0.12%
		Abnormal Defibrillation Impedance	0	0.00%	0	0.00%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	1	0.12%	0	0.00%	Other	0	0.00%
		Other	0	0.00%	0	0.00%	Crimps, Welds & Bonds	0	0.00%
		Total	2	0.23%	7	0.82%	Other	0	0.00%
		Total Returned for Analysis	0		4		Extrinsic Factors	6	0.70%
							Total	7	0.82%



YEAR	1	2	AT 31 MONTHS
SURVIVAL PROBABILITY	98.88%	98.88%	98.50%
± 1 STANDARD ERROR	0.36%	0.39%	0.55%
SAMPLE SIZE	720	460	210

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

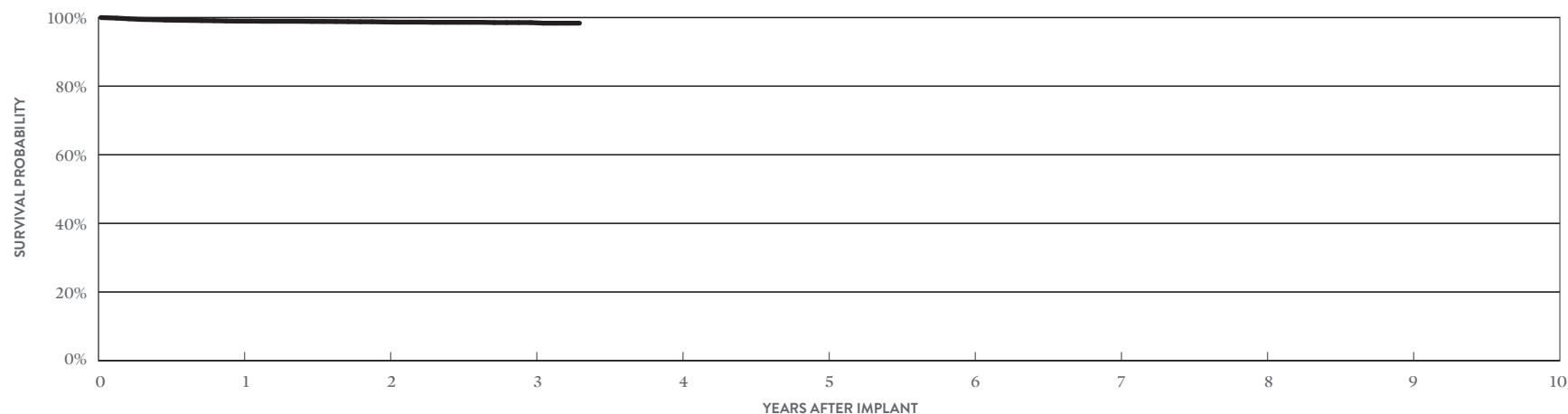
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Optisure™ DF4

MODEL LDA220Q

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	February 2014	Cardiac Perforation	4	0.06%	3	0.04%	Conductor Fracture	0	0.00%
Registered US Implants	7,163	Conductor Fracture	0	0.00%	2	0.03%	Clavicular Crush	0	0.00%
Estimated Active US Implants	5,736	Lead Dislodgement	23	0.32%	40	0.56%	In the Pocket	0	0.00%
Insulation	Optim™*	Failure to Capture	8	0.11%	24	0.34%	Intravascular	0	0.00%
Type and/or Fixation	Dual Coil, Active	Oversensing	3	0.04%	15	0.21%	Insulation Breach	2	0.03%
Polarity	Bipolar	Failure to Sense	1	0.01%	1	0.01%	Lead-to-Can Contact	1	0.01%
Steroid	Yes	Insulation Breach	0	0.00%	1	0.01%	Lead-to-Lead Contact	0	0.00%
Number of US Advisories (see pg. 342)	One	Abnormal Pacing Impedance	0	0.00%	0	0.00%	Clavicular Crush	0	0.00%
		Abnormal Defibrillation Impedance	4	0.06%	4	0.06%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	1	0.01%	0	0.00%	Other	1	0.01%
		Other	2	0.03%	2	0.03%	Crimps, Welds & Bonds	0	0.00%
		Total	46	0.64%	92	1.28%	Other	0	0.00%
		Total Returned for Analysis	18		34		Extrinsic Factors	36	0.50%
							Total	38	0.53%



YEAR	1	2	3	AT 40 MONTHS
SURVIVAL PROBABILITY	98.96%	98.71%	98.54%	98.37%
± 1 STANDARD ERROR	0.13%	0.15%	0.19%	0.25%
SAMPLE SIZE	5,970	3,720	1,660	280

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

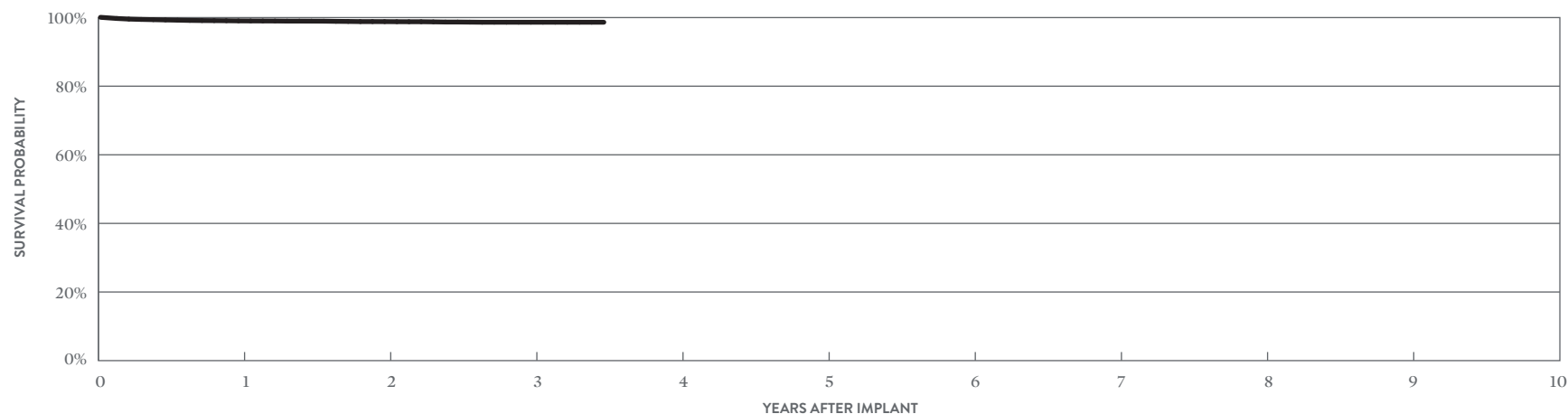
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Optisure™ DF4

MODEL LDA210Q

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	February 2014	Cardiac Perforation	26	0.12%	10	0.05%	Conductor Fracture	2	<0.01%
Registered US Implants	22,051	Conductor Fracture	1	<0.01%	2	<0.01%	Clavicular Crush	0	0.00%
Estimated Active US Implants	17,894	Lead Dislodgement	63	0.29%	107	0.49%	In the Pocket	1	<0.01%
Insulation	Optim™*	Failure to Capture	24	0.11%	42	0.19%	Intravascular	1	<0.01%
Type and/or Fixation	Single Coil, Active	Oversensing	10	0.05%	36	0.16%	Insulation Breach	3	0.01%
Polarity	Bipolar	Failure to Sense	6	0.03%	8	0.04%	Lead-to-Can Contact	1	<0.01%
Steroid	Yes	Insulation Breach	1	<0.01%	1	<0.01%	Lead-to-Lead Contact	1	<0.01%
Number of US Advisories	None	Abnormal Pacing Impedance	1	<0.01%	2	<0.01%	Clavicular Crush	0	0.00%
		Abnormal Defibrillation Impedance	5	0.02%	10	0.05%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	0	0.00%	2	<0.01%	Other	1	<0.01%
		Other	8	0.04%	9	0.04%	Crimps, Welds & Bonds	0	0.00%
		Total	145	0.66%	229	1.04%	Other	1	<0.01%
		Total Returned for Analysis	47		83		Extrinsic Factors	88	0.40%
							Total	94	0.43%



YEAR	1	2	3	at 42 months
SURVIVAL PROBABILITY	99.04%	98.82%	98.64%	98.64%
± 1 STANDARD ERROR	0.07%	0.09%	0.11%	0.11%
SAMPLE SIZE	17,720	10,050	4,090	320

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

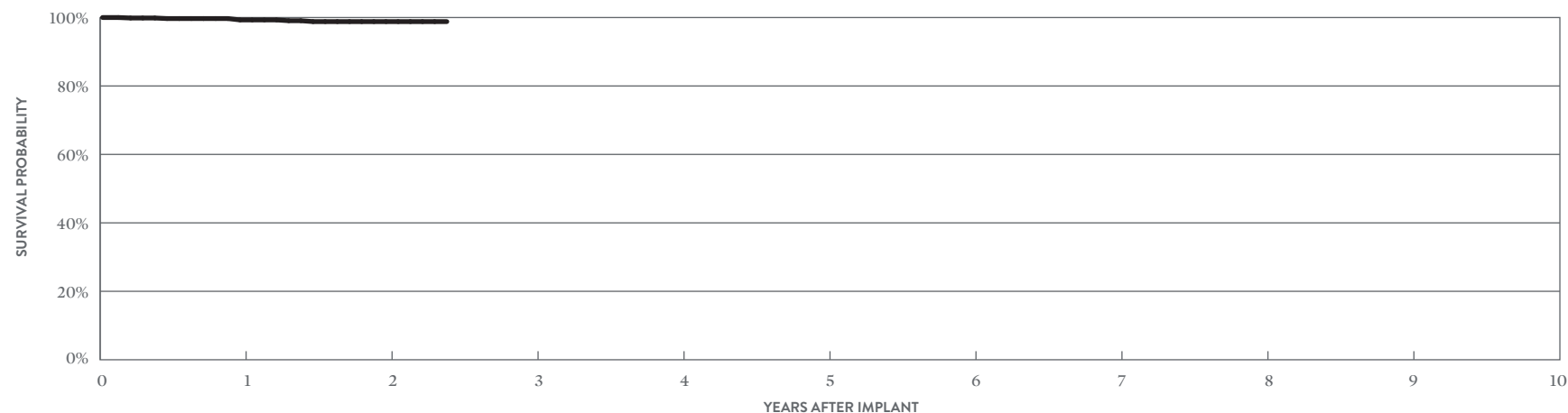
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Optisure™ DF4

MODEL LDA210

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	February 2014	Cardiac Perforation	0	0.00%	0	0.00%	Conductor Fracture	0	0.00%
Registered US Implants	797	Conductor Fracture	0	0.00%	1	0.13%	Clavicular Crush	0	0.00%
Estimated Active US Implants	656	Lead Dislodgement	2	0.25%	1	0.13%	In the Pocket	0	0.00%
Insulation	Optim™*	Failure to Capture	0	0.00%	2	0.25%	Intravascular	0	0.00%
Type and/or Fixation	Single Coil, Active	Oversensing	0	0.00%	2	0.25%	Insulation Breach	0	0.00%
Polarity	Bipolar	Failure to Sense	0	0.00%	0	0.00%	Lead-to-Can Contact	0	0.00%
Steroid	Yes	Insulation Breach	0	0.00%	0	0.00%	Lead-to-Lead Contact	0	0.00%
Number of US Advisories	None	Abnormal Pacing Impedance	0	0.00%	1	0.13%	Clavicular Crush	0	0.00%
		Abnormal Defibrillation Impedance	0	0.00%	0	0.00%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	0	0.00%	0	0.00%	Other	0	0.00%
		Other	0	0.00%	0	0.00%	Crimps, Welds & Bonds	0	0.00%
		Total	2	0.25%	7	0.88%	Other	0	0.00%
		Total Returned for Analysis	1		2		Extrinsic Factors	4	0.50%
							Total	4	0.50%



YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	99.30%	98.81%	98.81%
± 1 STANDARD ERROR	0.21%	0.49%	0.49%
SAMPLE SIZE	640	390	220

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

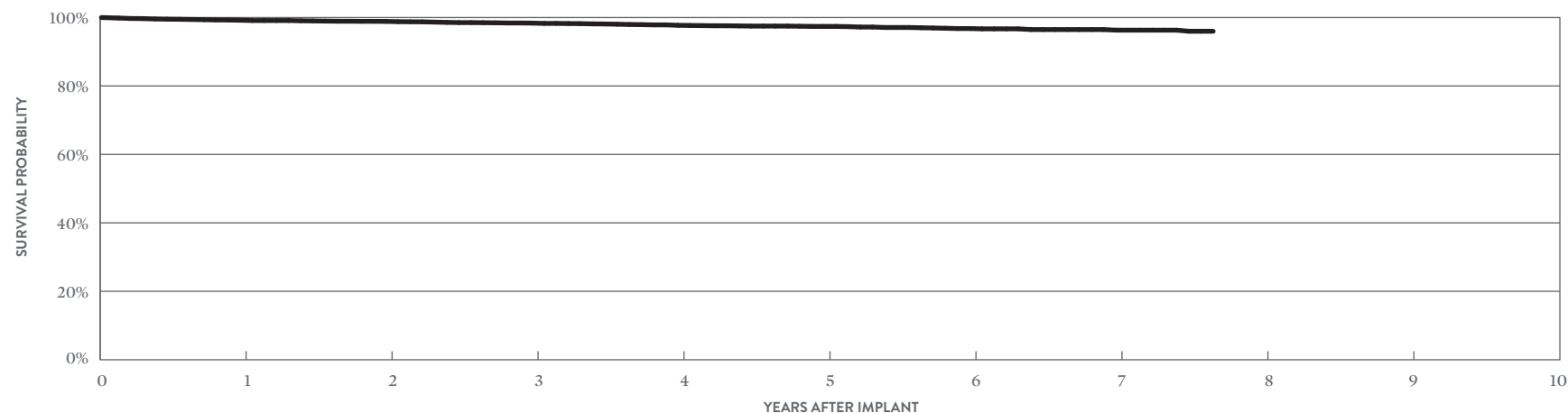
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Durata™ DF4

MODELS 7170Q & 7171Q

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE		QTY	RATE	
US Regulatory Approval	July 2009	Cardiac Perforation	6	0.10%	5	0.08%	Conductor Fracture	2	0.03%
Registered US Implants	6,170	Conductor Fracture	1	0.02%	9	0.15%	Clavicular Crush	0	0.00%
Estimated Active US Implants	3,661	Lead Dislodgement	15	0.24%	24	0.39%	In the Pocket	1	0.02%
Insulation	Optim™*	Failure to Capture	9	0.15%	48	0.78%	Intravascular	1	0.02%
Type and/or Fixation	Dual Coil, Passive	Oversensing	3	0.05%	27	0.44%	Insulation Breach	5	0.08%
Polarity	Bipolar	Failure to Sense	0	0.00%	0	0.00%	Lead-to-Can Contact	3	0.05%
Steroid	Yes	Insulation Breach	0	0.00%	2	0.03%	Lead-to-Lead Contact	2	0.03%
Number of US Advisories	None	Abnormal Pacing Impedance	1	0.02%	12	0.19%	Clavicular Crush	0	0.00%
		Abnormal Defibrillation Impedance	0	0.00%	11	0.18%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	0	0.00%	0	0.00%	Other	0	0.00%
		Other	1	0.02%	0	0.00%	Crimps, Welds & Bonds	0	0.00%
		Total	36	0.58%	138	2.24%	Other	0	0.00%
		Total Returned for Analysis	16		39		Extrinsic Factors	37	0.60%
							Total	44	0.71%



YEAR	1	2	3	4	5	6	7	AT 92 MONTHS
SURVIVAL PROBABILITY	99.21%	98.87%	98.37%	97.73%	97.40%	96.78%	96.31%	95.99%
± 1 STANDARD ERROR	0.12%	0.15%	0.19%	0.23%	0.27%	0.34%	0.38%	0.52%
SAMPLE SIZE	5,460	4,330	3,520	2,770	2,040	1,410	830	220

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

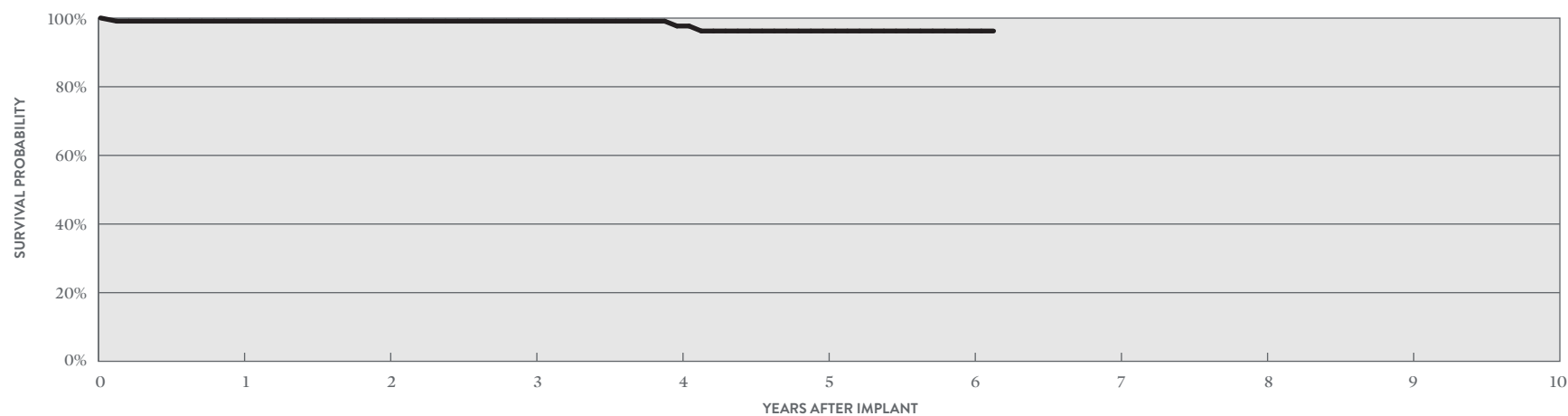
Durata™ DF4

MODELS 7170Q & 7171Q

US Regulatory Approval	July 2009
Number of Devices Enrolled in Study	116
Active Devices Enrolled in Study	55
Cumulative Months of Follow-up	6,504
Insulation	Optim™*
Type and/or Fixation	Dual Coil, Passive
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Pacing Impedance	1	0.86%
Conductor Fracture	1	0.86%
Lead Dislodgement	1	0.86%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	0	0.00%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	2	1.72%
Total	2	1.72%



YEAR	1	2	3	4	5	6	AT 74 MONTHS
SURVIVAL PROBABILITY	99.11%	99.11%	99.11%	97.69%	96.24%	96.24%	96.24%
± 1 STANDARD ERROR	0.89%	0.89%	0.89%	0.89%	2.17%	2.17%	2.17%
SAMPLE SIZE	110	100	90	70	60	60	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

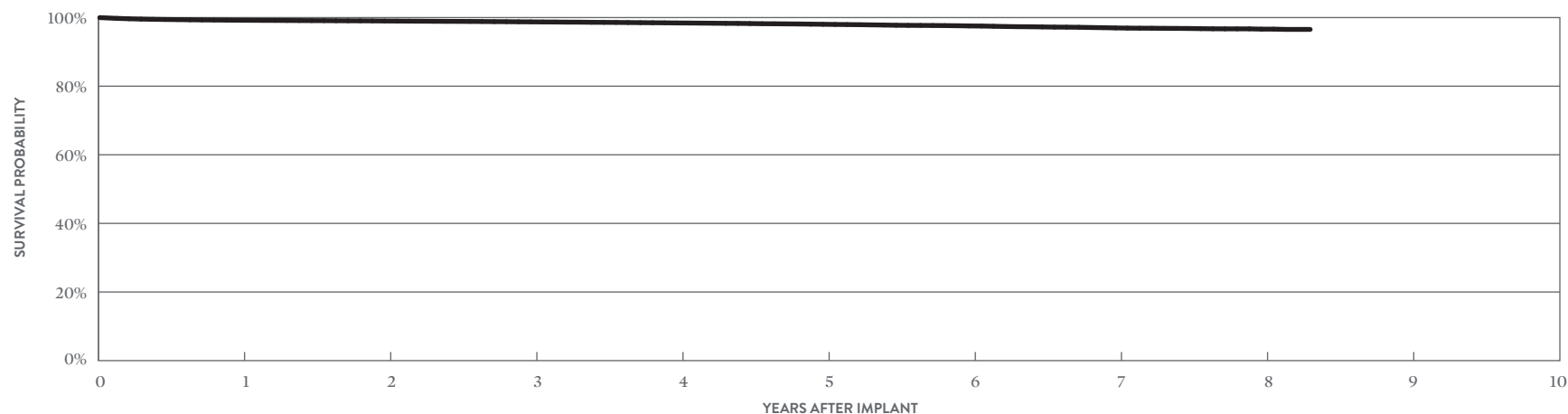
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Durata™ DF4

MODELS 7120Q & 7121Q

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	January 2009	Cardiac Perforation	81	0.06%	35	0.03%	Conductor Fracture	25	0.02%
Registered US Implants	125,371	Conductor Fracture	2	<0.01%	123	0.10%	Clavicular Crush	4	<0.01%
Estimated Active US Implants	72,386	Lead Dislodgement	232	0.19%	566	0.45%	In the Pocket	6	<0.01%
Insulation	Optim™*	Failure to Capture	100	0.08%	609	0.49%	Intravascular	15	0.01%
Type and/or Fixation	Dual Coil, Active	Oversensing	45	0.04%	499	0.40%	Insulation Breach	187	0.15%
Polarity	Bipolar	Failure to Sense	13	0.01%	69	0.06%	Lead-to-Can Contact	91	0.07%
Steroid	Yes	Insulation Breach	0	0.00%	26	0.02%	Lead-to-Lead Contact	23	0.02%
Number of US Advisories	None	Abnormal Pacing Impedance	5	<0.01%	101	0.08%	Clavicular Crush	26	0.02%
		Abnormal Defibrillation Impedance	9	<0.01%	249	0.20%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	3	<0.01%	7	<0.01%	Other	47	0.04%
		Other	34	0.03%	61	0.05%	Crimps, Welds & Bonds	2	<0.01%
		Total	524	0.42%	2345	1.87%	Other	36	0.03%
		Total Returned for Analysis	270		893		Extrinsic Factors	752	0.60%
							Total	1002	0.80%



YEAR	1	2	3	4	5	6	7	8	AT 100 MONTHS
SURVIVAL PROBABILITY	99.22%	99.00%	98.75%	98.43%	98.04%	97.56%	96.98%	96.63%	96.55%
± 1 STANDARD ERROR	0.03%	0.03%	0.03%	0.04%	0.05%	0.06%	0.08%	0.10%	0.14%
SAMPLE SIZE	114,350	95,620	80,390	65,150	49,910	35,260	21,190	8,180	470

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

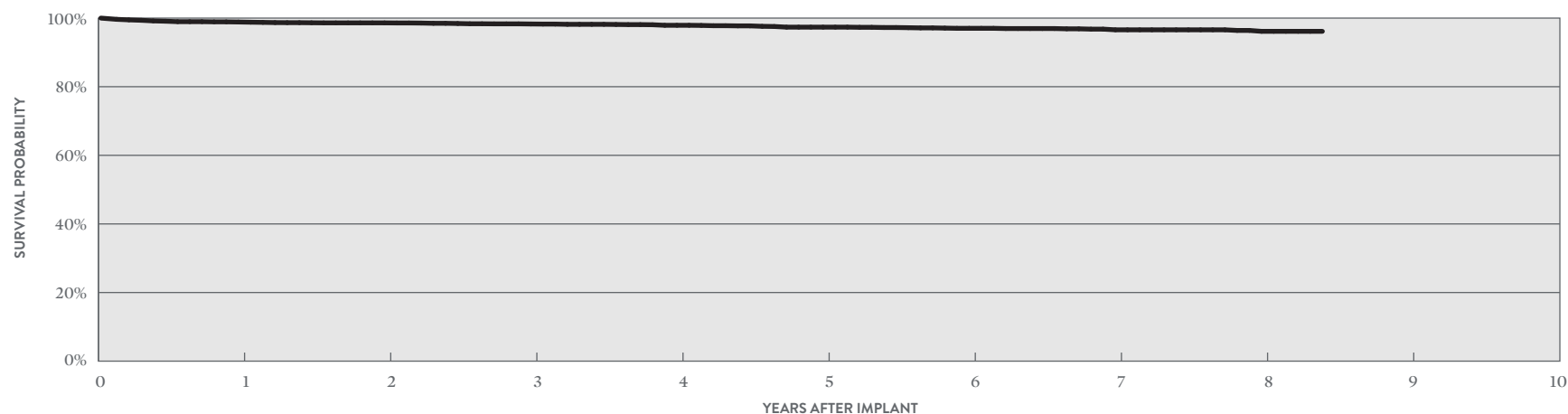
Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

Durata™ DF4

MODELS 7120Q & 7121Q

		QUALIFYING COMPLICATIONS			MALFUNCTIONS		
			QTY	RATE		QTY	RATE
US Regulatory Approval	January 2009	Abnormal Defibrillation Impedance	5	0.12%	Conductor Fracture	5	0.12%
Number of Devices Enrolled in Study	4,315	Abnormal Pacing Impedance	3	0.07%	Clavicular Crush	1	0.02%
Active Devices Enrolled in Study	1,891	Cardiac Perforation	1	0.02%	In the Pocket	2	0.05%
Cumulative Months of Follow-up	226,061	Conductor Fracture	15	0.35%	Intravascular	2	0.05%
Insulation	Optim™*	Failure to Capture	19	0.44%	Insulation Breach	6	0.14%
Type and/or Fixation	Dual Coil, Active	Failure to Sense	5	0.12%	Lead-to-Can Contact	3	0.07%
Polarity	Bipolar	Inappropriate Shock	4	0.09%	Lead-to-Lead Contact	2	0.05%
Steroid	Yes	Insulation Breach	3	0.07%	Clavicular Crush	0	0.00%
		Lead Dislodgement	39	0.90%	Externalized Conductors	0	0.00%
		Oversensing	7	0.16%	Other	1	0.02%
					Crimps, Welds & Bonds	0	0.00%
					Other	1	0.02%
					Extrinsic Factors	46	1.07%
					Total	58	1.34%



YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	98.86%	98.61%	98.28%	97.90%	97.36%	97.02%	96.58%	96.14%	96.14%
± 1 STANDARD ERROR	0.16%	0.18%	0.21%	0.24%	0.29%	0.32%	0.35%	0.42%	0.49%
SAMPLE SIZE	4,030	3,500	3,030	2,630	2,190	1,730	1,240	640	60

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

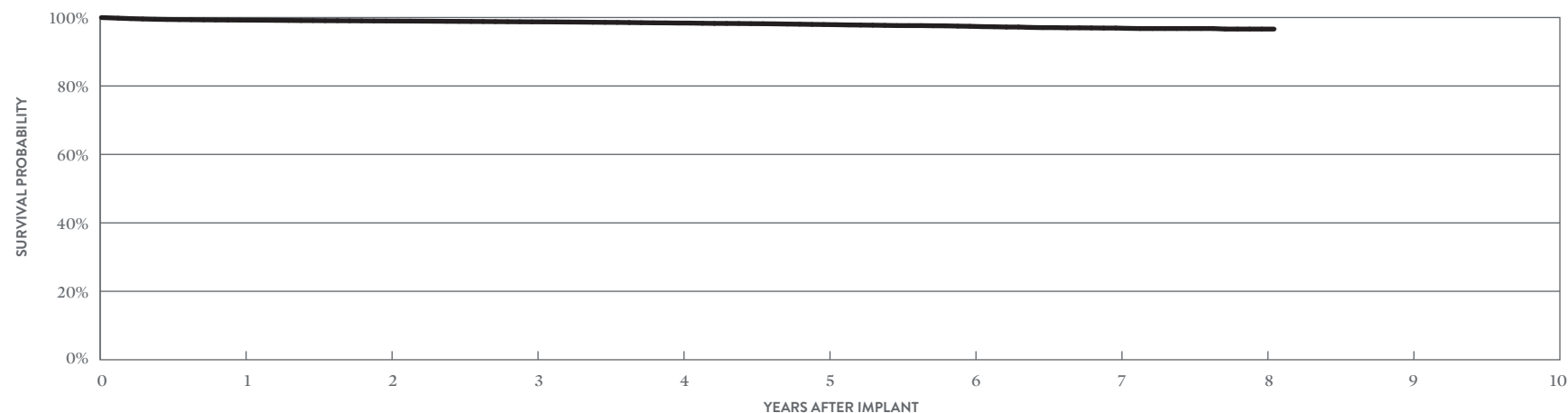
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Durata™ DF4

MODEL 7122Q

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	January 2009	Cardiac Perforation	109	0.13%	40	0.05%	Conductor Fracture	11	0.01%
Registered US Implants	85,535	Conductor Fracture	3	<0.01%	41	0.05%	Clavicular Crush	0	0.00%
Estimated Active US Implants	56,595	Lead Dislodgement	185	0.22%	358	0.42%	In the Pocket	7	<0.01%
Insulation	Optim™*	Failure to Capture	81	0.09%	262	0.31%	Intravascular	4	<0.01%
Type and/or Fixation	Single Coil, Active	Oversensing	25	0.03%	238	0.28%	Insulation Breach	87	0.10%
Polarity	Bipolar	Failure to Sense	7	<0.01%	32	0.04%	Lead-to-Can Contact	45	0.05%
Steroid	Yes	Insulation Breach	0	0.00%	14	0.02%	Lead-to-Lead Contact	11	0.01%
Number of US Advisories	None	Abnormal Pacing Impedance	5	<0.01%	44	0.05%	Clavicular Crush	11	0.01%
		Abnormal Defibrillation Impedance	7	<0.01%	69	0.08%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	3	<0.01%	8	<0.01%	Other	20	0.02%
		Other	33	0.04%	31	0.04%	Crimps, Welds & Bonds	0	0.00%
		Total	458	0.54%	1137	1.33%	Other	15	0.02%
		Total Returned for Analysis	205		517		Extrinsic Factors	475	0.56%
							Total	588	0.69%



YEAR	1	2	3	4	5	6	7	8	AT 97 MONTHS
SURVIVAL PROBABILITY	99.25%	98.99%	98.74%	98.39%	97.95%	97.44%	96.92%	96.63%	96.63%
± 1 STANDARD ERROR	0.03%	0.04%	0.04%	0.06%	0.08%	0.11%	0.16%	0.23%	0.23%
SAMPLE SIZE	74,000	54,480	39,520	26,210	15,450	8,320	3,950	1,260	240

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

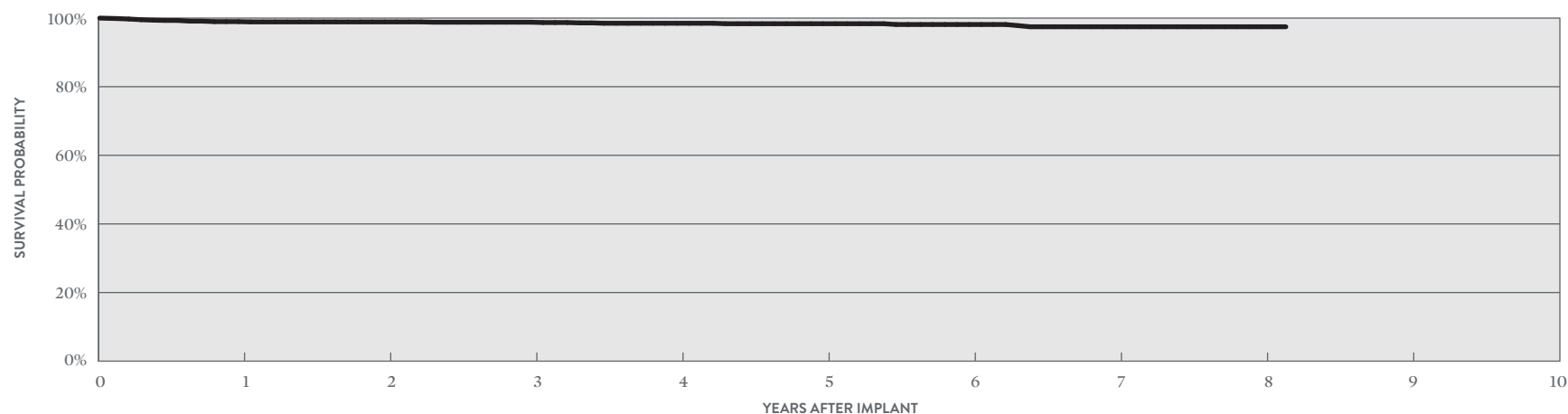
Durata™ DF4

MODEL 7122Q

US Regulatory Approval	January 2009
Number of Devices Enrolled in Study	1,542
Active Devices Enrolled in Study	755
Cumulative Months of Follow-up	73,384
Insulation	Optim™*
Type and/or Fixation	Single Coil, Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Defibrillation Impedance	2	0.13%
Conductor Fracture	4	0.26%
Failure to Capture	7	0.45%
Failure to Sense	1	0.06%
Lead Dislodgement	7	0.45%
Oversensing	1	0.06%
Pericardial Effusion	2	0.13%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	2	0.13%
Clavicular Crush	1	0.06%
In the Pocket	1	0.06%
Intravascular	0	0.00%
Insulation Breach	5	0.32%
Lead-to-Can Contact	4	0.26%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	1	0.06%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	14	0.91%
Total	21	1.36%



YEAR	1	2	3	4	5	6	7	8	AT 98 MONTHS
SURVIVAL PROBABILITY	98.97%	98.89%	98.80%	98.49%	98.36%	98.13%	97.47%	97.47%	97.47%
± 1 STANDARD ERROR	0.27%	0.28%	0.29%	0.34%	0.36%	0.43%	0.63%	0.63%	0.63%
SAMPLE SIZE	1,440	1,240	1,070	920	680	440	290	130	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

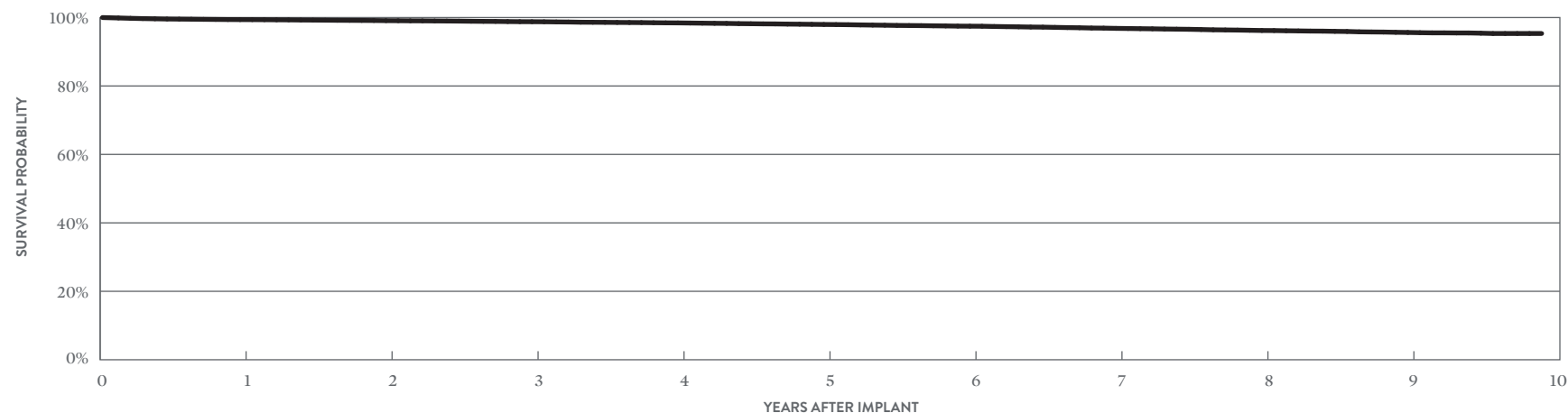
Durata™

MODELS 7120 & 7121

US Regulatory Approval	September 2007
Registered US Implants	59,761
Estimated Active US Implants	26,233
Insulation	Optim™*
Type and/or Fixation	Dual Coil, Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	40	0.07%	16	0.03%
Conductor Fracture	1	<0.01%	128	0.21%
Lead Dislodgement	69	0.12%	183	0.31%
Failure to Capture	24	0.04%	314	0.53%
Oversensing	49	0.08%	548	0.92%
Failure to Sense	5	<0.01%	63	0.11%
Insulation Breach	0	0.00%	61	0.10%
Abnormal Pacing Impedance	1	<0.01%	168	0.28%
Abnormal Defibrillation Impedance	19	0.03%	249	0.42%
Extracardiac Stimulation	0	0.00%	3	<0.01%
Other	21	0.04%	46	0.08%
Total	229	0.38%	1779	2.98%
Total Returned for Analysis	92		499	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	32	0.05%
Clavicular Crush	2	<0.01%
In the Pocket	22	0.04%
Intravascular	8	0.01%
Insulation Breach	136	0.23%
Lead-to-Can Contact	70	0.12%
Lead-to-Lead Contact	24	0.04%
Clavicular Crush	17	0.03%
Externalized Conductors	0	0.00%
Other	25	0.04%
Crimps, Welds & Bonds	1	<0.01%
Other	9	0.02%
Extrinsic Factors	393	0.66%
Total	571	0.96%



YEAR	1	2	3	4	5	6	7	8	9	AT 119 MONTHS
SURVIVAL PROBABILITY	99.39%	99.09%	98.78%	98.43%	97.97%	97.47%	96.83%	96.21%	95.63%	95.33%
± 1 STANDARD ERROR	0.03%	0.04%	0.05%	0.06%	0.07%	0.08%	0.09%	0.11%	0.13%	0.16%
SAMPLE SIZE	55,310	47,940	42,620	38,070	33,800	29,620	24,990	19,660	11,810	370

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

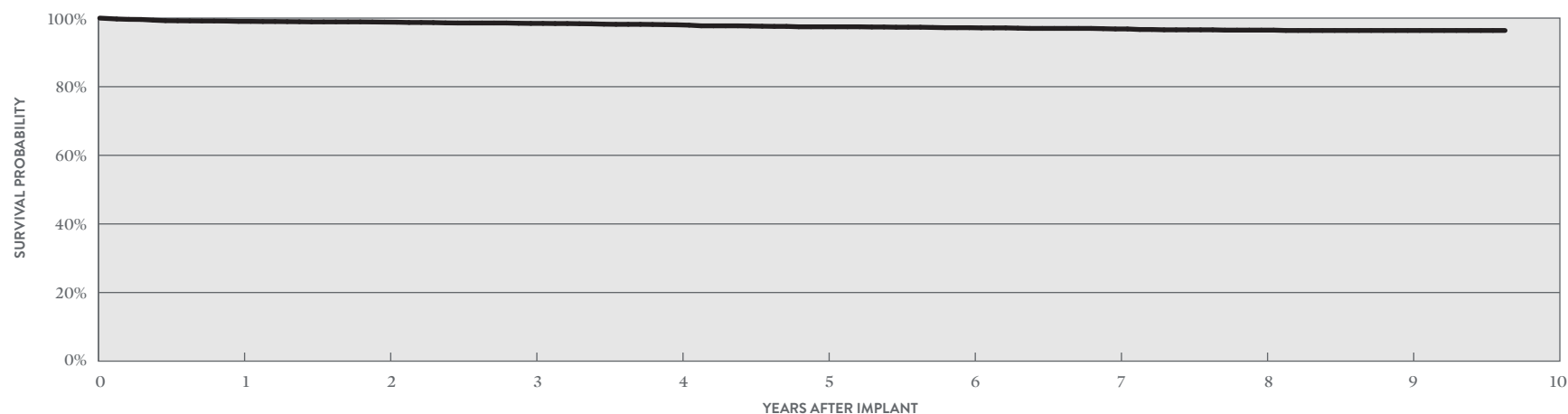
Durata™

MODELS 7120 & 7121

US Regulatory Approval	September 2007
Number of Devices Enrolled in Study	3,561
Active Devices Enrolled in Study	1,066
Cumulative Months of Follow-up	209,866
Insulation	Optim™*
Type and/or Fixation	Dual Coil, Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Defibrillation Impedance	4	0.11%
Abnormal Pacing Impedance	10	0.28%
Conductor Fracture	11	0.31%
Failure to Capture	13	0.37%
Failure to Sense	2	0.06%
Inappropriate Shock	2	0.06%
Insulation Breach	10	0.28%
Lead Dislodgement	20	0.56%
Oversensing	10	0.28%
Skin Erosion	1	0.03%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	1	0.03%
Clavicular Crush	0	0.00%
In the Pocket	1	0.03%
Intravascular	0	0.00%
Insulation Breach	12	0.34%
Lead-to-Can Contact	7	0.20%
Lead-to-Lead Contact	4	0.11%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	1	0.03%
Crimps, Welds & Bonds	0	0.00%
Other	1	0.03%
Extrinsic Factors	30	0.84%
Total	44	1.24%



YEAR	1	2	3	4	5	6	7	8	9	AT 116 MONTHS
SURVIVAL PROBABILITY	99.04%	98.83%	98.45%	98.04%	97.46%	97.20%	96.83%	96.49%	96.39%	96.39%
± 1 STANDARD ERROR	0.16%	0.18%	0.22%	0.25%	0.31%	0.34%	0.37%	0.41%	0.42%	0.42%
SAMPLE SIZE	3,360	2,950	2,560	2,180	1,820	1,540	1,310	1,100	680	70

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

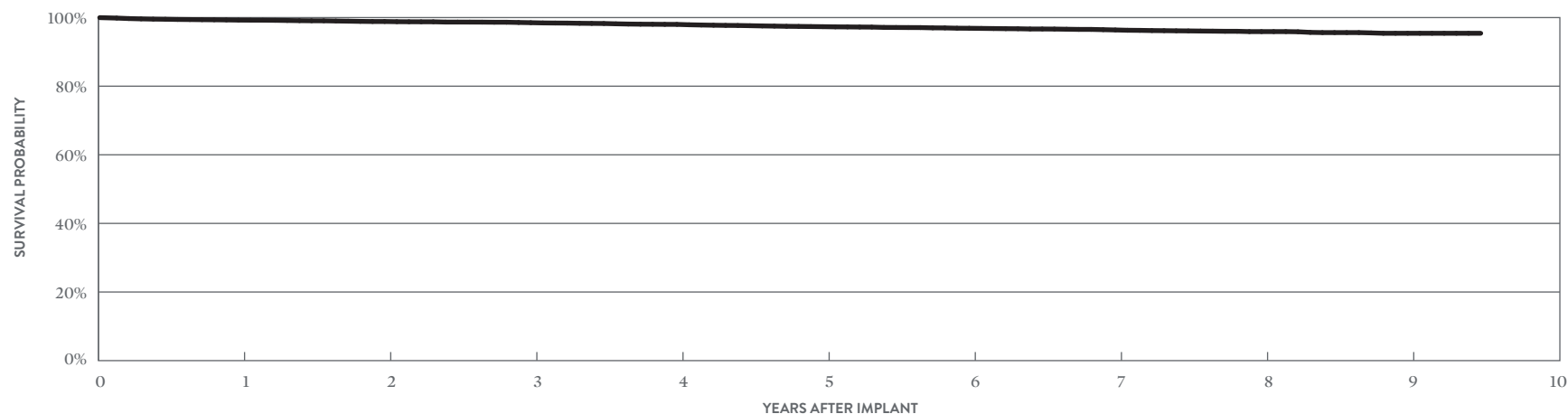
Durata™

MODEL 7122

US Regulatory Approval	September 2007
Registered US Implants	15,003
Estimated Active US Implants	7,673
Insulation	Optim™*
Type and/or Fixation	Single Coil, Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	11	0.07%	3	0.02%
Conductor Fracture	1	<0.01%	29	0.19%
Lead Dislodgement	20	0.13%	61	0.41%
Failure to Capture	18	0.12%	75	0.50%
Oversensing	12	0.08%	110	0.73%
Failure to Sense	0	0.00%	11	0.07%
Insulation Breach	0	0.00%	21	0.14%
Abnormal Pacing Impedance	3	0.02%	36	0.24%
Abnormal Defibrillation Impedance	1	<0.01%	28	0.19%
Extracardiac Stimulation	2	0.01%	2	0.01%
Other	4	0.03%	7	0.05%
Total	72	0.48%	383	2.55%
Total Returned for Analysis	32		170	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	16	0.11%
Clavicular Crush	1	<0.01%
In the Pocket	12	0.08%
Intravascular	3	0.02%
Insulation Breach	57	0.38%
Lead-to-Can Contact	31	0.21%
Lead-to-Lead Contact	16	0.11%
Clavicular Crush	2	0.01%
Externalized Conductors	1	<0.01%
Other	7	0.05%
Crimps, Welds & Bonds	0	0.00%
Other	4	0.03%
Extrinsic Factors	126	0.84%
Total	203	1.35%



YEAR	1	2	3	4	5	6	7	8	9	AT 114 MONTHS
SURVIVAL PROBABILITY	99.26%	98.85%	98.51%	98.01%	97.34%	96.88%	96.39%	95.91%	95.42%	95.42%
± 1 STANDARD ERROR	0.07%	0.09%	0.11%	0.13%	0.16%	0.19%	0.21%	0.25%	0.32%	0.32%
SAMPLE SIZE	13,720	11,530	9,780	8,130	6,640	5,400	4,130	2,740	1,360	250

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

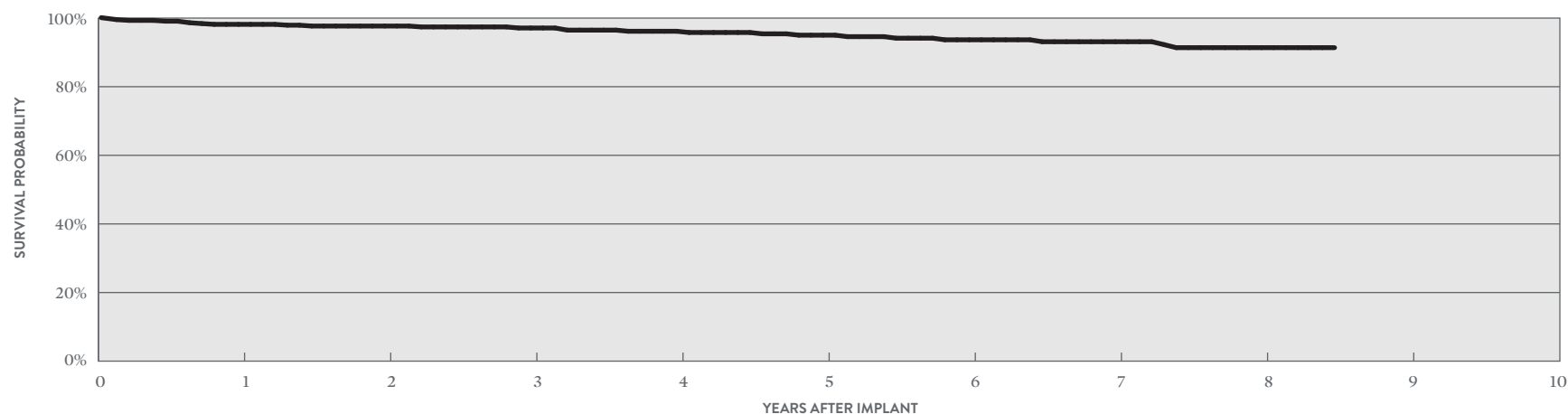
Durata™

MODEL 7122

US Regulatory Approval	September 2007
Number of Devices Enrolled in Study	450
Active Devices Enrolled in Study	176
Cumulative Months of Follow-up	26,801
Insulation	Optim™*
Type and/or Fixation	Single Coil, Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Defibrillation Impedance	1	0.22%
Abnormal Pacing Impedance	4	0.89%
Conductor Fracture	6	1.33%
Failure to Capture	5	1.11%
Failure to Sense	1	0.22%
Lead Dislodgement	5	1.11%
Oversensing	2	0.44%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	2	0.44%
Clavicular Crush	0	0.00%
In the Pocket	1	0.22%
Intravascular	1	0.22%
Insulation Breach	2	0.44%
Lead-to-Can Contact	1	0.22%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	1	0.22%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	8	1.78%
Total	12	2.67%



YEAR	1	2	3	4	5	6	7	8	AT 102 MONTHS
SURVIVAL PROBABILITY	98.17%	97.68%	97.11%	96.16%	95.02%	93.67%	93.11%	91.38%	91.38%
± 1 STANDARD ERROR	0.64%	0.72%	0.83%	0.98%	1.17%	1.39%	1.49%	1.90%	1.90%
SAMPLE SIZE	430	400	350	290	250	210	160	100	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

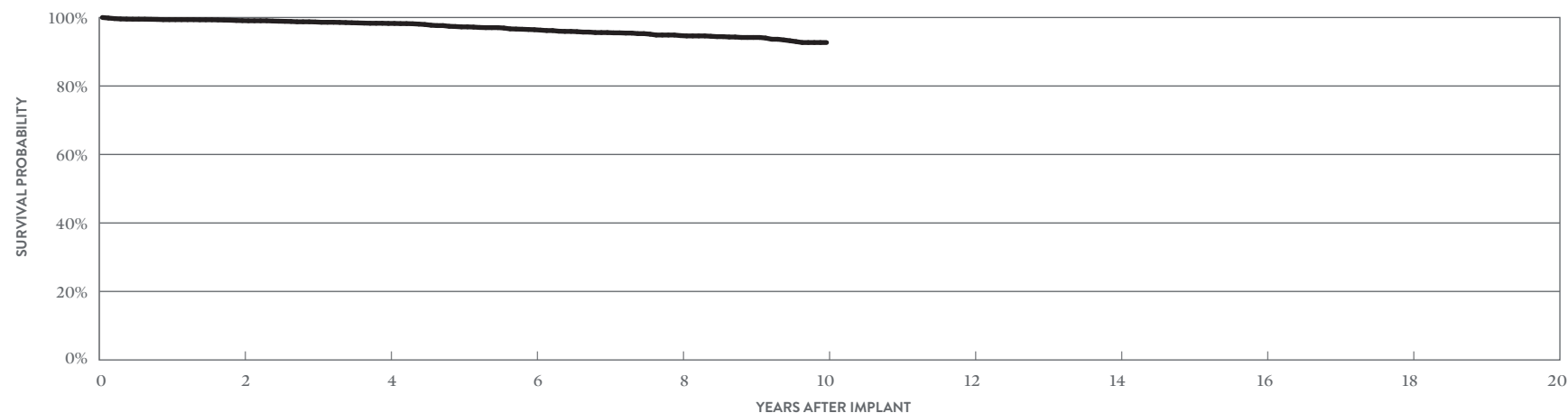
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Riata™ ST Optim™

MODELS 7070 & 7071

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	July 2006	Cardiac Perforation	3	0.09%	2	0.06%	Conductor Fracture	1	0.03%
Registered US Implants	3,311	Conductor Fracture	1	0.03%	17	0.51%	Clavicular Crush	0	0.00%
Estimated Active US Implants	1,311	Lead Dislodgement	3	0.09%	12	0.36%	In the Pocket	0	0.00%
Insulation	Optim™*	Failure to Capture	5	0.15%	34	1.03%	Intravascular	1	0.03%
Type and/or Fixation	Dual Coil, Passive	Oversensing	4	0.12%	49	1.48%	Insulation Breach	13	0.39%
Polarity	Bipolar	Failure to Sense	3	0.09%	2	0.06%	Lead-to-Can Contact	4	0.12%
Steroid	Yes	Insulation Breach	0	0.00%	6	0.18%	Lead-to-Lead Contact	3	0.09%
Number of US Advisories	None	Abnormal Pacing Impedance	0	0.00%	13	0.39%	Clavicular Crush	2	0.06%
		Abnormal Defibrillation Impedance	0	0.00%	14	0.42%	Externalized Conductors	1	0.03%
		Extracardiac Stimulation	0	0.00%	1	0.03%	Other	3	0.09%
		Other	0	0.00%	2	0.06%	Crimps, Welds & Bonds	0	0.00%
		Total	19	0.57%	152	4.59%	Other	0	0.00%
		Total Returned for Analysis	6		33		Extrinsic Factors	21	0.63%
							Total	35	1.06%



YEAR	2	4	6	8	10
SURVIVAL PROBABILITY	99.09%	98.27%	96.44%	94.71%	92.69%
± 1 STANDARD ERROR	0.17%	0.26%	0.40%	0.52%	0.81%
SAMPLE SIZE	2590	2070	1660	1190	210

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

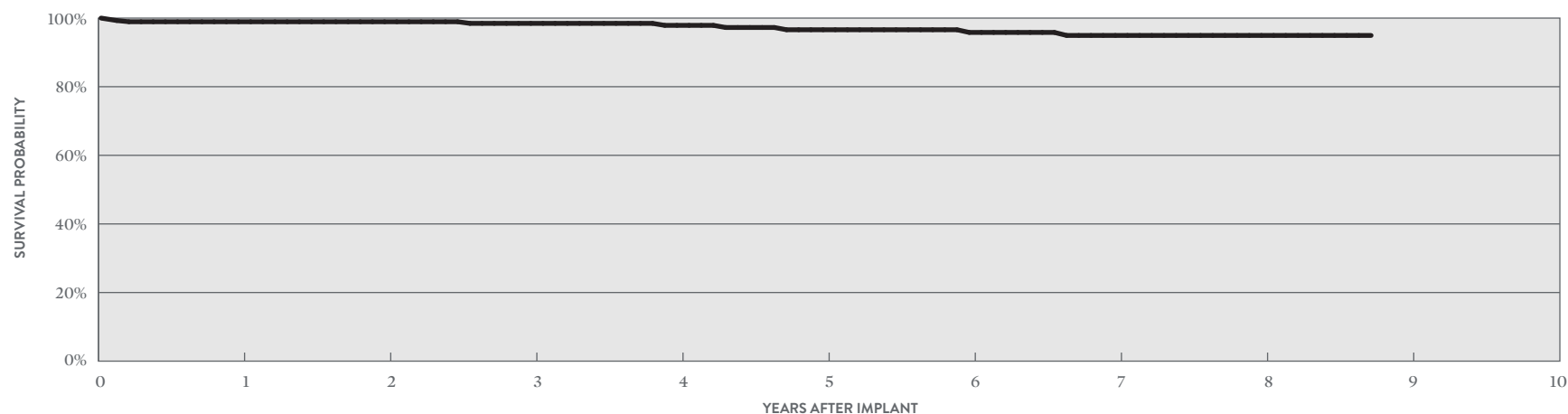
Riata™ ST Optim™

MODELS 7070 & 7071

US Regulatory Approval	July 2006
Number of Devices Enrolled in Study	288
Active Devices Enrolled in Study	81
Cumulative Months of Follow-up	17,458
Insulation	Optim™*
Type and/or Fixation	Dual Coil, Passive
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Defibrillation Impedance	1	0.35%
Abnormal Pacing Impedance	2	0.69%
Cardiac Perforation	1	0.35%
Conductor Fracture	2	0.69%
Failure to Capture	1	0.35%
Lead Dislodgement	1	0.35%
Oversensing	1	0.35%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	0	0.00%
Lead-to-Can Contact	0	0.00%
Lead-to-Lead Contact	0	0.00%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	1	0.35%
Total	1	0.35%



YEAR	1	2	3	4	5	6	7	8	AT 105 MONTHS
SURVIVAL PROBABILITY	98.94%	98.94%	98.46%	97.88%	96.60%	95.82%	94.94%	94.94%	94.94%
± 1 STANDARD ERROR	0.61%	0.61%	0.77%	0.96%	1.31%	1.31%	1.73%	1.73%	1.73%
SAMPLE SIZE	270	240	210	180	150	130	110	100	60

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

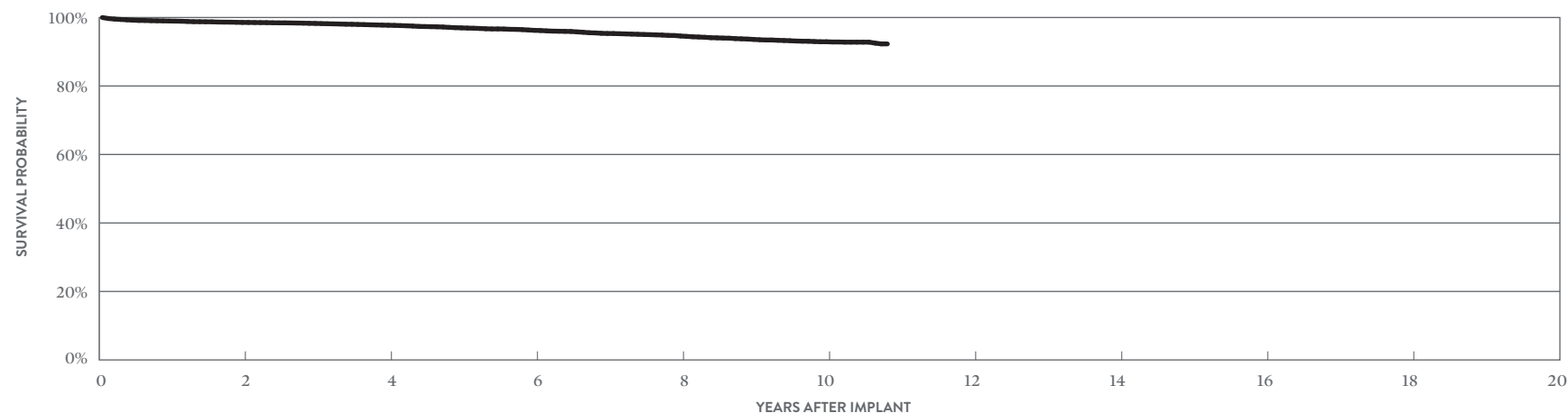
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Riata™ ST Optim™

MODELS 7020 & 7021

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)					
		QTY	RATE	QTY	RATE	MALFUNCTIONS	QTY	RATE	
US Regulatory Approval	July 2006	Cardiac Perforation	33	0.23%	16	0.11%	Conductor Fracture	10	0.07%
Registered US Implants	14,245	Conductor Fracture	0	0.00%	54	0.38%	Clavicular Crush	1	<0.01%
Estimated Active US Implants	5,126	Lead Dislodgement	27	0.19%	62	0.44%	In the Pocket	4	0.03%
Insulation	Optim™*	Failure to Capture	17	0.12%	147	1.03%	Intravascular	5	0.04%
Type and/or Fixation	Dual Coil, Active	Oversensing	19	0.13%	235	1.65%	Insulation Breach	50	0.35%
Polarity	Bipolar	Failure to Sense	8	0.06%	20	0.14%	Lead-to-Can Contact	21	0.15%
Steroid	Yes	Insulation Breach	0	0.00%	24	0.17%	Lead-to-Lead Contact	7	0.05%
Number of US Advisories	None	Abnormal Pacing Impedance	1	<0.01%	44	0.31%	Clavicular Crush	4	0.03%
		Abnormal Defibrillation Impedance	4	0.03%	91	0.64%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	3	0.02%	2	0.01%	Other	18	0.13%
		Other	0	0.00%	28	0.20%	Crimps, Welds & Bonds	0	0.00%
		Total	112	0.79%	723	5.08%	Other	0	0.00%
		Total Returned for Analysis	53		204		Extrinsic Factors	172	1.21%
							Total	232	1.63%



YEAR	2	4	6	8	10	AT 130 MONTHS
SURVIVAL PROBABILITY	98.56%	97.74%	96.25%	94.59%	92.92%	92.29%
± 1 STANDARD ERROR	0.10%	0.14%	0.19%	0.24%	0.30%	0.43%
SAMPLE SIZE	11,260	8,920	7,320	6,040	3,820	330

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

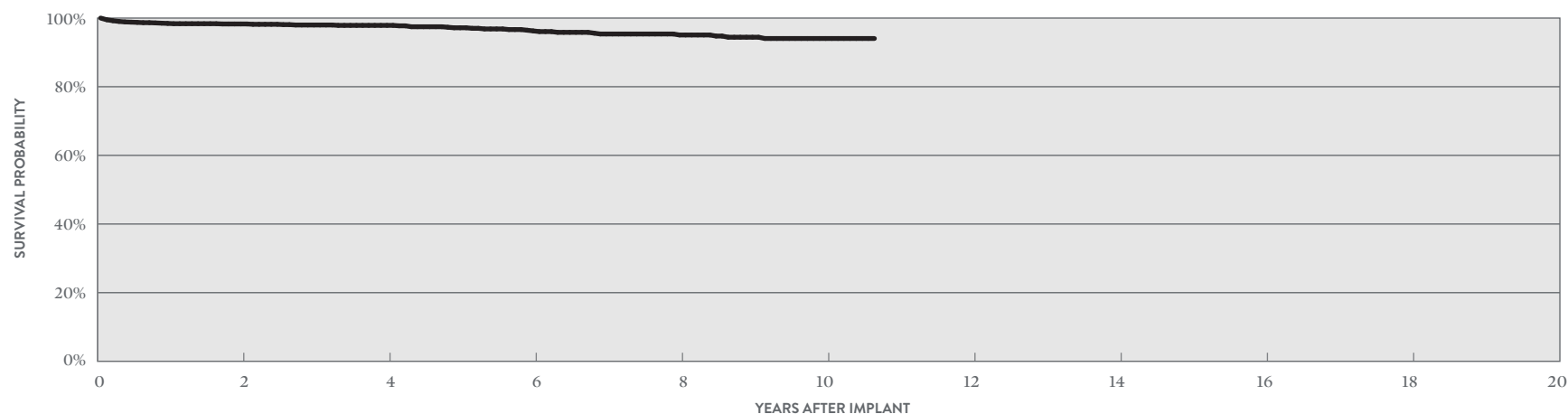
Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

Riata™ ST Optim™

MODELS 7020 & 7021

		QUALIFYING COMPLICATIONS			MALFUNCTIONS		
			QTY	RATE		QTY	RATE
US Regulatory Approval	July 2006	Abnormal Pacing Impedance	6	0.41%	Conductor Fracture	3	0.20%
Number of Devices Enrolled in Study	1,470	Cardiac Perforation	1	0.07%	Clavicular Crush	0	0.00%
Active Devices Enrolled in Study	258	Conductor Fracture	8	0.54%	In the Pocket	3	0.20%
Cumulative Months of Follow-up	83,312	Failure to Capture	14	0.95%	Intravascular	0	0.00%
Insulation	Optim™*	Failure to Sense	1	0.07%	Insulation Breach	3	0.20%
Type and/or Fixation	Dual Coil, Active	Insulation Breach	2	0.14%	Lead-to-Can Contact	1	0.07%
Polarity	Bipolar	Lead Dislodgement	9	0.61%	Lead-to-Lead Contact	0	0.00%
Steroid	Yes	Oversensing	4	0.27%	Clavicular Crush	2	0.14%
		Skin Erosion	1	0.07%	Externalized Conductors	0	0.00%
					Other	0	0.00%
					Crimps, Welds & Bonds	0	0.00%
					Other	0	0.00%
					Extrinsic Factors	14	0.95%
					Total	20	1.36%



YEAR	2	4	6	8	10	AT 128 MONTHS
SURVIVAL PROBABILITY	98.28%	97.87%	96.26%	95.07%	94.05%	94.05%
± 1 STANDARD ERROR	0.35%	0.40%	0.62%	0.78%	1.01%	1.01%
SAMPLE SIZE	1,180	840	550	360	210	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

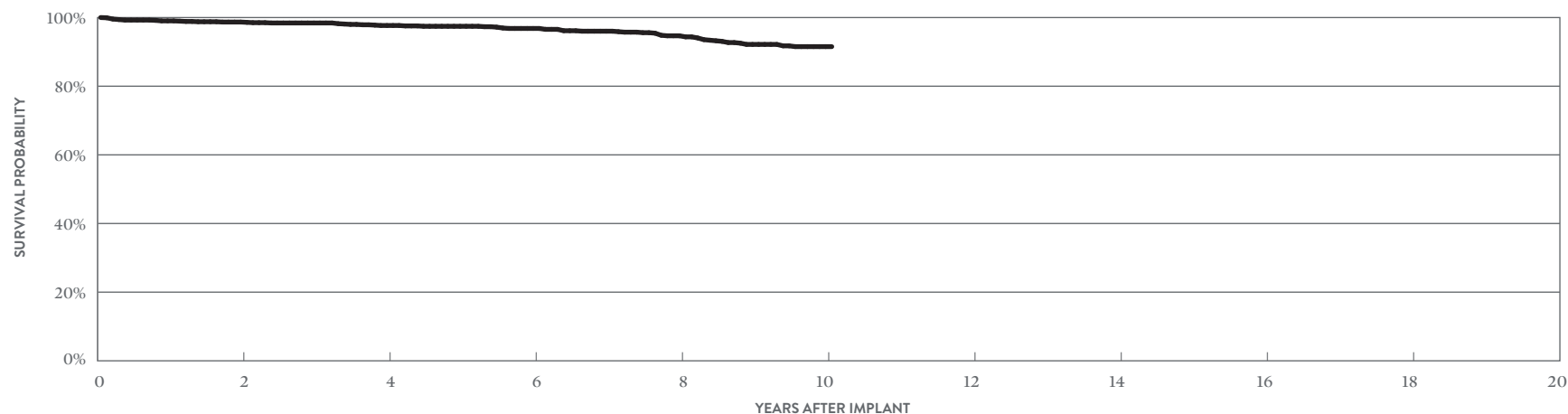
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Riata™ ST Optim™

MODEL 7022

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE		QTY	RATE	
US Regulatory Approval	July 2006	Cardiac Perforation	5	0.34%	3	0.20%	Conductor Fracture	3	0.20%
Registered US Implants	1,471	Conductor Fracture	0	0.00%	10	0.68%	Clavicular Crush	0	0.00%
Estimated Active US Implants	555	Lead Dislodgement	3	0.20%	11	0.75%	In the Pocket	2	0.14%
Insulation	Optim™*	Failure to Capture	1	0.07%	12	0.82%	Intravascular	1	0.07%
Type and/or Fixation	Single Coil, Active	Oversensing	0	0.00%	22	1.50%	Insulation Breach	7	0.48%
Polarity	Bipolar	Failure to Sense	0	0.00%	1	0.07%	Lead-to-Can Contact	6	0.41%
Steroid	Yes	Insulation Breach	0	0.00%	7	0.48%	Lead-to-Lead Contact	0	0.00%
Number of US Advisories	None	Abnormal Pacing Impedance	2	0.14%	4	0.27%	Clavicular Crush	0	0.00%
		Abnormal Defibrillation Impedance	0	0.00%	3	0.20%	Externalized Conductors	0	0.00%
		Extracardiac Stimulation	0	0.00%	1	0.07%	Other	1	0.07%
		Other	0	0.00%	1	0.07%	Crimps, Welds & Bonds	0	0.00%
		Total	11	0.75%	75	5.10%	Other	0	0.00%
		Total Returned for Analysis	4		26		Extrinsic Factors	21	1.43%
							Total	31	2.11%



YEAR	2	4	6	8	10	AT 121 MONTHS
SURVIVAL PROBABILITY	98.69%	97.68%	96.82%	94.68%	91.53%	91.53%
± 1 STANDARD ERROR	0.32%	0.45%	0.55%	0.77%	1.04%	1.04%
SAMPLE SIZE	1,170	930	780	640	370	230

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

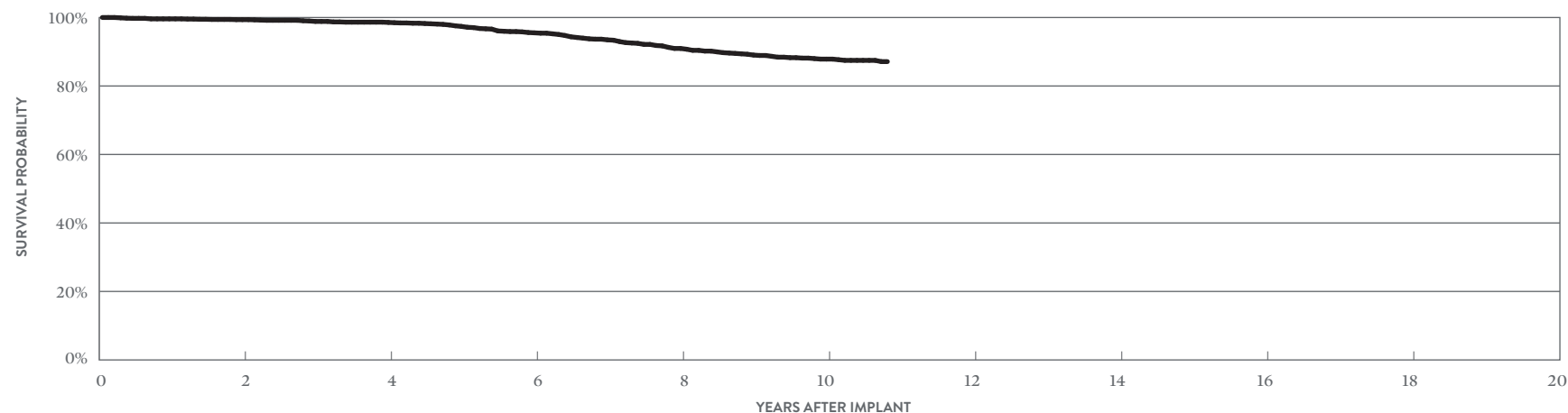
Riata™ ST

MODELS 7010 & 7011

US Regulatory Approval	March 2006
Registered US Implants	2,200
Estimated Active US Implants	700
Insulation	Silicone
Type and/or Fixation	Dual Coil, Active
Polarity	Integrated Bipolar
Steroid	Yes
Number of US Advisories (see pg. 343)	One

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	3	0.14%	3	0.14%
Conductor Fracture	0	0.00%	5	0.23%
Lead Dislodgement	1	0.05%	8	0.36%
Failure to Capture	2	0.09%	10	0.45%
Oversensing	2	0.09%	43	1.95%
Failure to Sense	1	0.05%	3	0.14%
Insulation Breach	0	0.00%	40	1.82%
Abnormal Pacing Impedance	1	0.05%	26	1.18%
Abnormal Defibrillation Impedance	0	0.00%	19	0.86%
Extracardiac Stimulation	0	0.00%	0	0.00%
Other	1	0.05%	3	0.14%
Total	11	0.50%	160	7.27%
Total Returned for Analysis	4		37	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	2	0.09%
Clavicular Crush	0	0.00%
In the Pocket	2	0.09%
Intravascular	0	0.00%
Insulation Breach	38	1.73%
Lead-to-Can Contact	12	0.55%
Lead-to-Lead Contact	17	0.77%
Clavicular Crush	1	0.05%
Externalized Conductors	2	0.09%
Other	6	0.27%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	11	0.50%
Total	51	2.32%



YEAR	2	4	6	8	10	AT 130 MONTHS
SURVIVAL PROBABILITY	99.37%	98.56%	95.52%	90.96%	87.85%	87.12%
± 1 STANDARD ERROR	0.18%	0.28%	0.57%	0.86%	1.03%	1.11%
SAMPLE SIZE	1,750	1,370	1,070	880	650	220

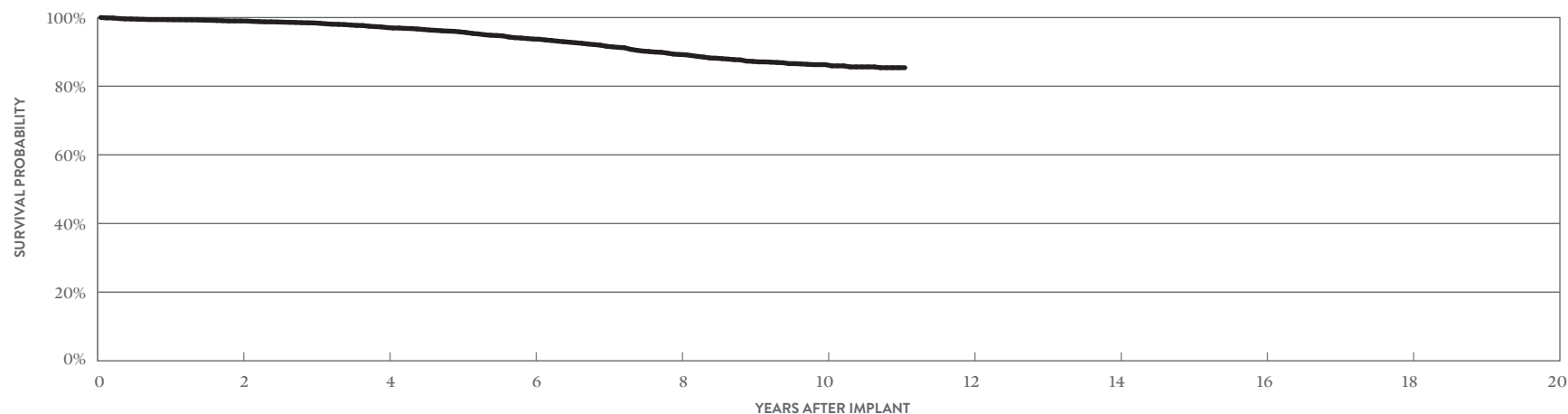
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Riata™ ST

MODELS 7040 & 7041

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	March 2006	Cardiac Perforation	4	0.10%	3	0.07%	Conductor Fracture	4	0.10%
Registered US Implants	4,055	Conductor Fracture	0	0.00%	34	0.84%	Clavicular Crush	0	0.00%
Estimated Active US Implants	1,307	Lead Dislodgement	5	0.12%	5	0.12%	In the Pocket	1	0.02%
Insulation	Silicone	Failure to Capture	1	0.02%	49	1.21%	Intravascular	3	0.07%
Type and/or Fixation	Dual Coil, Passive	Oversensing	3	0.07%	96	2.37%	Insulation Breach	55	1.36%
Polarity	Bipolar	Failure to Sense	0	0.00%	14	0.35%	Lead-to-Can Contact	28	0.69%
Steroid	Yes	Insulation Breach	0	0.00%	55	1.36%	Lead-to-Lead Contact	15	0.37%
Number of US Advisories (see pg. 343)	One	Abnormal Pacing Impedance	2	0.05%	19	0.47%	Clavicular Crush	0	0.00%
		Abnormal Defibrillation Impedance	0	0.00%	24	0.59%	Externalized Conductors	2	0.05%
		Extracardiac Stimulation	0	0.00%	1	0.02%	Other	10	0.25%
		Other	1	0.02%	7	0.17%	Crimps, Welds & Bonds	0	0.00%
		Total	16	0.39%	307	7.57%	Other	0	0.00%
		Total Returned for Analysis	3		68		Extrinsic Factors	29	0.72%
							Total	88	2.17%



YEAR	2	4	6	8	10	AT 133 MONTHS
SURVIVAL PROBABILITY	99.01%	97.08%	93.73%	89.23%	86.28%	85.39%
± 1 STANDARD ERROR	0.17%	0.30%	0.48%	0.67%	0.80%	0.87%
SAMPLE SIZE	3,240	2,540	1,990	1,580	940	210

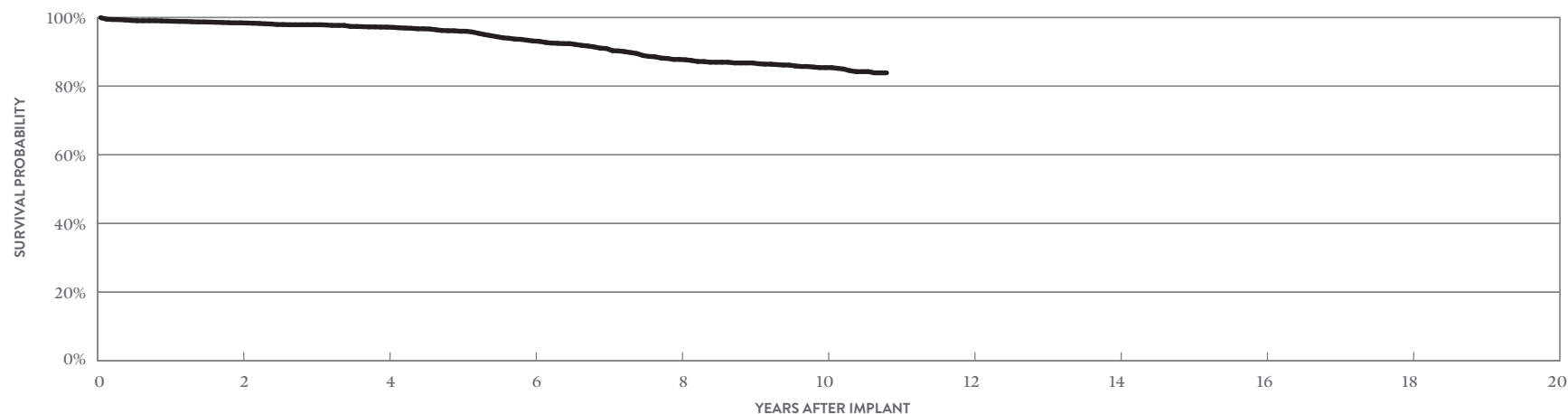
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Riata™ ST

MODEL 7002

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE	QTY	RATE		
US Regulatory Approval	June 2005	Cardiac Perforation	6	0.25%	5	0.21%	Conductor Fracture	5	0.21%
Registered US Implants	2,408	Conductor Fracture	0	0.00%	9	0.37%	Clavicular Crush	0	0.00%
Estimated Active US Implants	762	Lead Dislodgement	3	0.12%	9	0.37%	In the Pocket	2	0.08%
Insulation	Silicone	Failure to Capture	4	0.17%	22	0.91%	Intravascular	3	0.12%
Type and/or Fixation	Single Coil, Active	Oversensing	4	0.17%	62	2.57%	Insulation Breach	66	2.74%
Polarity	Bipolar	Failure to Sense	0	0.00%	2	0.08%	Lead-to-Can Contact	29	1.20%
Steroid	Yes	Insulation Breach	0	0.00%	69	2.87%	Lead-to-Lead Contact	17	0.71%
Number of US Advisories (see pg. 343)	One	Abnormal Pacing Impedance	2	0.08%	4	0.17%	Clavicular Crush	0	0.00%
		Abnormal Defibrillation Impedance	1	0.04%	9	0.37%	Externalized Conductors	9	0.37%
		Extracardiac Stimulation	0	0.00%	0	0.00%	Other	11	0.46%
		Other	1	0.04%	8	0.33%	Crimps, Welds & Bonds	0	0.00%
		Total	21	0.87%	199	8.26%	Other	0	0.00%
		Total Returned for Analysis	11		68		Extrinsic Factors	23	0.96%
							Total	94	3.90%



YEAR	2	4	6	8	10	AT 130 MONTHS
SURVIVAL PROBABILITY	98.46%	97.21%	93.16%	87.80%	85.41%	83.89%
± 1 STANDARD ERROR	0.27%	0.38%	0.64%	0.92%	1.04%	1.20%
SAMPLE SIZE	1,920	1,550	1,220	950	610	210

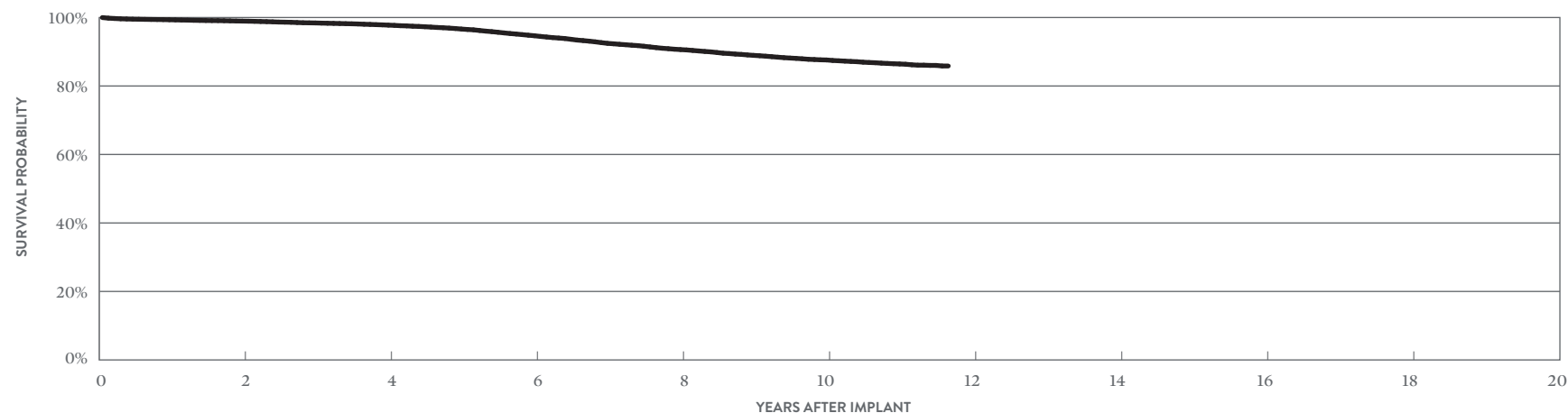
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

Riata™ ST

MODELS 7000 & 7001

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS			
		QTY	RATE	QTY	RATE		QTY	RATE	
US Regulatory Approval	June 2005	Cardiac Perforation	42	0.12%	32	0.09%	Conductor Fracture	24	0.07%
Registered US Implants	34,880	Conductor Fracture	0	0.00%	146	0.42%	Clavicular Crush	4	0.01%
Estimated Active US Implants	10,819	Lead Dislodgement	38	0.11%	58	0.17%	In the Pocket	7	0.02%
Insulation	Silicone	Failure to Capture	42	0.12%	329	0.94%	Intravascular	13	0.04%
Type and/or Fixation	Dual Coil, Active	Oversensing	40	0.11%	810	2.32%	Insulation Breach	587	1.68%
Polarity	Bipolar	Failure to Sense	7	0.02%	62	0.18%	Lead-to-Can Contact	308	0.88%
Steroid	Yes	Insulation Breach	1	<0.01%	722	2.07%	Lead-to-Lead Contact	157	0.45%
Number of US Advisories (see pg. 343)	One	Abnormal Pacing Impedance	8	0.02%	114	0.33%	Clavicular Crush	11	0.03%
		Abnormal Defibrillation Impedance	4	0.01%	197	0.56%	Externalized Conductors	36	0.10%
		Extracardiac Stimulation	3	<0.01%	5	0.01%	Other	75	0.22%
		Other	11	0.03%	93	0.27%	Crimps, Welds & Bonds	1	<0.01%
		Total	196	0.56%	2568	7.36%	Other	1	<0.01%
		Total Returned for Analysis	97		709		Extrinsic Factors	300	0.86%
							Total	913	2.62%



YEAR	2	4	6	8	10	AT 140 MONTHS
SURVIVAL PROBABILITY	98.95%	97.77%	94.67%	90.63%	87.60%	85.84%
± 1 STANDARD ERROR	0.06%	0.09%	0.15%	0.21%	0.26%	0.35%
SAMPLE SIZE	28,340	22,350	17,560	13,820	10,030	360

Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

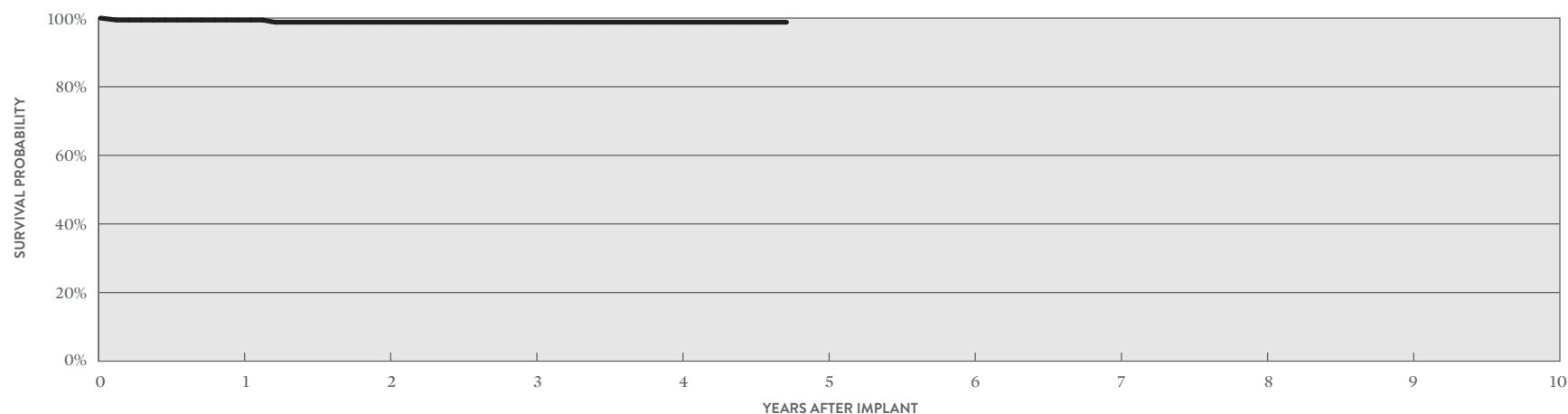
Riata™ ST

MODELS 7000 & 7001

US Regulatory Approval	June 2005
Number of Devices Enrolled in Study	179
Active Devices Enrolled in Study	30
Cumulative Months of Follow-up	8,070
Insulation	Silicone
Type and/or Fixation	Dual Coil, Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Insulation Breach	1	0.56%
Lead Dislodgement	1	0.56%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	0	0.00%
Insulation Breach	5	2.79%
Lead-to-Can Contact	3	1.68%
Lead-to-Lead Contact	1	0.56%
Clavicular Crush	0	0.00%
Externalized Conductors	0	0.00%
Other	1	0.56%
Crimps, Welds & Bonds	1	0.56%
Other	0	0.00%
Extrinsic Factors	0	0.00%
Total	6	3.35%



YEAR	1	2	3	4	AT 57 MONTHS
SURVIVAL PROBABILITY	99.43%	98.80%	98.80%	98.80%	98.80%
± 1 STANDARD ERROR	0.57%	0.85%	0.85%	0.85%	0.85%
SAMPLE SIZE	170	150	120	90	50

Defibrillation Leads

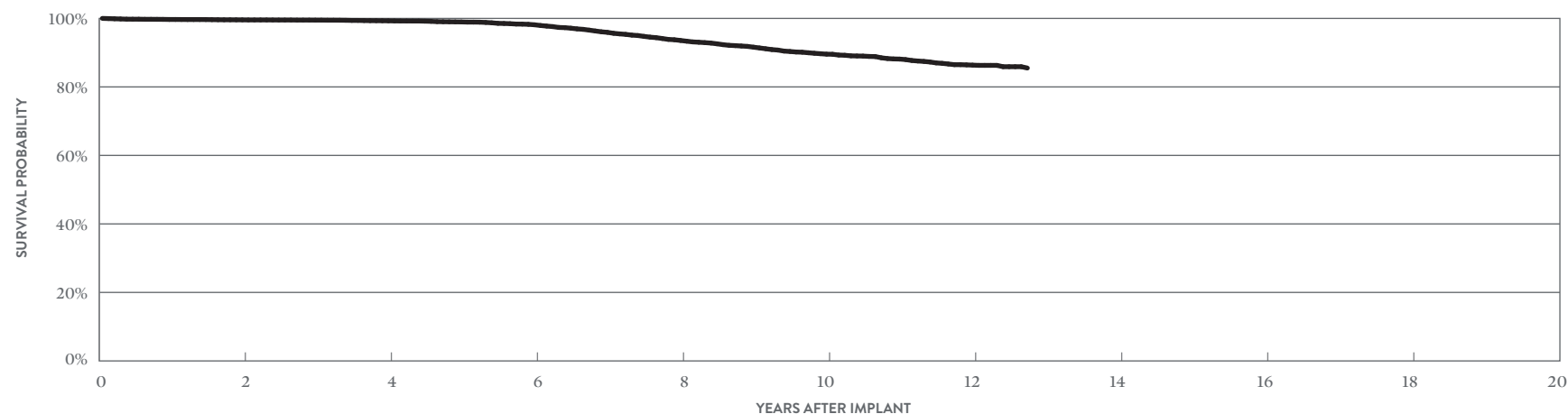
CUSTOMER REPORTED PERFORMANCE DATA

Riata™ i

MODELS 1590 & 1591

US Regulatory Approval	April 2004
Registered US Implants	9,701
Estimated Active US Implants	2,552
Insulation	Silicone
Type and/or Fixation	Dual Coil, Active
Polarity	Integrated Bipolar
Steroid	Yes
Number of US Advisories (see pg. 343)	One

MALFUNCTIONS	QTY	RATE
Conductor Fracture	7	0.07%
Clavicular Crush	1	0.01%
In the Pocket	1	0.01%
Intravascular	5	0.05%
Insulation Breach	182	1.88%
Lead-to-Can Contact	74	0.76%
Lead-to-Lead Contact	55	0.57%
Clavicular Crush	2	0.02%
Externalized Conductors	18	0.19%
Other	33	0.34%
Crimps, Welds & Bonds	0	0.00%
Other	1	0.01%
Extrinsic Factors	54	0.56%
Total	244	2.52%



YEAR	2	4	6	8	10	12	AT 153 MONTHS
SURVIVAL PROBABILITY	99.57%	99.29%	98.14%	93.56%	89.56%	86.36%	85.51%
± 1 STANDARD ERROR	0.07%	0.09%	0.17%	0.36%	0.48%	0.58%	0.62%
SAMPLE SIZE	8,030	6,360	4,930	3,850	3,000	1,800	260

Defibrillation Leads

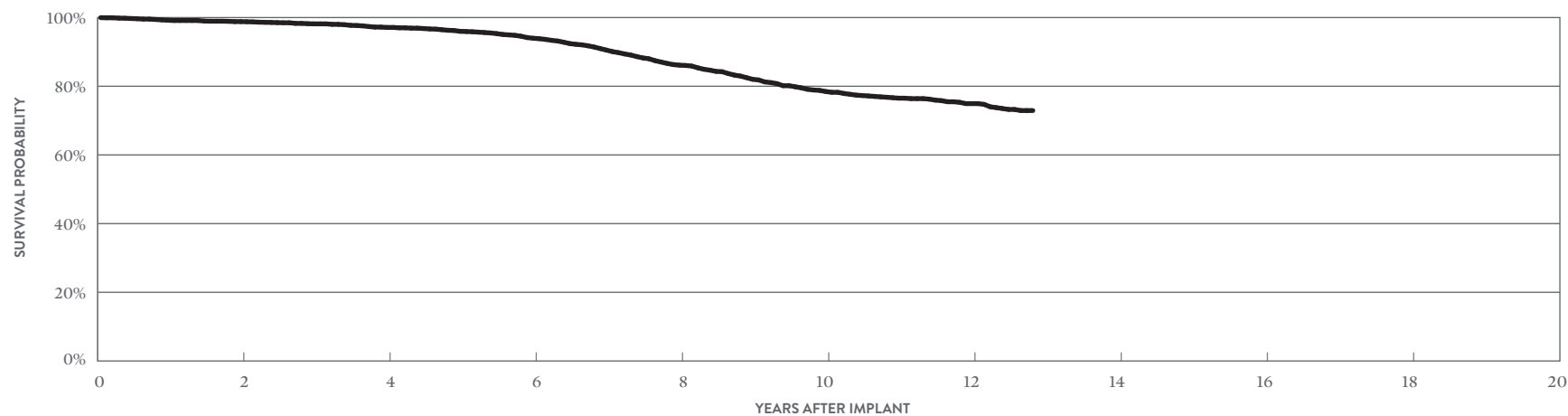
CUSTOMER REPORTED PERFORMANCE DATA

Riata™

MODEL 1582

US Regulatory Approval	March 2003
Registered US Implants	3,131
Estimated Active US Implants	703
Insulation	Silicone
Type and/or Fixation	Single Coil, Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories (see pg. 343)	One

MALFUNCTIONS	QTY	RATE
Conductor Fracture	3	0.10%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	3	0.10%
Insulation Breach	167	5.33%
Lead-to-Can Contact	51	1.63%
Lead-to-Lead Contact	30	0.96%
Clavicular Crush	2	0.06%
Externalized Conductors	49	1.56%
Other	35	1.12%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	35	1.12%
Total	205	6.55%



YEAR	2	4	6	8	10	12	AT 154 MONTHS
SURVIVAL PROBABILITY	98.82%	97.12%	93.97%	86.13%	78.46%	74.94%	72.92%
± 1 STANDARD ERROR	0.21%	0.34%	0.54%	0.88%	1.14%	1.30%	1.45%
SAMPLE SIZE	2,520	2,000	1,530	1,120	790	470	210

Defibrillation Leads

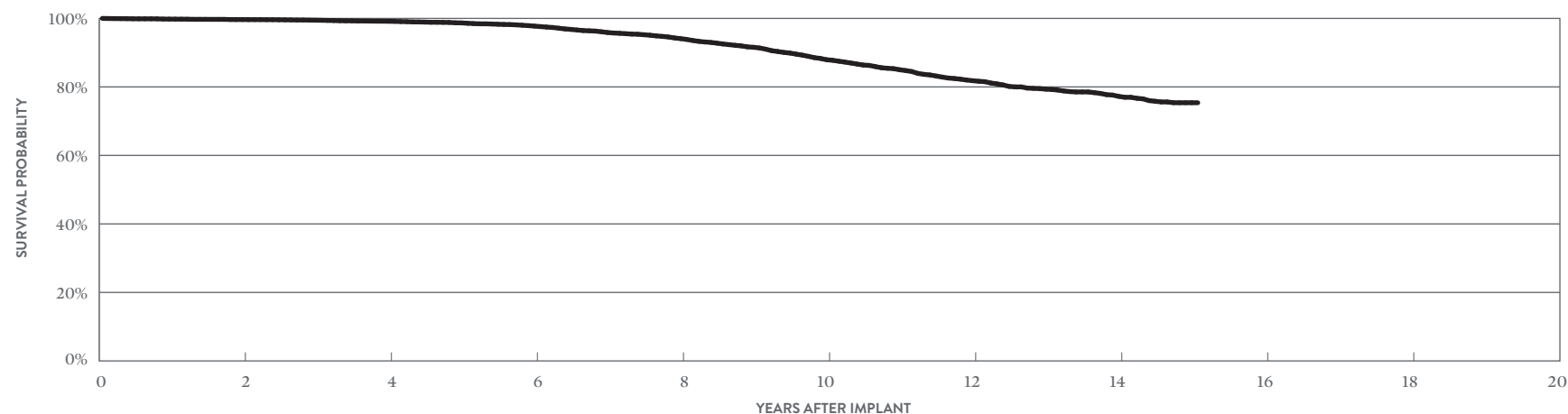
CUSTOMER REPORTED PERFORMANCE DATA

Riata™

MODELS 1570 & 1571

US Regulatory Approval	March 2002
Registered US Implants	10,280
Estimated Active US Implants	2,269
Insulation	Silicone
Type and/or Fixation	Dual Coil, Passive
Polarity	Bipolar
Steroid	Yes
Number of US Advisories (see pg. 343)	One

MALFUNCTIONS	QTY	RATE
Conductor Fracture	5	0.05%
Clavicular Crush	2	0.02%
In the Pocket	3	0.03%
Intravascular	0	0.00%
Insulation Breach	219	2.13%
Lead-to-Can Contact	103	1.00%
Lead-to-Lead Contact	40	0.39%
Clavicular Crush	2	0.02%
Externalized Conductors	40	0.39%
Other	34	0.33%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	59	0.57%
Total	283	2.75%



YEAR	2	4	6	8	10	12	14	AT 181 MONTHS
SURVIVAL PROBABILITY	99.66%	99.16%	97.75%	94.10%	87.94%	81.81%	77.24%	75.38%
± 1 STANDARD ERROR	0.06%	0.10%	0.18%	0.33%	0.51%	0.68%	0.86%	1.03%
SAMPLE SIZE	8,560	6,860	5,290	4,010	2,880	1,840	780	220

Defibrillation Leads

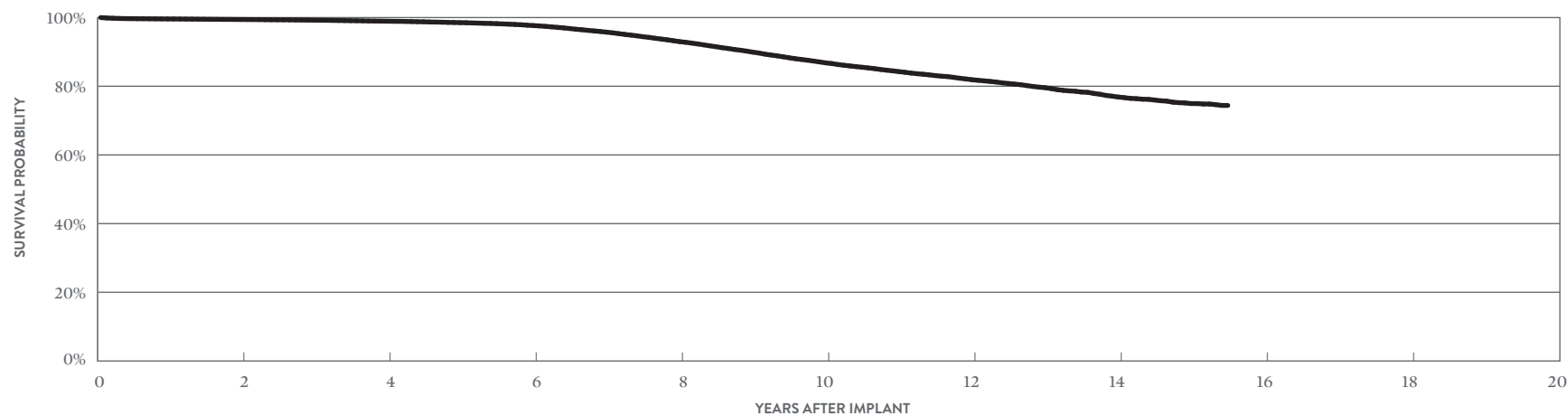
CUSTOMER REPORTED PERFORMANCE DATA

Riata™

MODELS 1580 & 1581

US Regulatory Approval	March 2002
Registered US Implants	68,398
Estimated Active US Implants	14,789
Insulation	Silicone
Type and/or Fixation	Dual Coil, Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories (see pg. 343)	One

MALFUNCTIONS	QTY	RATE
Conductor Fracture	32	0.05%
Clavicular Crush	4	<0.01%
In the Pocket	11	0.02%
Intravascular	17	0.02%
Insulation Breach	1756	2.57%
Lead-to-Can Contact	720	1.05%
Lead-to-Lead Contact	349	0.51%
Clavicular Crush	19	0.03%
Externalized Conductors	348	0.51%
Other	320	0.47%
Crimps, Welds & Bonds	3	<0.01%
Other	0	0.00%
Extrinsic Factors	538	0.79%
Total	2329	3.41%



YEAR	2	4	6	8	10	12	14	AT 186 MONTHS
SURVIVAL PROBABILITY	99.43%	98.94%	97.67%	92.99%	86.80%	81.90%	76.86%	74.41%
± 1 STANDARD ERROR	0.03%	0.04%	0.07%	0.14%	0.21%	0.25%	0.37%	0.58%
SAMPLE SIZE	56,210	44,570	34,250	25,880	18,840	12,460	3,760	270

Defibrillation Leads

ACTIVELY MONITORED STUDY DATA

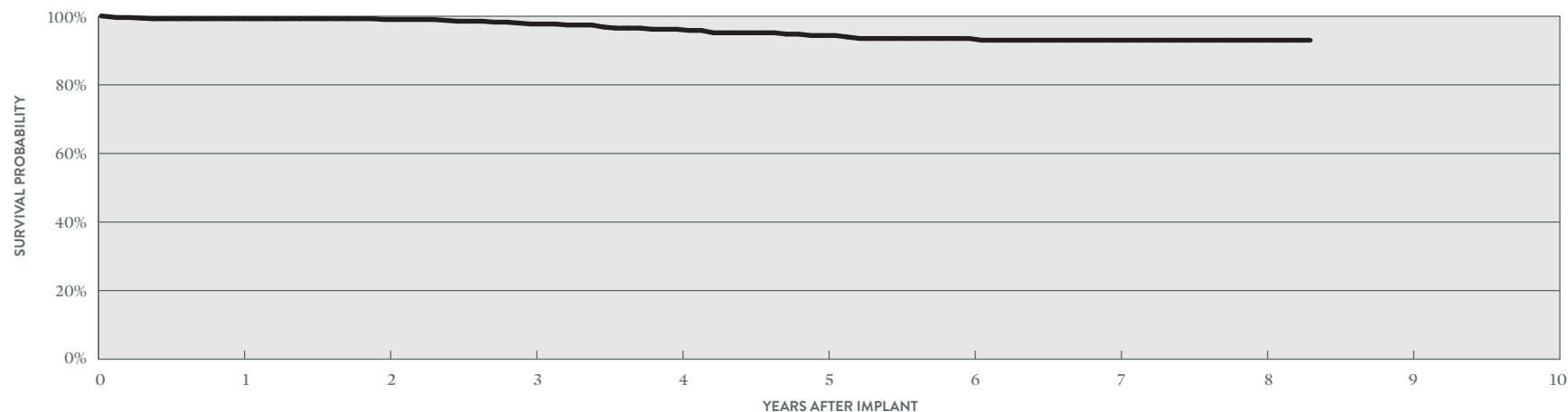
Riata™

MODELS 1580 & 1581

US Regulatory Approval	March 2002
Number of Devices Enrolled in Study	566
Active Devices Enrolled in Study	134
Cumulative Months of Follow-up	29,476
Insulation	Silicone
Type and/or Fixation	Dual Coil, Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Conductor Fracture	3	0.53%
Failure to Capture	1	0.18%
Insulation Breach	9	1.59%
Lead Dislodgement	2	0.35%
Oversensing	7	1.24%
Skin Erosion	1	0.18%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	1	0.18%
Clavicular Crush	0	0.00%
In the Pocket	0	0.00%
Intravascular	1	0.18%
Insulation Breach	22	3.89%
Lead-to-Can Contact	7	1.24%
Lead-to-Lead Contact	7	1.24%
Clavicular Crush	0	0.00%
Externalized Conductors	6	1.06%
Other	2	0.35%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	7	1.24%
Total	30	5.30%



YEAR	1	2	3	4	5	6	7	8	AT 100 MONTHS
SURVIVAL PROBABILITY	99.28%	99.05%	97.74%	96.21%	94.39%	93.52%	93.02%	93.02%	93.02%
± 1 STANDARD ERROR	0.36%	0.36%	0.66%	0.98%	1.26%	1.39%	1.47%	1.47%	1.47%
SAMPLE SIZE	530	470	390	320	260	200	160	100	50

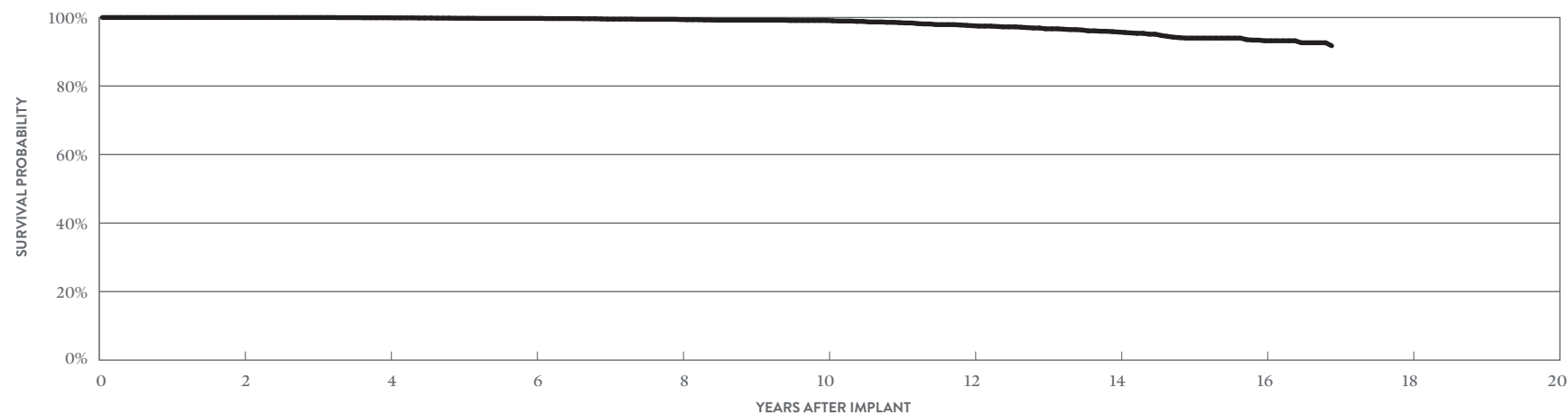
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

TVL™ ADX

MODEL 1559

US Regulatory Approval	November 1999
Registered US Implants	4,559
Estimated Active US Implants	685
Insulation	Silicone
Type and/or Fixation	Single Coil, Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None



YEAR	2	4	6	8	10	12	14	16	AT 203 MONTHS
SURVIVAL PROBABILITY	100.00%	99.93%	99.74%	99.41%	99.13%	97.60%	95.75%	93.21%	91.75%
± 1 STANDARD ERROR	0.00%	0.05%	0.10%	0.15%	0.22%	0.42%	0.62%	0.82%	0.93%
SAMPLE SIZE	3,720	2,940	2,270	1,700	1,250	980	810	610	220

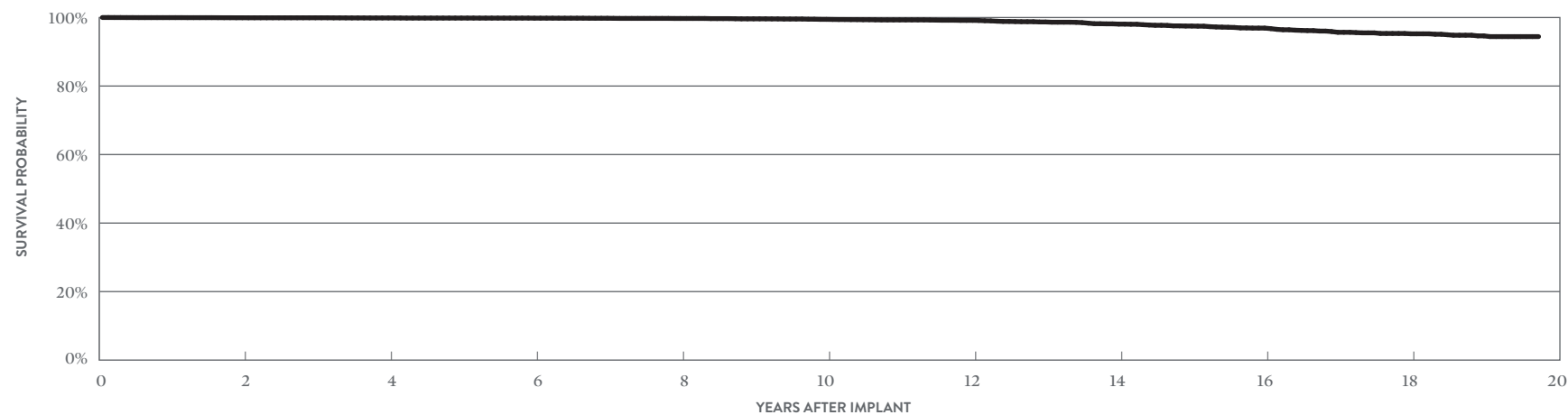
Defibrillation Leads

CUSTOMER REPORTED PERFORMANCE DATA

SPL™

MODELS SP01, SP02, SP03 & SP04

US Regulatory Approval	September 1997
Registered US Implants	12,374
Estimated Active US Implants	1,938
Insulation	Silicone
Type and/or Fixation	Dual Coil, Passive
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None



YEAR	2	4	6	8	10	12	14	16	18	AT 237 MONTHS
SURVIVAL PROBABILITY	99.93%	99.87%	99.81%	99.72%	99.46%	99.16%	98.07%	96.91%	95.24%	94.43%
± 1 STANDARD ERROR	0.03%	0.04%	0.04%	0.06%	0.10%	0.14%	0.24%	0.33%	0.46%	0.59%
SAMPLE SIZE	10,310	8,350	6,700	5,260	4,010	3,070	2,510	2,030	1,090	230

SUMMARY INFORMATION
Defibrillation Leads

Defibrillation Leads

Survival Probability Summary

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
LDA230Q	Optisure™ DF4	98.88%	98.88%								
LDA220Q	Optisure™ DF4	98.96%	98.71%	98.54%							
LDA210Q	Optisure™ DF4	99.04%	98.82%	98.64%							
LDA210	Optisure™ DF4	99.30%	98.81%								
7170Q/7171Q	Durata™ DF4	99.21%	98.87%	98.37%	97.73%	97.40%	96.78%	96.31%			
7120Q/7121Q	Durata™ DF4	99.22%	99.00%	98.75%	98.43%	98.04%	97.56%	96.98%	96.63%		
7122Q	Durata™ DF4	99.25%	98.99%	98.74%	98.39%	97.95%	97.44%	96.92%	96.63%		
7120/7121	Durata™	99.39%	99.09%	98.78%	98.43%	97.97%	97.47%	96.83%	96.21%	95.63%	
7122	Durata™	99.26%	98.85%	98.51%	98.01%	97.34%	96.88%	96.39%	95.91%	95.42%	
7070/7071	Riata™ ST Optim™	99.37%	99.09%	98.70%	98.27%	97.26%	96.44%	95.60%	94.71%	94.17%	92.69%
7020/7021	Riata™ ST Optim™	98.97%	98.56%	98.27%	97.74%	96.97%	96.25%	95.33%	94.59%	93.60%	92.92%
7022	Riata™ ST Optim™	99.03%	98.69%	98.42%	97.68%	97.45%	96.82%	96.02%	94.68%	92.17%	91.53%
7010/7011	Riata™ ST	99.60%	99.37%	98.85%	98.56%	97.40%	95.52%	93.47%	90.96%	89.03%	87.85%
7040/7041	Riata™ ST	99.38%	99.01%	98.43%	97.08%	95.82%	93.73%	91.63%	89.23%	87.23%	86.28%
7002	Riata™ ST	98.97%	98.46%	97.90%	97.21%	96.01%	93.16%	90.99%	87.80%	86.77%	85.41%
7000/7001	Riata™ ST	99.34%	98.95%	98.41%	97.77%	96.63%	94.67%	92.45%	90.63%	88.94%	87.60%
1590/1591	Riata™ i	99.69%	99.57%	99.50%	99.29%	98.97%	98.14%	95.93%	93.56%	91.60%	89.56%
1582	Riata™	99.21%	98.82%	98.16%	97.12%	95.99%	93.97%	90.53%	86.13%	82.02%	78.46%
1570/1571	Riata™	99.80%	99.66%	99.48%	99.16%	98.69%	97.75%	95.88%	94.10%	91.57%	87.94%
1580/1581	Riata™	99.59%	99.43%	99.24%	98.94%	98.52%	97.67%	95.75%	92.99%	89.93%	86.80%
1559	TVL™ ADX	100.00%	100.00%	100.00%	99.93%	99.78%	99.74%	99.53%	99.41%	99.21%	99.13%
SP01/SP02/SP03/SP04	SPL™	99.97%	99.93%	99.92%	99.87%	99.84%	99.81%	99.78%	99.72%	99.61%	99.46%

Defibrillation Leads

Acute Observation Summary

POST IMPLANT ≤30 DAYS

MODELS	US REGULATORY APPROVAL	REGISTERED US IMPLANTS	ESTIMATED ACTIVE US IMPLANTS	CARDIAC PERFORATION		CONDUCTOR FRACTURE		LEAD DISLODGEMENT		FAILURE TO CAPTURE		OVERSENSING		FAILURE TO SENSE		INSULATION BREACH		ABNORMAL PACING IMPEDANCE		ABNORMAL DEFIBRILLATION IMPEDANCE		EXTRACARDIAC STIMULATION		OTHER		TOTAL		TOTAL RETURNED FOR ANALYSIS
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	
LDA230Q	Feb-14	857	674	0	0.00%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.12%	0	0.00%	2	0.23%	0
LDA220Q	Feb-14	7,163	5,736	4	0.06%	0	0.00%	23	0.32%	8	0.11%	3	0.04%	1	0.01%	0	0.00%	0	0.00%	4	0.06%	1	0.01%	2	0.03%	46	0.64%	18
LDA210Q	Feb-14	22,051	17,894	26	0.12%	1	<0.01%	63	0.29%	24	0.11%	10	0.05%	6	0.03%	1	<0.01%	1	<0.01%	5	0.02%	0	0.00%	8	0.04%	145	0.66%	47
LDA210	Feb-14	797	656	0	0.00%	0	0.00%	2	0.25%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.25%	1
7170Q/7171Q	Jul-09	6,170	3,661	6	0.10%	1	0.02%	15	0.24%	9	0.15%	3	0.05%	0	0.00%	0	0.00%	1	0.02%	0	0.00%	0	0.00%	1	0.02%	36	0.58%	16
7120Q/7121Q	Jan-09	125,371	72,386	81	0.06%	2	<0.01%	232	0.19%	100	0.08%	45	0.04%	13	0.01%	0	0.00%	5	<0.01%	9	<0.01%	3	<0.01%	34	0.03%	524	0.42%	270
7122Q	Jan-09	85,535	56,595	109	0.13%	3	<0.01%	185	0.22%	81	0.09%	25	0.03%	7	<0.01%	0	0.00%	5	<0.01%	7	<0.01%	3	<0.01%	33	0.04%	458	0.54%	205
7120/7121	Sep-07	59,761	26,233	40	0.07%	1	<0.01%	69	0.12%	24	0.04%	49	0.08%	5	<0.01%	0	0.00%	1	<0.01%	19	0.03%	0	0.00%	21	0.04%	229	0.38%	92
7122	Sep-07	15,003	7,673	11	0.07%	1	<0.01%	20	0.13%	18	0.12%	12	0.08%	0	0.00%	0	0.00%	3	0.02%	1	<0.01%	2	0.01%	4	0.03%	72	0.48%	32
7070/7071	Jul-06	3,311	1,311	3	0.09%	1	0.03%	3	0.09%	5	0.15%	4	0.12%	3	0.09%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	19	0.57%	6
7020/7021	Jul-06	14,245	5,126	33	0.23%	0	0.00%	27	0.19%	17	0.12%	19	0.13%	8	0.06%	0	0.00%	1	<0.01%	4	0.03%	3	0.02%	0	0.00%	112	0.79%	53
7022	Jul-06	1,471	555	5	0.34%	0	0.00%	3	0.20%	1	0.07%	0	0.00%	0	0.00%	0	0.00%	2	0.14%	0	0.00%	0	0.00%	0	0.00%	11	0.75%	4
7010/7011	Mar-06	2,200	700	3	0.14%	0	0.00%	1	0.05%	2	0.09%	2	0.09%	1	0.05%	0	0.00%	1	0.05%	0	0.00%	0	0.00%	1	0.05%	11	0.50%	4
7040/7041	Mar-06	4,055	1,307	4	0.10%	0	0.00%	5	0.12%	1	0.02%	3	0.07%	0	0.00%	0	0.00%	2	0.05%	0	0.00%	0	0.00%	1	0.02%	16	0.39%	3
7002	Jun-05	2,408	762	6	0.25%	0	0.00%	3	0.12%	4	0.17%	4	0.17%	0	0.00%	0	0.00%	2	0.08%	1	0.04%	0	0.00%	1	0.04%	21	0.87%	11
7000/7001	Jun-05	34,880	10,819	42	0.12%	0	0.00%	38	0.11%	42	0.12%	40	0.11%	7	0.02%	1	<0.01%	8	0.02%	4	0.01%	3	<0.01%	11	0.03%	196	0.56%	97

Definitions of observations and complications can be found on [page 7](#).

Defibrillation Leads

Chronic Complication Summary

>30 DAYS

MODELS	US REGULATORY APPROVAL	REGISTERED US IMPLANTS	ESTIMATED ACTIVE US IMPLANTS	CARDIAC PERFORATION		CONDUCTOR FRACTURE		LEAD DISLODGE		FAILURE TO CAPTURE		OVERSENSING		FAILURE TO SENSE		INSULATION BREACH		ABNORMAL PACING IMPEDANCE		ABNORMAL DEFIBRILLATION IMPEDANCE		EXTRACARDIAC STIMULATION		OTHER		TOTAL		TOTAL RETURNED FOR ANALYSIS
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	
LDA230Q	Feb-14	857	674	0	0.00%	0	0.00%	2	0.23%	3	0.35%	2	0.23%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	7	0.82%	4
LDA220Q	Feb-14	7,163	5,736	3	0.04%	2	0.03%	40	0.56%	24	0.34%	15	0.21%	1	0.01%	1	0.01%	0	0.00%	4	0.06%	0	0.00%	2	0.03%	92	1.28%	34
LDA210Q	Feb-14	22,051	17,894	10	0.05%	2	<0.01%	107	0.49%	42	0.19%	36	0.16%	8	0.04%	1	<0.01%	2	<0.01%	10	0.05%	2	<0.01%	9	0.04%	229	1.04%	83
LDA210	Feb-14	797	656	0	0.00%	1	0.13%	1	0.13%	2	0.25%	2	0.25%	0	0.00%	0	0.00%	1	0.13%	0	0.00%	0	0.00%	0	0.00%	7	0.88%	2
7170Q/7171Q	Jul-09	6,170	3,661	5	0.08%	9	0.15%	24	0.39%	48	0.78%	27	0.44%	0	0.00%	2	0.03%	12	0.19%	11	0.18%	0	0.00%	0	0.00%	138	2.24%	39
7120Q/7121Q	Jan-09	125,371	72,386	35	0.03%	123	0.10%	566	0.45%	609	0.49%	499	0.40%	69	0.06%	26	0.02%	101	0.08%	249	0.20%	7	<0.01%	61	0.05%	2345	1.87%	893
7122Q	Jan-09	85,535	56,595	40	0.05%	41	0.05%	358	0.42%	262	0.31%	238	0.28%	32	0.04%	14	0.02%	44	0.05%	69	0.08%	8	<0.01%	31	0.04%	1137	1.33%	517
7120/7121	Sep-07	59,761	26,233	16	0.03%	128	0.21%	183	0.31%	314	0.53%	548	0.92%	63	0.11%	61	0.10%	168	0.28%	249	0.42%	3	<0.01%	46	0.08%	1779	2.98%	499
7122	Sep-07	15,003	7,673	3	0.02%	29	0.19%	61	0.41%	75	0.50%	110	0.73%	11	0.07%	21	0.14%	36	0.24%	28	0.19%	2	0.01%	7	0.05%	383	2.55%	170
7070/7071	Jul-06	3,311	1,311	2	0.06%	17	0.51%	12	0.36%	34	1.03%	49	1.48%	2	0.06%	6	0.18%	13	0.39%	14	0.42%	1	0.03%	2	0.06%	152	4.59%	33
7020/7021	Jul-06	14,245	5,126	16	0.11%	54	0.38%	62	0.44%	147	1.03%	235	1.65%	20	0.14%	24	0.17%	44	0.31%	91	0.64%	2	0.01%	28	0.20%	723	5.08%	204
7022	Jul-06	1,471	555	3	0.20%	10	0.68%	11	0.75%	12	0.82%	22	1.50%	1	0.07%	7	0.48%	4	0.27%	3	0.20%	1	0.07%	1	0.07%	75	5.10%	26
7010/7011	Mar-06	2,200	700	3	0.14%	5	0.23%	8	0.36%	10	0.45%	43	1.95%	3	0.14%	40	1.82%	26	1.18%	19	0.86%	0	0.00%	3	0.14%	160	7.27%	37
7040/7041	Mar-06	4,055	1,307	3	0.07%	34	0.84%	5	0.12%	49	1.21%	96	2.37%	14	0.35%	55	1.36%	19	0.47%	24	0.59%	1	0.02%	7	0.17%	307	7.57%	68
7002	Jun-05	2,408	762	5	0.21%	9	0.37%	9	0.37%	22	0.91%	62	2.57%	2	0.08%	69	2.87%	4	0.17%	9	0.37%	0	0.00%	8	0.33%	199	8.26%	68
7000/7001	Jun-05	34,880	10,819	32	0.09%	146	0.42%	58	0.17%	329	0.94%	810	2.32%	62	0.18%	722	2.07%	114	0.33%	197	0.56%	5	0.01%	93	0.27%	2568	7.36%	709

Definitions of observations and complications can be found on [page 7](#).

Defibrillation Leads

U.S. Malfunction Summary

MODELS	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
LDA230Q	857	2.70%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	6	0.70%	7	0.82%
LDA220Q	7,163	2.90%	0	0.00%	2	0.03%	0	0.00%	0	0.00%	36	0.50%	38	0.53%
LDA210Q	22,051	2.70%	2	<0.01%	3	0.01%	0	0.00%	1	<0.01%	88	0.40%	94	0.43%
LDA210	797	3.60%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	0.50%	4	0.50%
7170Q/7171Q	6,170	4.20%	2	0.03%	5	0.08%	0	0.00%	0	0.00%	37	0.60%	44	0.71%
7120Q/7121Q	125,371	4.40%	25	0.02%	187	0.15%	2	<0.01%	36	0.03%	752	0.60%	1002	0.80%
7122Q	85,535	4.50%	11	0.01%	87	0.10%	0	0.00%	15	0.02%	475	0.56%	588	0.69%
7120/7121	59,761	5.50%	32	0.05%	136	0.23%	1	<0.01%	9	0.02%	393	0.66%	571	0.96%
7122	15,003	7.20%	16	0.11%	57	0.38%	0	0.00%	4	0.03%	126	0.84%	203	1.35%
7070/7071	3,311	7.40%	1	0.03%	13	0.39%	0	0.00%	0	0.00%	21	0.63%	35	1.06%
7020/7021	14,245	7.10%	10	0.07%	50	0.35%	0	0.00%	0	0.00%	172	1.21%	232	1.63%
7022	1,471	9.90%	3	0.20%	7	0.48%	0	0.00%	0	0.00%	21	1.43%	31	2.11%
7010/7011	2,200	8.80%	2	0.09%	38	1.73%	0	0.00%	0	0.00%	11	0.50%	51	2.32%
7040/7041	4,055	8.20%	4	0.10%	55	1.36%	0	0.00%	0	0.00%	29	0.72%	88	2.17%
7002	2,408	9.80%	5	0.21%	66	2.74%	0	0.00%	0	0.00%	23	0.96%	94	3.90%
7000/7001	34,880	7.40%	24	0.07%	587	1.68%	1	<0.01%	1	<0.01%	300	0.86%	913	2.62%
1590/1591	9,701	7.50%	7	0.07%	182	1.88%	0	0.00%	1	0.01%	54	0.56%	244	2.52%
1582	3,131	11.50%	3	0.10%	167	5.33%	0	0.00%	0	0.00%	35	1.12%	205	6.55%
1570/1571	10,280	8.40%	5	0.05%	219	2.13%	0	0.00%	0	0.00%	59	0.57%	283	2.75%
1580/1581	68,398	8.00%	32	0.05%	1756	2.57%	3	<0.01%	0	0.00%	538	0.79%	2329	3.41%

Definitions of malfunction categories can be found on [pages 8-9](#).

Defibrillation Leads

Worldwide Malfunction Summary

MODELS	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
LDA230Q	887	2.6%	0	0.00%	1	0.11%	0	0.00%	0	0.00%	6	0.68%	7	0.79%
LDA220Q	9,875	2.3%	0	0.00%	2	0.02%	0	0.00%	1	0.01%	45	0.46%	48	0.49%
LDA210Q	38,906	1.7%	3	0.01%	6	0.02%	0	0.00%	6	0.02%	159	0.41%	174	0.45%
LDA210	878	3.4%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	5	0.57%	5	0.57%
7170Q/7171Q	17,866	2.3%	7	0.04%	15	0.08%	2	0.01%	0	0.00%	68	0.38%	92	0.51%
7120Q/7121Q	213,064	3.2%	55	0.03%	262	0.12%	3	<0.01%	93	0.04%	1209	0.57%	1622	0.76%
7122Q	233,385	2.3%	42	0.02%	197	0.08%	2	<0.01%	133	0.06%	1179	0.51%	1553	0.67%
7120/7121	144,039	3.1%	114	0.08%	228	0.16%	1	<0.01%	25	0.02%	776	0.54%	1144	0.79%
7122	64,505	2.9%	114	0.18%	134	0.21%	1	<0.01%	23	0.04%	468	0.73%	740	1.15%

Definitions of malfunction categories can be found on [pages 8-9](#).

Defibrillation Leads

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	ABNORMAL DEFIBRILLATION IMPEDANCE		ABNORMAL PACING IMPEDANCE		CARDIAC PERFORATION		CONDUCTOR FRACTURE		EXTRACARDIAC STIMULATION		FAILURE TO CAPTURE		FAILURE TO SENSE		INAPPROPRIATE SHOCK		INSULATION BREACH		LEAD DISLODGEEMENT		OVERSENSING		PERICARDIAL EFFUSION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
7170Q/7171Q	116	55	6,504	0	0.00%	1	0.86%	0	0.00%	1	0.86%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.86%	0	0.00%	0	0.00%	0	0.00%	3	2.59%
7120Q/7121Q	4,315	1,891	226,061	5	0.12%	3	0.07%	1	0.02%	15	0.35%	0	0.00%	19	0.44%	5	0.12%	4	0.09%	3	0.07%	39	0.90%	7	0.16%	0	0.00%	0	0.00%	101	2.34%
7122Q	1,542	755	73,384	2	0.13%	0	0.00%	0	0.00%	4	0.26%	0	0.00%	7	0.45%	1	0.06%	0	0.00%	0	0.00%	7	0.45%	1	0.06%	2	0.13%	0	0.00%	24	1.56%
7120/7121	3,561	1,066	209,866	4	0.11%	10	0.28%	0	0.00%	11	0.31%	0	0.00%	13	0.37%	2	0.06%	2	0.06%	10	0.28%	20	0.56%	10	0.28%	0	0.00%	1	0.03%	83	2.33%
7122	450	176	26,801	1	0.22%	4	0.89%	0	0.00%	6	1.33%	0	0.00%	5	1.11%	1	0.22%	0	0.00%	0	0.00%	5	1.11%	2	0.44%	0	0.00%	0	0.00%	24	5.33%
7070/7071	288	81	17,458	1	0.35%	2	0.69%	1	0.35%	2	0.69%	0	0.00%	1	0.35%	0	0.00%	0	0.00%	0	0.00%	1	0.35%	1	0.35%	0	0.00%	0	0.00%	9	3.13%
7020/7021	1,470	258	83,312	0	0.00%	6	0.41%	1	0.07%	8	0.54%	0	0.00%	14	0.95%	1	0.07%	0	0.00%	2	0.14%	9	0.61%	4	0.27%	0	0.00%	1	0.07%	46	3.13%
7000/7001	179	30	8,070	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.56%	1	0.56%	0	0.00%	0	0.00%	0	0.00%	2	1.12%
1580/1581	566	134	29,476	0	0.00%	0	0.00%	0	0.00%	3	0.53%	0	0.00%	1	0.18%	0	0.00%	0	0.00%	9	1.59%	2	0.35%	7	1.24%	0	0.00%	1	0.18%	23	4.06%

A list of complications can be found on [page 12](#).

Defibrillation Leads

Actively Monitored Study Data Summary

MALFUNCTIONS

MODELS	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
7170Q/7171Q	116	5.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	1.72%	2	1.72%
7120Q/7121Q	4,315	5.70%	5	0.12%	6	0.14%	0	0.00%	1	0.02%	46	1.07%	58	1.34%
7122Q	1,542	5.60%	2	0.13%	5	0.32%	0	0.00%	0	0.00%	14	0.91%	21	1.36%
7120/7121	3,561	4.90%	1	0.03%	12	0.34%	0	0.00%	1	0.03%	30	0.84%	44	1.24%
7122	450	5.60%	2	0.44%	2	0.44%	0	0.00%	0	0.00%	8	1.78%	12	2.67%
7070/7071	288	2.80%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.35%	1	0.35%
7020/7021	1,470	5.60%	3	0.20%	3	0.20%	0	0.00%	0	0.00%	14	0.95%	20	1.36%
7000/7001	179	8.40%	0	0.00%	5	2.79%	1	0.56%	0	0.00%	0	0.00%	6	3.35%
1580/1581	566	7.10%	1	0.18%	22	3.89%	0	0.00%	0	0.00%	7	1.24%	30	5.30%

Definitions of malfunction categories can be found on [pages 8-9](#).

Dual-Chamber Pacemakers

Dual-Chamber Pacemakers

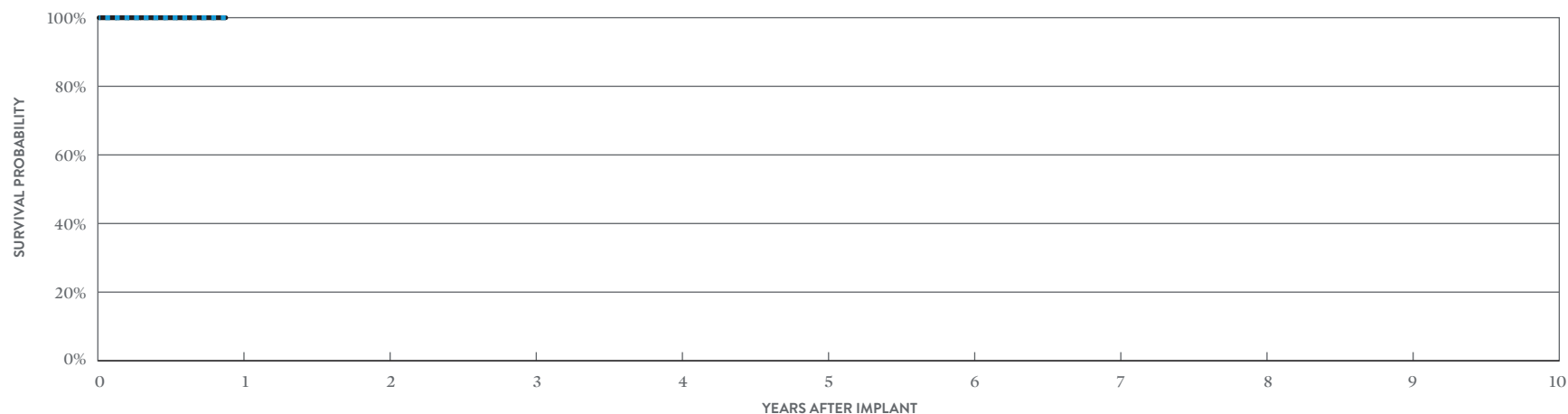
CUSTOMER REPORTED PERFORMANCE DATA

Assurity MRI™

MODEL PM2272

US Regulatory Approval	January 2017
Registered US Implants	36,177
Estimated Active US Implants	34,323
Estimated Longevity	9.4 Years
Normal Battery Depletion	0
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	1	<0.01%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	AT 11 MONTHS
SURVIVAL PROBABILITY	99.99%
± 1 STANDARD ERROR	0.00%
SAMPLE SIZE	1,230

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	AT 11 MONTHS
SURVIVAL PROBABILITY	99.99%
± 1 STANDARD ERROR	0.00%

Dual-Chamber Pacemakers

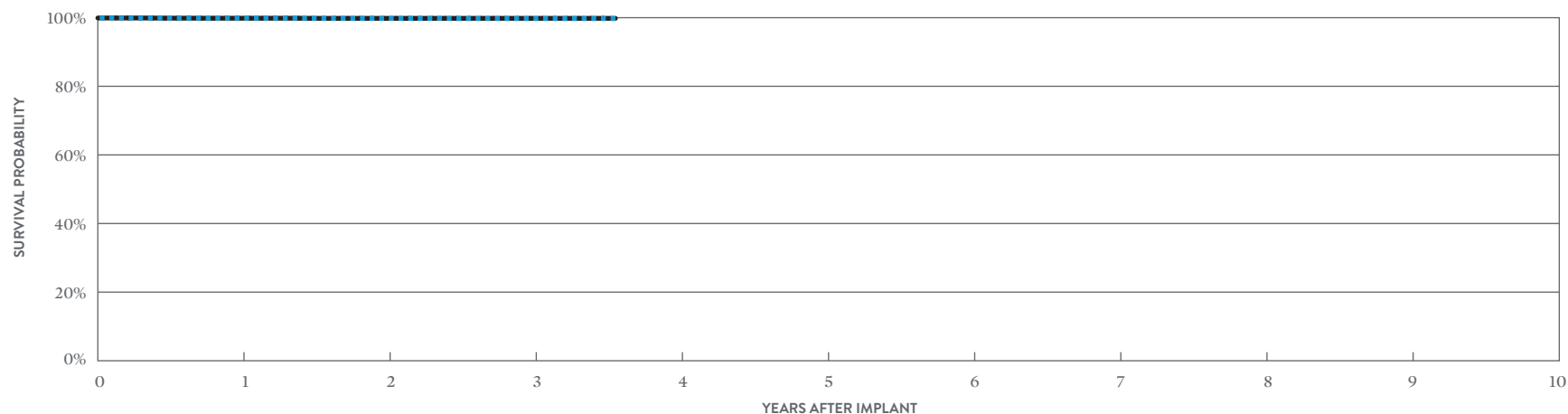
CUSTOMER REPORTED PERFORMANCE DATA

Endurity™ DR

MODEL PM2160

US Regulatory Approval	March 2014
Registered US Implants	8,929
Estimated Active US Implants	6,924
Estimated Longevity	9.7 Years
Normal Battery Depletion	2
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	5	0.06%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	2	0.02%
Total	0	0.00%	7	0.08%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 43 MONTHS
SURVIVAL PROBABILITY	99.84%	99.78%	99.73%	99.73%
± 1 STANDARD ERROR	0.04%	0.05%	0.06%	0.06%
SAMPLE SIZE	8,250	6,670	4,150	300

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 43 MONTHS
SURVIVAL PROBABILITY	99.84%	99.81%	99.81%	99.81%
± 1 STANDARD ERROR	0.04%	0.05%	0.05%	0.05%

Dual-Chamber Pacemakers

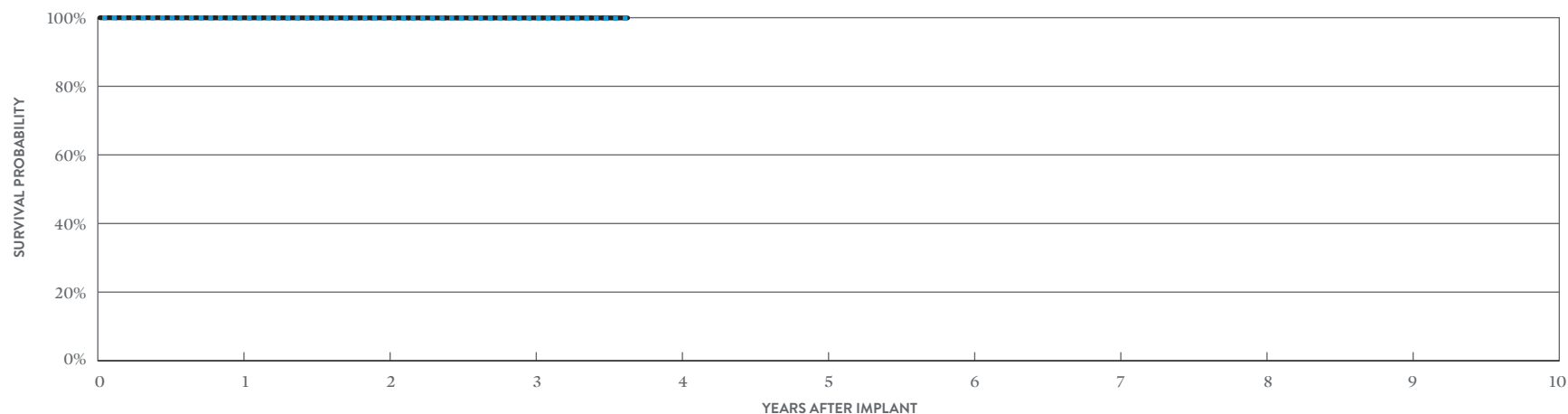
CUSTOMER REPORTED PERFORMANCE DATA

Assurity™ DR RF

MODEL PM2240

US Regulatory Approval	March 2014
Registered US Implants	168,347
Estimated Active US Implants	134,324
Estimated Longevity	9.4 Years
Normal Battery Depletion	11
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	3	<0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	1	<0.01%	22	0.01%
Possible Early Battery Depletion	2	<0.01%	0	0.00%
Other	0	0.00%	7	<0.01%
Total	5	<0.01%	33	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 44 MONTHS
SURVIVAL PROBABILITY	99.95%	99.92%	99.90%	99.86%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.02%
SAMPLE SIZE	147,110	96,690	42,460	620

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 44 MONTHS
SURVIVAL PROBABILITY	99.95%	99.94%	99.93%	99.93%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.01%

Dual-Chamber Pacemakers

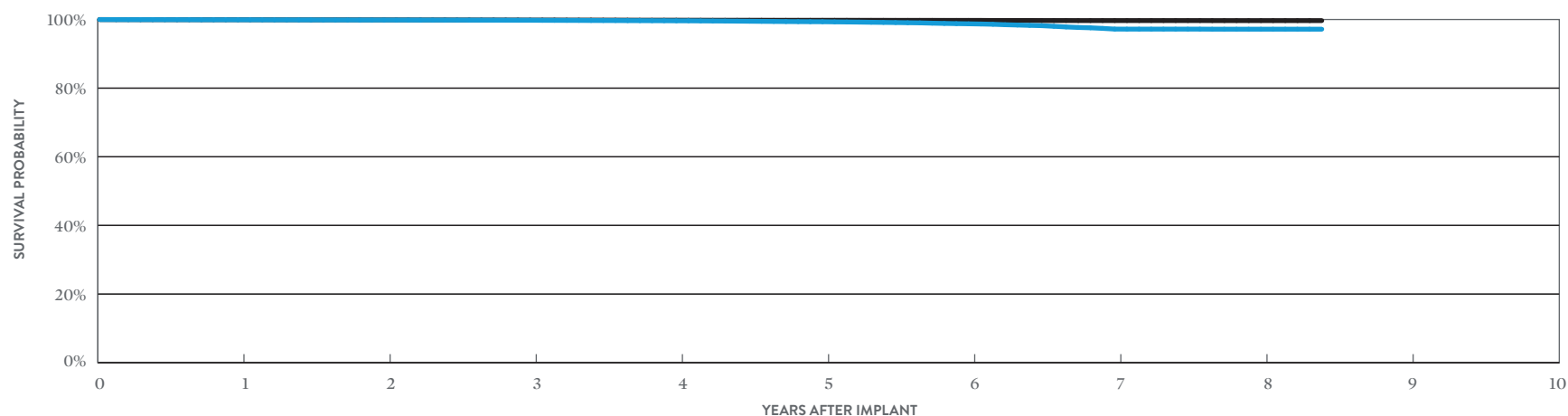
CUSTOMER REPORTED PERFORMANCE DATA

Accent™ DR RF

MODEL PM2210

US Regulatory Approval	July 2009
Registered US Implants	243,042
Estimated Active US Implants	131,570
Estimated Longevity	8 Years
Normal Battery Depletion	634
Number of US Advisories (see pgs. 334, 336)	Two

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	15	<0.01%	44	0.02%
Electrical Interconnect	7	<0.01%	31	0.01%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	4	<0.01%
Mechanical	0	0.00%	20	<0.01%
Possible Early Battery Depletion	7	<0.01%	22	<0.01%
Other	5	<0.01%	37	0.02%
Total	34	0.01%	158	0.07%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	99.92%	99.86%	99.78%	99.62%	99.35%	98.75%	97.21%	97.18%	97.18%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.01%	0.02%	0.03%	0.07%	0.07%	0.07%
SAMPLE SIZE	228,250	202,270	180,990	152,670	113,070	74,100	41,890	16,120	660

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 101 MONTHS
SURVIVAL PROBABILITY	99.95%	99.90%	99.84%	99.80%	99.77%	99.74%	99.69%	99.68%	99.68%
± 1 STANDARD ERROR	0.00%	0.01%	0.01%	0.01%	0.01%	0.01%	0.02%	0.02%	0.02%

Dual-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

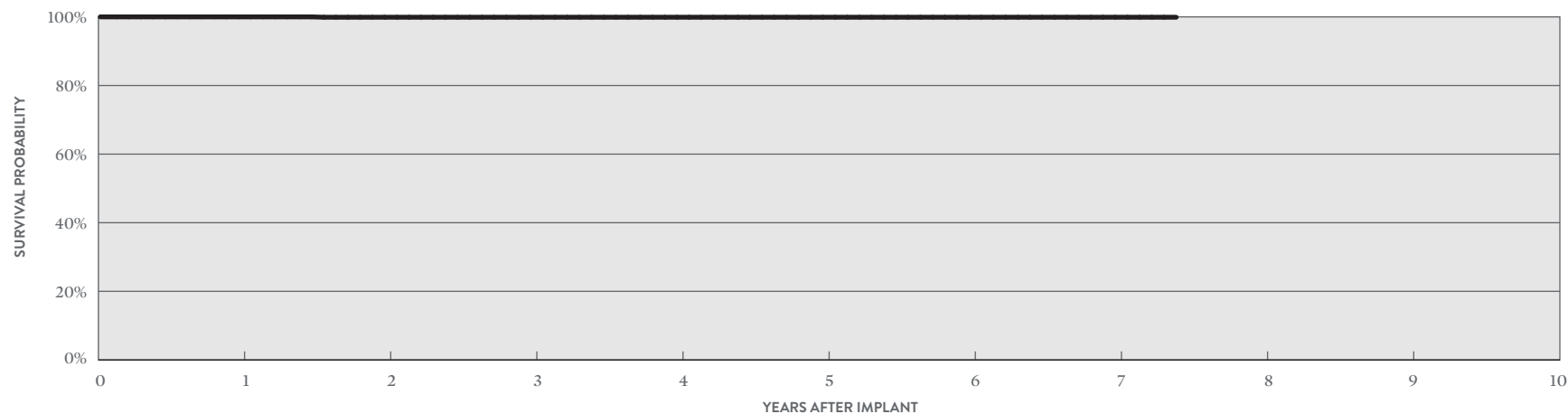
Accent™ DR RF

MODEL PM2210

US Regulatory Approval	July 2009
Number of Devices Enrolled in Study	1,774
Active Devices Enrolled in Study	323
Cumulative Months of Follow-up	55,167
Estimated Longevity	8 Years

QUALIFYING COMPLICATIONS	QTY	RATE
Premature Battery Depletion	1	0.06%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.06%
Electrical Interconnect	0	0.00%	1	0.06%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	2	0.11%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	7	AT 89 MONTHS
SURVIVAL PROBABILITY	100.00%	99.90%	99.90%	99.90%	99.90%	99.90%	99.90%	99.90%
± 1 STANDARD ERROR	0.00%	0.10%	0.10%	0.10%	0.10%	0.10%	0.10%	0.10%
SAMPLE SIZE	1,540	1,060	650	460	400	350	220	50

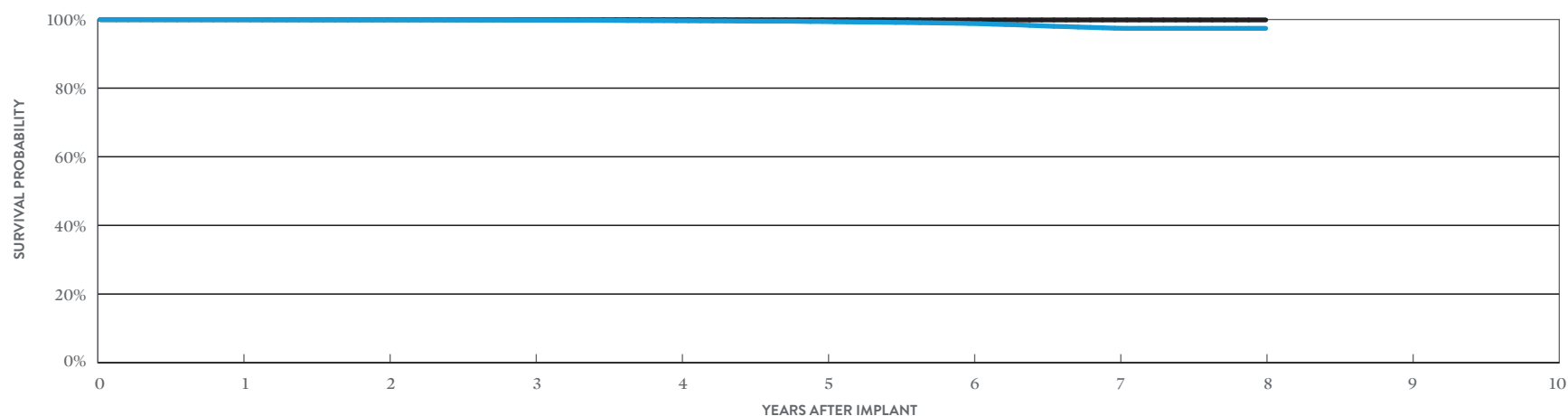
Dual-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

Accent™ DR MODEL PM2110

US Regulatory Approval	July 2009
Registered US Implants	48,906
Estimated Active US Implants	27,342
Estimated Longevity	9.2 Years
Normal Battery Depletion	129
Number of US Advisories (see pg. 336)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	3	<0.01%
Electrical Interconnect	2	<0.01%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	3	<0.01%
Mechanical	0	0.00%	5	0.01%
Possible Early Battery Depletion	0	0.00%	2	<0.01%
Other	0	0.00%	0	0.00%
Total	4	<0.01%	13	0.03%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8
SURVIVAL PROBABILITY	99.96%	99.91%	99.83%	99.64%	99.42%	98.85%	97.47%	97.42%
± 1 STANDARD ERROR	0.01%	0.01%	0.02%	0.03%	0.04%	0.07%	0.15%	0.17%
SAMPLE SIZE	46,000	40,860	36,570	30,860	22,800	14,490	7,220	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8
SURVIVAL PROBABILITY	99.97%	99.95%	99.93%	99.93%	99.92%	99.88%	99.88%	99.88%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.01%	0.01%	0.02%	0.02%	0.02%

Dual-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

Accent™ DR

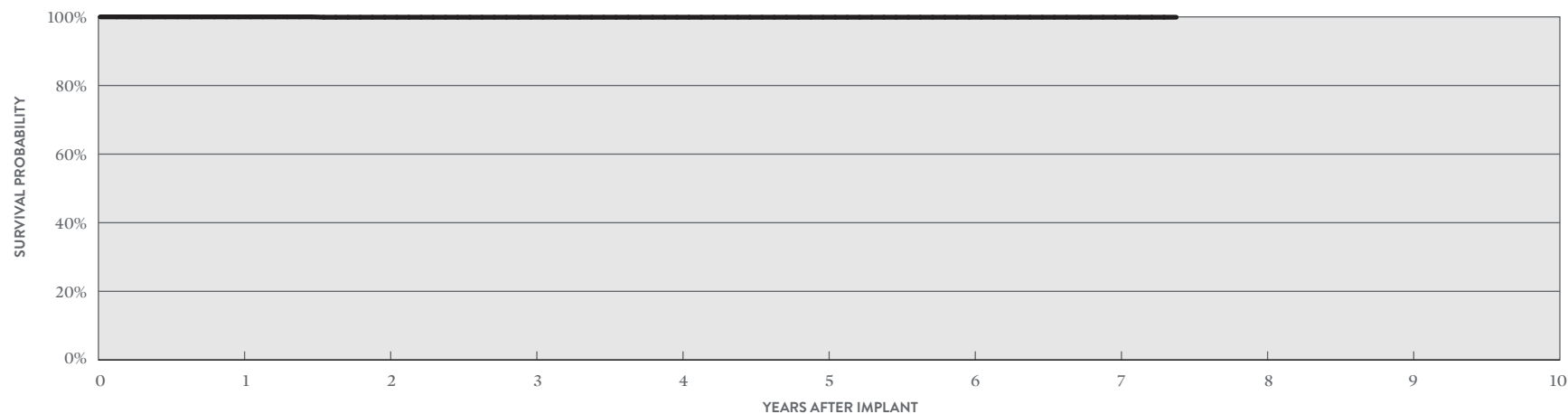
MODEL PM2110

US Regulatory Approval	June 2013
Number of Devices Enrolled in Study	227
Active Devices Enrolled in Study	61
Cumulative Months of Follow-up	9,045
Estimated Longevity	9.2 Years

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	5	6	AT 78 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
SAMPLE SIZE	210	150	100	90	80	70	50

Dual-Chamber Pacemakers

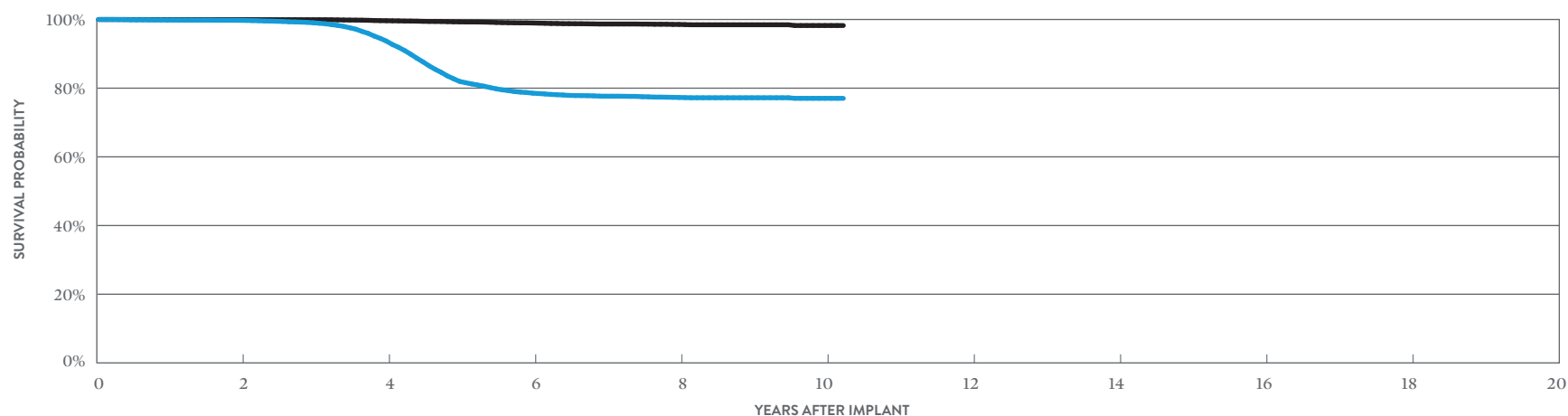
CUSTOMER REPORTED PERFORMANCE DATA

Zephyr™ DR

MODEL 5820

US Regulatory Approval	March 2007
Registered US Implants	53,978
Estimated Active US Implants	17,194
Estimated Longevity	6.5 Years
Normal Battery Depletion	2,260
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	34	0.06%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	9	0.02%
Mechanical	0	0.00%	2	<0.01%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	0	0.00%	73	0.14%
Total	2	<0.01%	119	0.22%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 123 MONTHS
SURVIVAL PROBABILITY	99.75%	93.70%	78.54%	77.29%	77.04%	77.04%
± 1 STANDARD ERROR	0.02%	0.13%	0.25%	0.27%	0.30%	0.30%
SAMPLE SIZE	42,230	30,350	15,890	5,580	1,030	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 123 MONTHS
SURVIVAL PROBABILITY	99.96%	99.64%	98.97%	98.56%	98.24%	98.24%
± 1 STANDARD ERROR	0.01%	0.03%	0.07%	0.10%	0.19%	0.19%

Dual-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

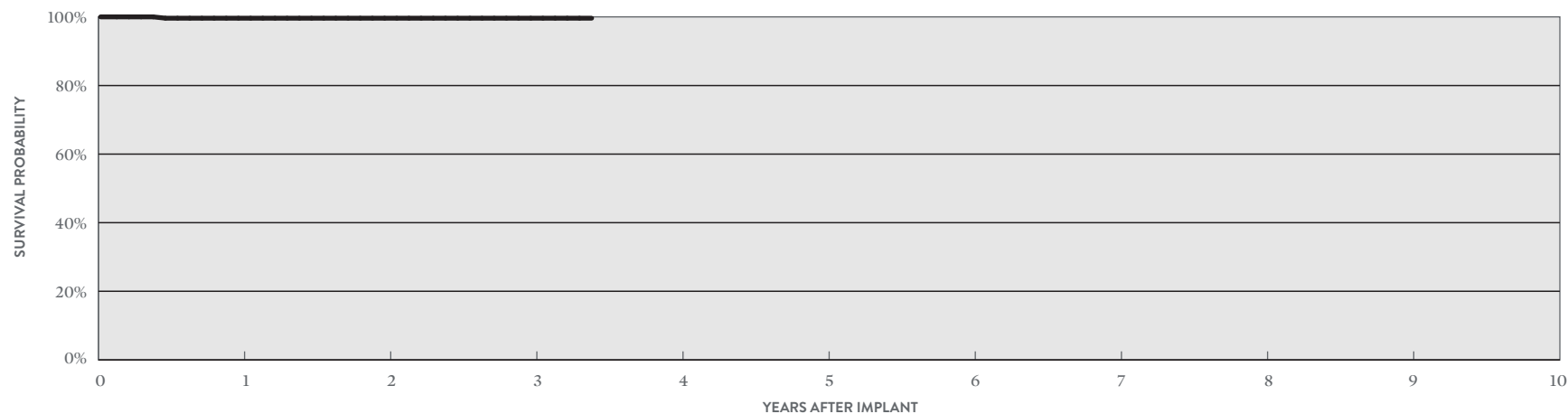
Zephyr™ DR

MODEL 5820

US Regulatory Approval	March 2007
Number of Devices Enrolled in Study	283
Active Devices Enrolled in Study	8
Cumulative Months of Follow-up	7,892
Estimated Longevity	6.5 Years

QUALIFYING COMPLICATIONS	QTY	RATE
Skin Erosion	1	0.35%

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	AT 41 MONTHS
SURVIVAL PROBABILITY	99.62%	99.62%	99.62%	99.62%
± 1 STANDARD ERROR	0.38%	0.38%	0.38%	0.38%
SAMPLE SIZE	260	200	120	50

Dual-Chamber Pacemakers

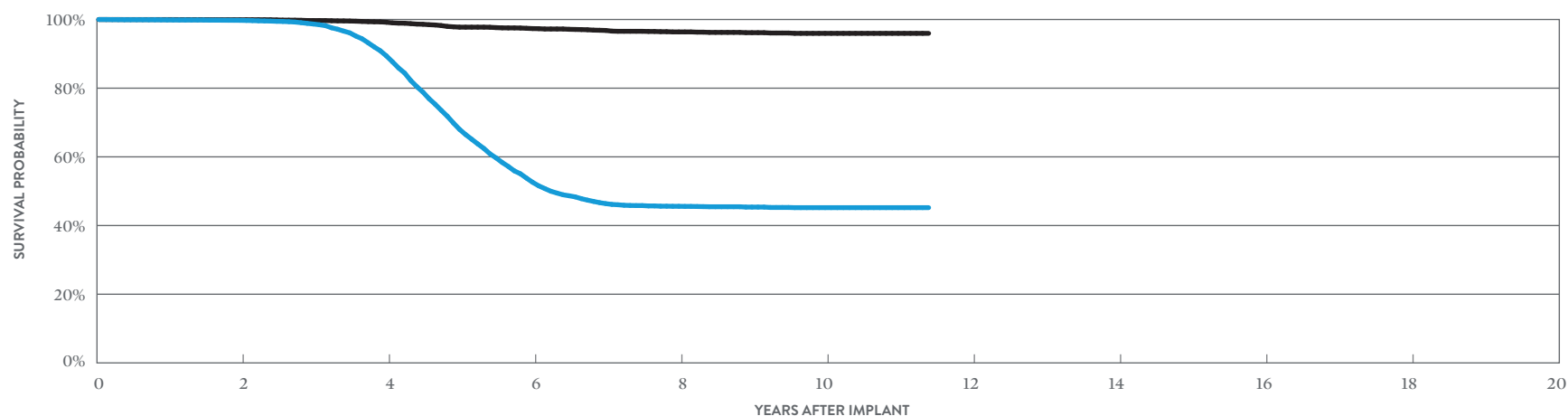
CUSTOMER REPORTED PERFORMANCE DATA

Victory™ DR

MODEL 5810

US Regulatory Approval	December 2005
Registered US Implants	26,311
Estimated Active US Implants	2,756
Estimated Longevity	6.5 Years
Normal Battery Depletion	2,775
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	<0.01%	89	0.34%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	8	0.03%
Mechanical	0	0.00%	2	<0.01%
Possible Early Battery Depletion	0	0.00%	17	0.06%
Other	0	0.00%	37	0.14%
Total	1	<0.01%	153	0.58%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 137 MONTHS
SURVIVAL PROBABILITY	99.75%	89.39%	52.59%	45.60%	45.21%	45.21%
± 1 STANDARD ERROR	0.03%	0.23%	0.43%	0.45%	0.45%	0.45%
SAMPLE SIZE	20,910	14,940	7,620	3,500	1,850	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 137 MONTHS
SURVIVAL PROBABILITY	99.93%	99.19%	97.35%	96.38%	95.96%	95.96%
± 1 STANDARD ERROR	0.02%	0.07%	0.15%	0.21%	0.25%	0.25%

Dual-Chamber Pacemakers

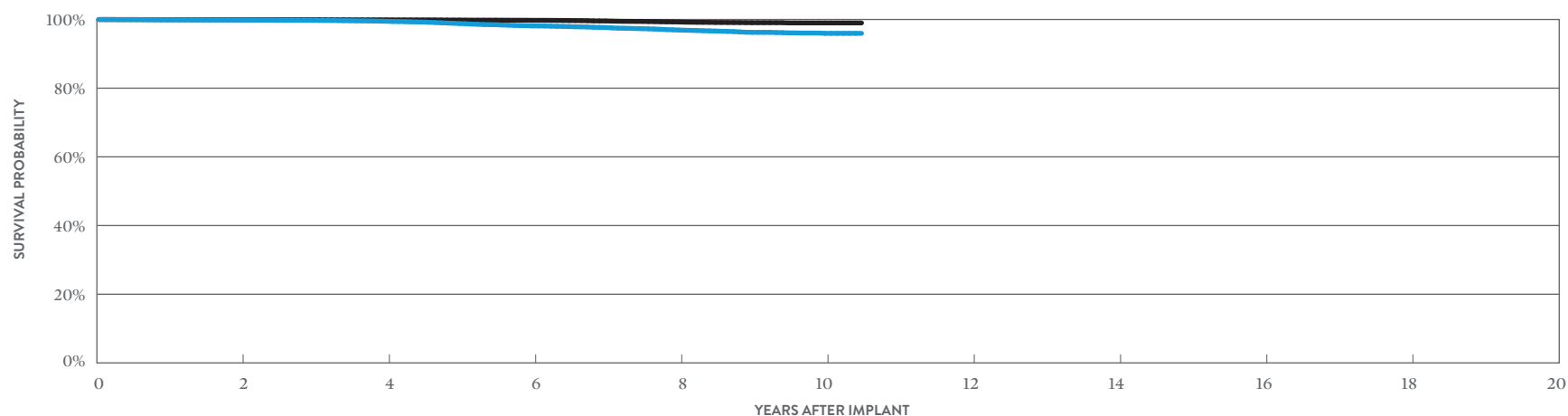
CUSTOMER REPORTED PERFORMANCE DATA

Zephyr™ XL DR

MODEL 5826

US Regulatory Approval	March 2007
Registered US Implants	112,224
Estimated Active US Implants	33,698
Estimated Longevity	11.7 Years
Normal Battery Depletion	568
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	<0.01%	19	0.02%
Electrical Interconnect	4	<0.01%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	14	0.01%
Mechanical	1	<0.01%	9	<0.01%
Possible Early Battery Depletion	0	0.00%	3	<0.01%
Other	1	<0.01%	122	0.11%
Total	7	<0.01%	167	0.15%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 126 MONTHS
SURVIVAL PROBABILITY	99.84%	99.48%	98.13%	96.93%	95.96%	95.96%
± 1 STANDARD ERROR	0.01%	0.02%	0.05%	0.08%	0.11%	0.13%
SAMPLE SIZE	92,170	72,260	55,060	33,560	6,780	350

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 126 MONTHS
SURVIVAL PROBABILITY	99.93%	99.89%	99.75%	99.26%	98.99%	98.99%
± 1 STANDARD ERROR	0.01%	0.01%	0.02%	0.04%	0.06%	0.06%

Dual-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

Zephyr™ XL DR

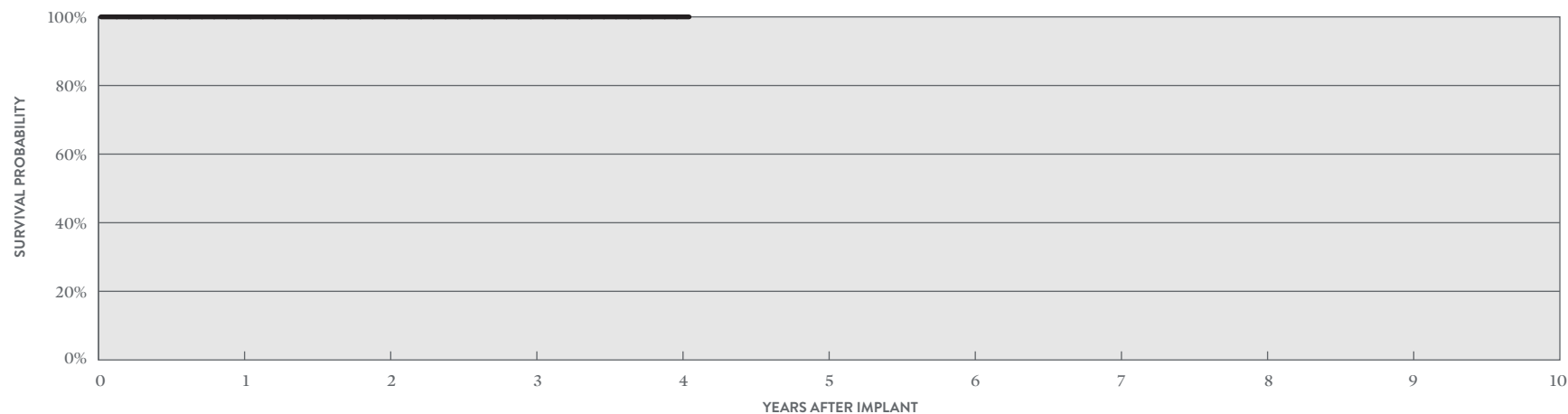
MODEL 5826

US Regulatory Approval	March 2007
Number of Devices Enrolled in Study	1,516
Active Devices Enrolled in Study	14
Cumulative Months of Follow-up	47,883
Estimated Longevity	11.7 Years

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.07%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	1	0.07%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	4	AT 49 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%	0.00%
SAMPLE SIZE	1,440	1,270	900	360	70

Dual-Chamber Pacemakers

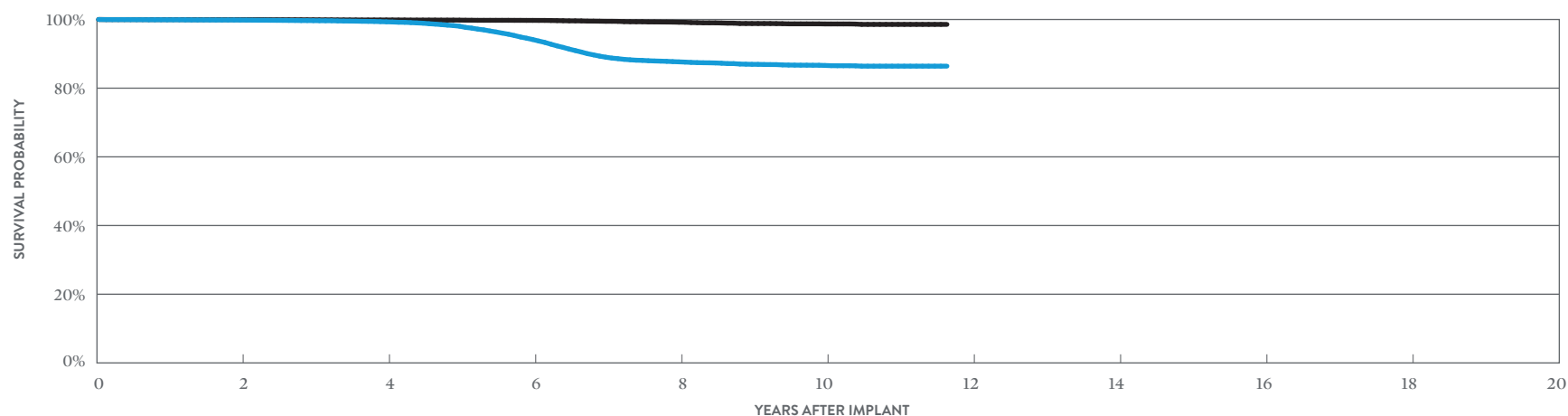
CUSTOMER REPORTED PERFORMANCE DATA

Victory™ XL DR

MODEL 5816

US Regulatory Approval	December 2005
Registered US Implants	62,690
Estimated Active US Implants	12,050
Estimated Longevity	11.7 Years
Normal Battery Depletion	1,496
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	25	0.04%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	8	0.01%
Mechanical	0	0.00%	9	0.01%
Possible Early Battery Depletion	0	0.00%	5	<0.01%
Other	1	<0.01%	82	0.13%
Total	3	<0.01%	129	0.21%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 140 MONTHS
SURVIVAL PROBABILITY	99.83%	99.32%	94.17%	87.66%	86.61%	86.41%
± 1 STANDARD ERROR	0.02%	0.04%	0.12%	0.19%	0.20%	0.21%
SAMPLE SIZE	52,020	40,740	31,840	20,260	9,600	250

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 140 MONTHS
SURVIVAL PROBABILITY	99.95%	99.86%	99.74%	99.19%	98.70%	98.57%
± 1 STANDARD ERROR	0.01%	0.02%	0.03%	0.05%	0.08%	0.10%

Dual-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

Victory™ XL DR

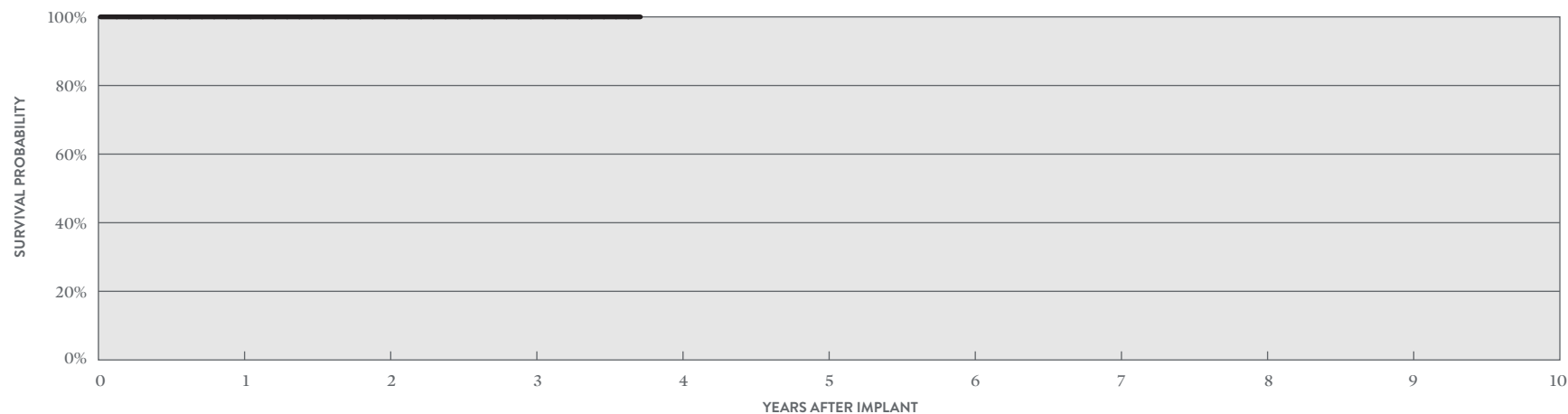
MODEL 5816

US Regulatory Approval	December 2005
Number of Devices Enrolled in Study	332
Active Devices Enrolled in Study	0
Cumulative Months of Follow-up	10,627
Estimated Longevity	11.7 Years

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	AT 45 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%
SAMPLE SIZE	320	280	200	50

Dual-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

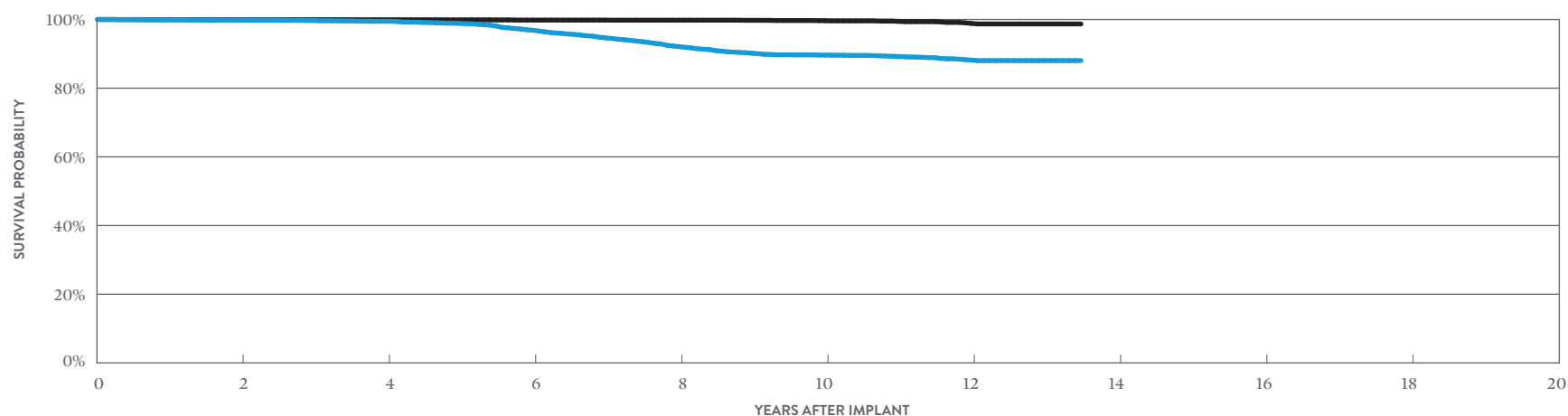
Verity ADx™ XL DR MODEL 5356

Verity ADx™ XL DR M/S MODEL 5357M/S

Verity ADx™ XL DC MODEL 5256

US Regulatory Approval	May 2003
Registered US Implants	17,366
Estimated Active US Implants	3,864
Estimated Longevity	6.9 Years
Normal Battery Depletion	306
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	10	0.06%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	0	0.00%	1	<0.01%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	0	0.00%	10	0.06%
Total	1	<0.01%	22	0.13%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 162 MONTHS
SURVIVAL PROBABILITY	99.83%	99.47%	96.87%	92.10%	89.64%	88.17%	88.03%
± 1 STANDARD ERROR	0.03%	0.06%	0.18%	0.31%	0.37%	0.43%	0.45%
SAMPLE SIZE	14,250	11,040	8,290	6,120	4,300	1,890	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 162 MONTHS
SURVIVAL PROBABILITY	99.95%	99.91%	99.82%	99.79%	99.65%	98.86%	98.70%
± 1 STANDARD ERROR	0.02%	0.02%	0.04%	0.05%	0.07%	0.19%	0.25%

Dual-Chamber Pacemakers

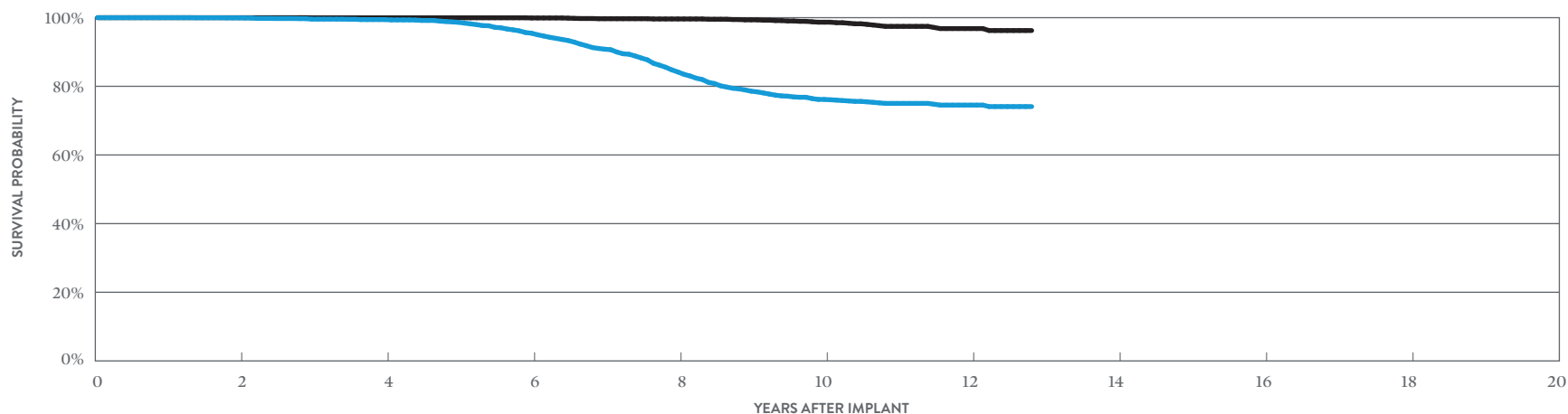
CUSTOMER REPORTED PERFORMANCE DATA

Integrity ADx™ DR

MODEL 5366

US Regulatory Approval	May 2003
Registered US Implants	8,083
Estimated Active US Implants	1,099
Estimated Longevity	6.9 Years
Normal Battery Depletion	320
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	9	0.11%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	2	0.02%
Mechanical	0	0.00%	1	0.01%
Possible Early Battery Depletion	0	0.00%	1	0.01%
Other	0	0.00%	14	0.17%
Total	0	0.00%	27	0.33%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 154 MONTHS
SURVIVAL PROBABILITY	99.94%	99.44%	95.47%	84.17%	76.18%	74.53%	74.10%
± 1 STANDARD ERROR	0.03%	0.10%	0.30%	0.58%	0.75%	0.82%	0.87%
SAMPLE SIZE	6,760	5,330	4,140	3,030	1,770	610	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 154 MONTHS
SURVIVAL PROBABILITY	100.00%	99.96%	99.91%	99.62%	98.68%	96.82%	96.25%
± 1 STANDARD ERROR	0.00%	0.02%	0.02%	0.10%	0.25%	0.52%	0.65%

Dual-Chamber Pacemakers

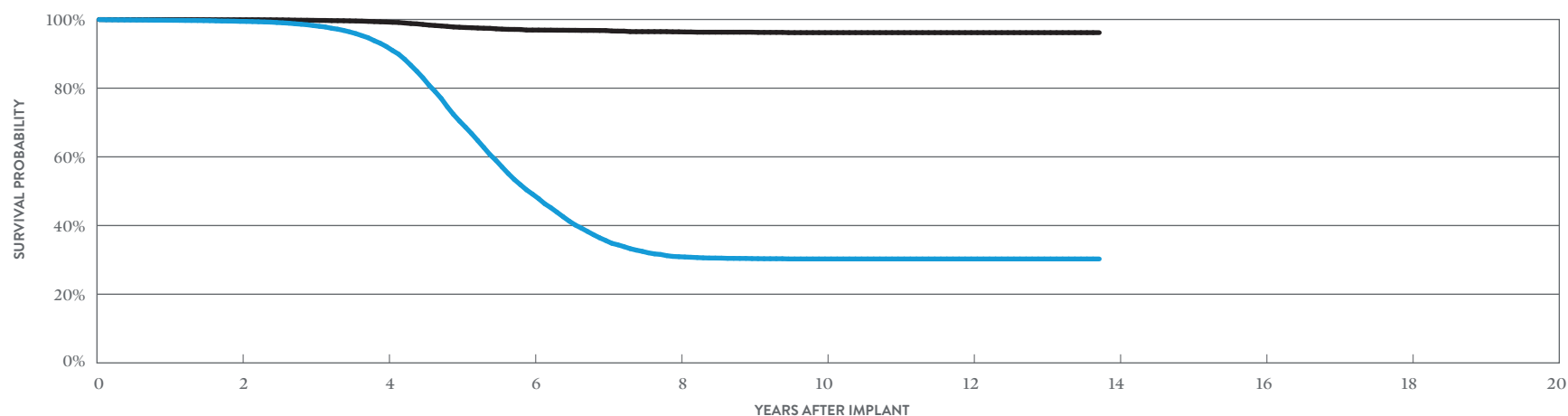
CUSTOMER REPORTED PERFORMANCE DATA

Identity ADx™ DR

MODEL 5380

US Regulatory Approval	March 2003
Registered US Implants	54,047
Estimated Active US Implants	3,032
Estimated Longevity	3.8 Years
Normal Battery Depletion	6,220
Number of US Advisories (see pg. 337)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	4	<0.01%	262	0.48%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	2	<0.01%
Mechanical	0	0.00%	6	0.01%
Possible Early Battery Depletion	0	0.00%	11	0.02%
Other	0	0.00%	17	0.03%
Total	5	<0.01%	298	0.55%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 165 MONTHS
SURVIVAL PROBABILITY	99.45%	92.12%	49.13%	30.92%	30.27%	30.27%	30.27%
± 1 STANDARD ERROR	0.03%	0.14%	0.32%	0.34%	0.34%	0.34%	0.34%
SAMPLE SIZE	43,620	31,500	13,450	4,770	2,840	1,680	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 165 MONTHS
SURVIVAL PROBABILITY	99.93%	99.26%	96.92%	96.41%	96.17%	96.17%	96.17%
± 1 STANDARD ERROR	0.01%	0.04%	0.12%	0.15%	0.17%	0.17%	0.17%

Dual-Chamber Pacemakers

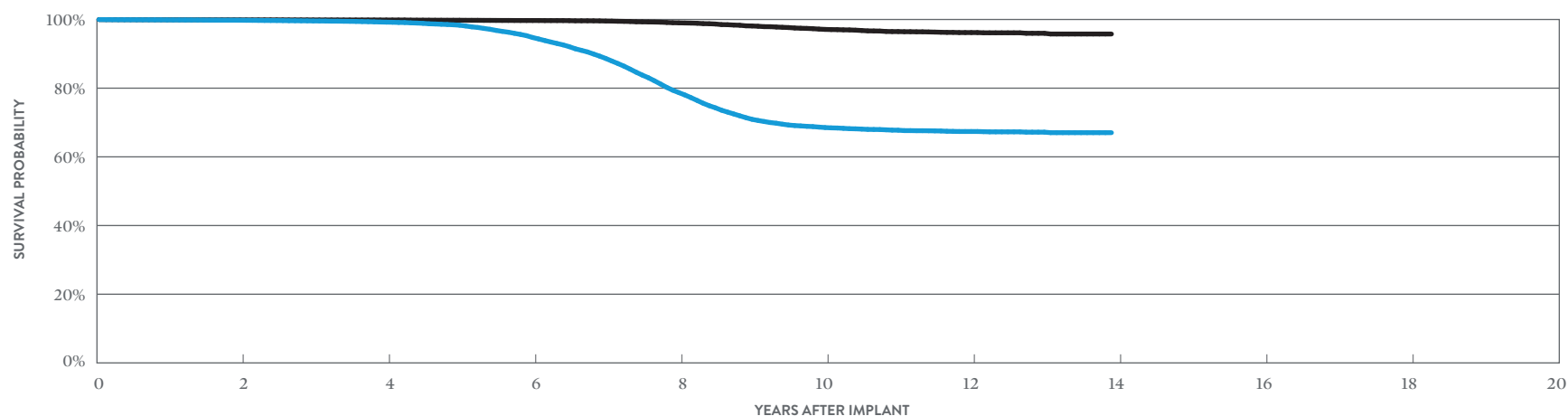
CUSTOMER REPORTED PERFORMANCE DATA

Identity ADx™ XL DR MODEL 5386

Identity ADx™ XL DC MODEL 5286

US Regulatory Approval	March 2003
Registered US Implants	67,384
Estimated Active US Implants	9,665
Estimated Longevity	6.9 Years
Normal Battery Depletion	3,302
Number of US Advisories (see pg. 337)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	132	0.20%
Electrical Interconnect	0	0.00%	2	<0.01%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	7	0.01%
Mechanical	0	0.00%	10	0.01%
Possible Early Battery Depletion	0	0.00%	6	<0.01%
Other	0	0.00%	106	0.16%
Total	2	<0.01%	263	0.39%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 167 MONTHS
SURVIVAL PROBABILITY	99.78%	99.22%	94.75%	78.70%	68.53%	67.38%	67.04%
± 1 STANDARD ERROR	0.02%	0.04%	0.11%	0.23%	0.29%	0.31%	0.33%
SAMPLE SIZE	56,170	44,130	33,580	23,150	11,470	4,550	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 167 MONTHS
SURVIVAL PROBABILITY	99.90%	99.85%	99.70%	99.02%	97.11%	96.19%	95.77%
± 1 STANDARD ERROR	0.01%	0.02%	0.03%	0.06%	0.13%	0.18%	0.26%

Dual-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

Identity ADx™ XL DR

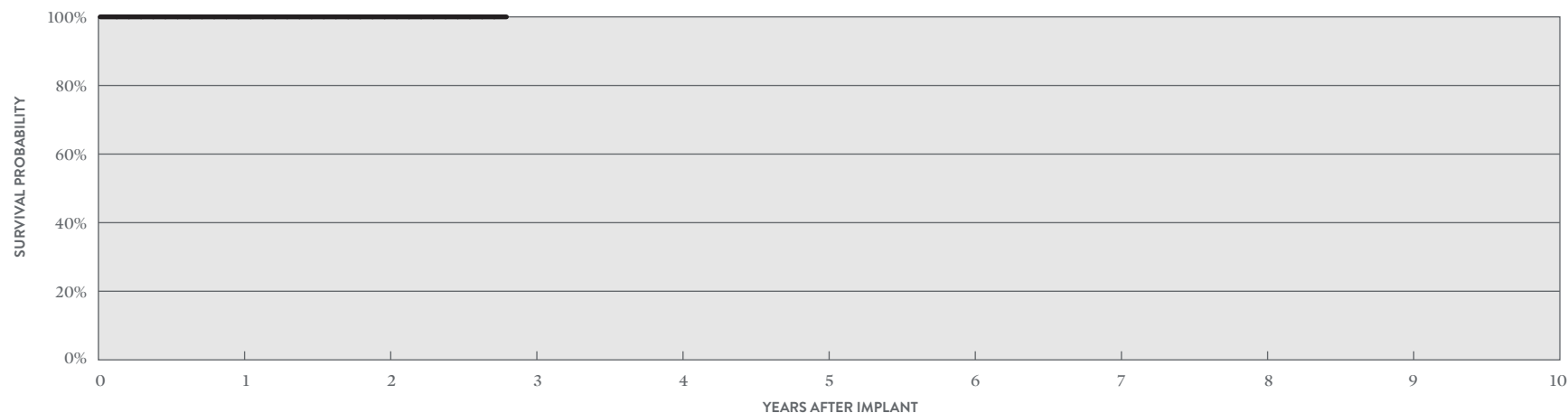
MODEL 5386

US Regulatory Approval	March 2003
Number of Devices Enrolled in Study	102
Active Devices Enrolled in Study	0
Cumulative Months of Follow-up	3,251
Estimated Longevity	6.9 Years

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	AT 34 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%
SAMPLE SIZE	100	80	50

Dual-Chamber Pacemakers

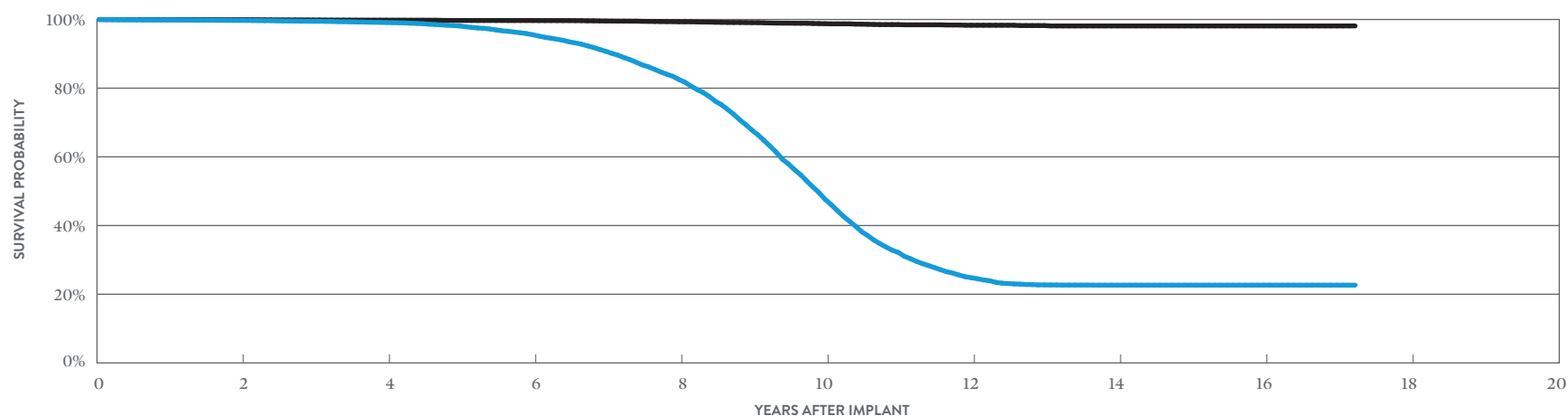
CUSTOMER REPORTED PERFORMANCE DATA

Integrity AFx™ DR

MODELS 5342 & 5346

US Regulatory Approval	(5342) April 2000 (5346) July 2001
Registered US Implants	47,442
Estimated Active US Implants	1,624
Estimated Longevity	6.3 Years
Normal Battery Depletion	4,611
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	92	0.19%
Electrical Interconnect	3	<0.01%	1	<0.01%
Battery	0	0.00%	2	<0.01%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	1	<0.01%	3	<0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	6	0.01%
Total	6	0.01%	104	0.22%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 207 MONTHS
SURVIVAL PROBABILITY	99.73%	99.12%	95.56%	82.53%	47.64%	24.80%	22.64%	22.64%	22.64%
± 1 STANDARD ERROR	0.03%	0.05%	0.12%	0.25%	0.41%	0.38%	0.38%	0.38%	0.38%
SAMPLE SIZE	39,960	32,370	24,810	16,380	7,920	3,260	1,760	1,150	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 207 MONTHS
SURVIVAL PROBABILITY	99.92%	99.81%	99.70%	99.33%	98.77%	98.32%	98.12%	98.12%	98.12%
± 1 STANDARD ERROR	0.01%	0.02%	0.03%	0.05%	0.09%	0.14%	0.17%	0.17%	0.17%

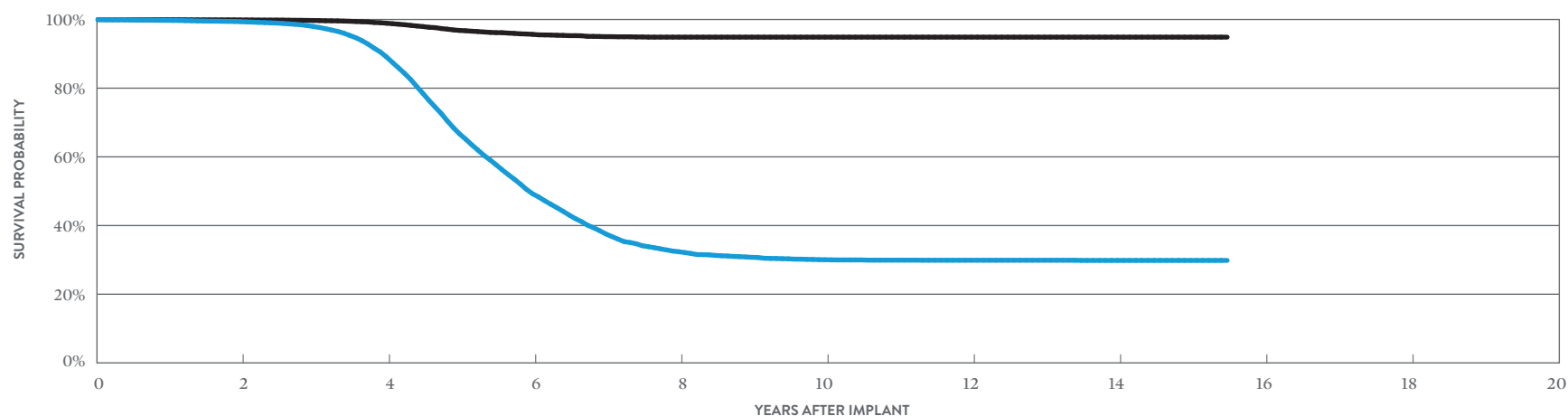
Dual-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

Identity™ MODEL 5370

US Regulatory Approval	November 2001
Registered US Implants	58,366
Estimated Active US Implants	1,956
Estimated Longevity	3.8 Years
Normal Battery Depletion	6,084
Number of US Advisories (see pg. 337)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	3	<0.01%	398	0.68%
Electrical Interconnect	2	<0.01%	2	<0.01%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	5	<0.01%
Possible Early Battery Depletion	0	0.00%	12	0.02%
Other	0	0.00%	12	0.02%
Total	5	<0.01%	430	0.74%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	AT 186 MONTHS
SURVIVAL PROBABILITY	99.37%	89.11%	49.24%	32.37%	30.04%	29.89%	29.84%	29.84%
± 1 STANDARD ERROR	0.03%	0.15%	0.33%	0.37%	0.38%	0.38%	0.38%	0.38%
SAMPLE SIZE	47,520	34,030	12,120	3,890	2,470	1,900	1,190	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	AT 186 MONTHS
SURVIVAL PROBABILITY	99.87%	98.91%	95.67%	94.86%	94.86%	94.86%	94.86%	94.86%
± 1 STANDARD ERROR	0.01%	0.05%	0.14%	0.19%	0.19%	0.19%	0.19%	0.19%

Dual-Chamber Pacemakers

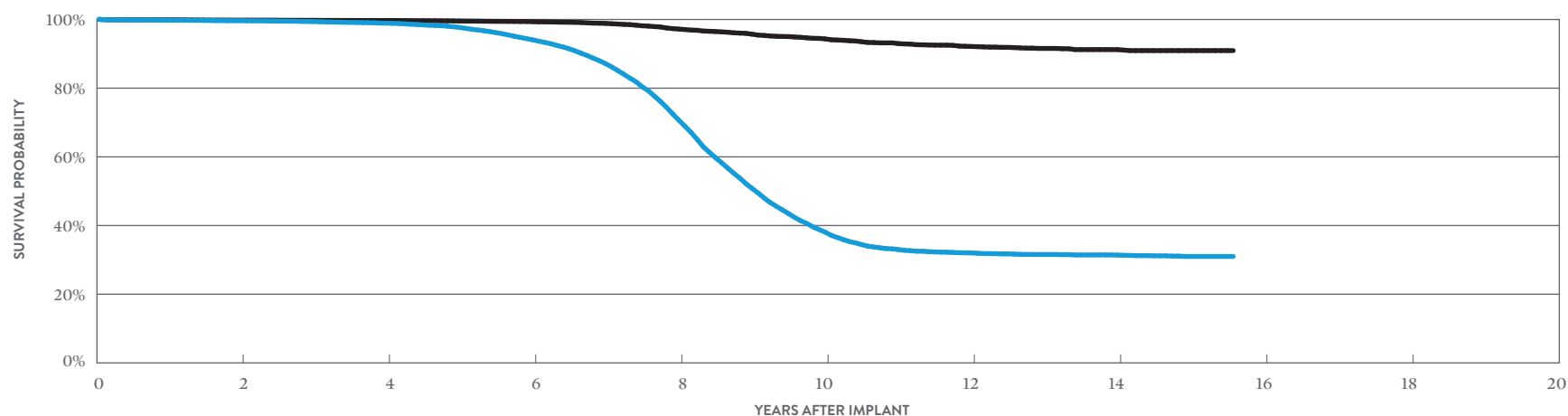
CUSTOMER REPORTED PERFORMANCE DATA

Identity™ XL

MODEL 5376

US Regulatory Approval	November 2001
Registered US Implants	51,521
Estimated Active US Implants	3,357
Estimated Longevity	6.9 Years
Normal Battery Depletion	5,329
Number of US Advisories (see pg. 337)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	311	0.60%
Electrical Interconnect	4	<0.01%	2	<0.01%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	12	0.02%
Mechanical	2	<0.01%	6	0.01%
Possible Early Battery Depletion	0	0.00%	5	<0.01%
Other	0	0.00%	95	0.18%
Total	8	0.02%	431	0.84%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	AT 187 MONTHS
SURVIVAL PROBABILITY	99.64%	98.92%	94.08%	70.66%	38.09%	32.01%	31.39%	31.00%
± 1 STANDARD ERROR	0.03%	0.05%	0.13%	0.29%	0.34%	0.34%	0.35%	0.36%
SAMPLE SIZE	43,560	34,720	26,340	17,440	8,280	3,830	1,720	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	AT 187 MONTHS
SURVIVAL PROBABILITY	99.80%	99.71%	99.35%	97.21%	94.35%	92.14%	91.23%	90.94%
± 1 STANDARD ERROR	0.02%	0.03%	0.04%	0.11%	0.19%	0.28%	0.34%	0.37%

Dual-Chamber Pacemakers

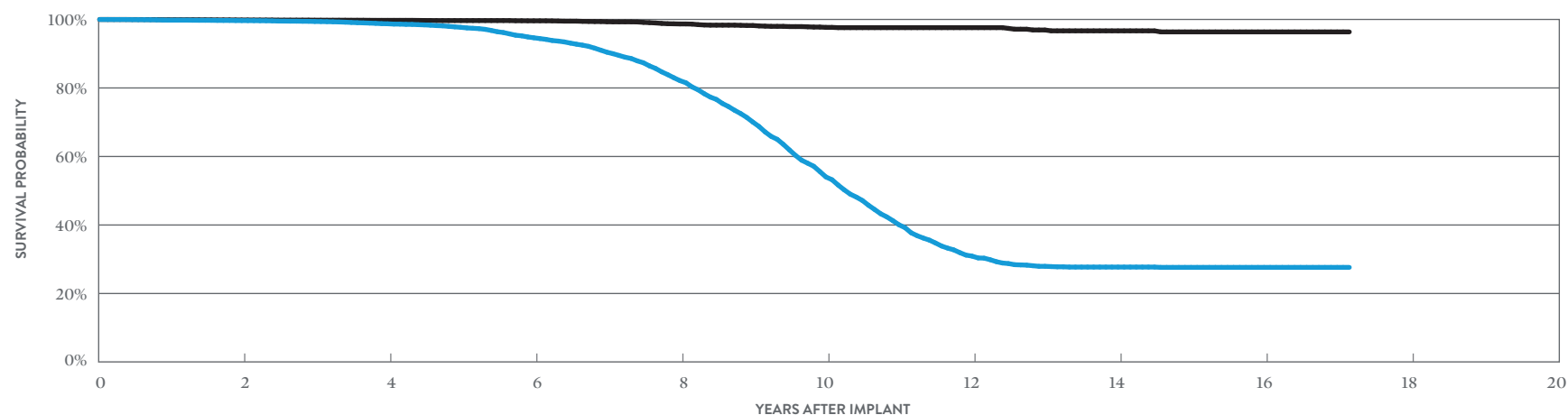
CUSTOMER REPORTED PERFORMANCE DATA

Entity™ DR MODEL 5326

Entity™ DC MODEL 5226

US Regulatory Approval	June 1999
Registered US Implants	21,828
Estimated Active US Implants	652
Estimated Longevity	6.3 Years
Normal Battery Depletion	1,546
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	<0.01%	65	0.30%
Electrical Interconnect	2	<0.01%	2	<0.01%
Battery	0	0.00%	1	<0.01%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	0	0.00%	3	0.01%
Total	3	0.01%	74	0.34%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 206 MONTHS
SURVIVAL PROBABILITY	99.66%	98.73%	94.64%	82.14%	54.02%	30.92%	27.72%	27.63%	27.63%
± 1 STANDARD ERROR	0.04%	0.09%	0.20%	0.41%	0.66%	0.71%	0.70%	0.70%	0.70%
SAMPLE SIZE	17,830	14,030	10,260	6,310	2,990	1,280	740	480	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 206 MONTHS
SURVIVAL PROBABILITY	99.85%	99.74%	99.60%	98.68%	97.68%	97.60%	96.67%	96.36%	96.36%
± 1 STANDARD ERROR	0.03%	0.04%	0.05%	0.13%	0.20%	0.22%	0.39%	0.45%	0.45%

Dual-Chamber Pacemakers

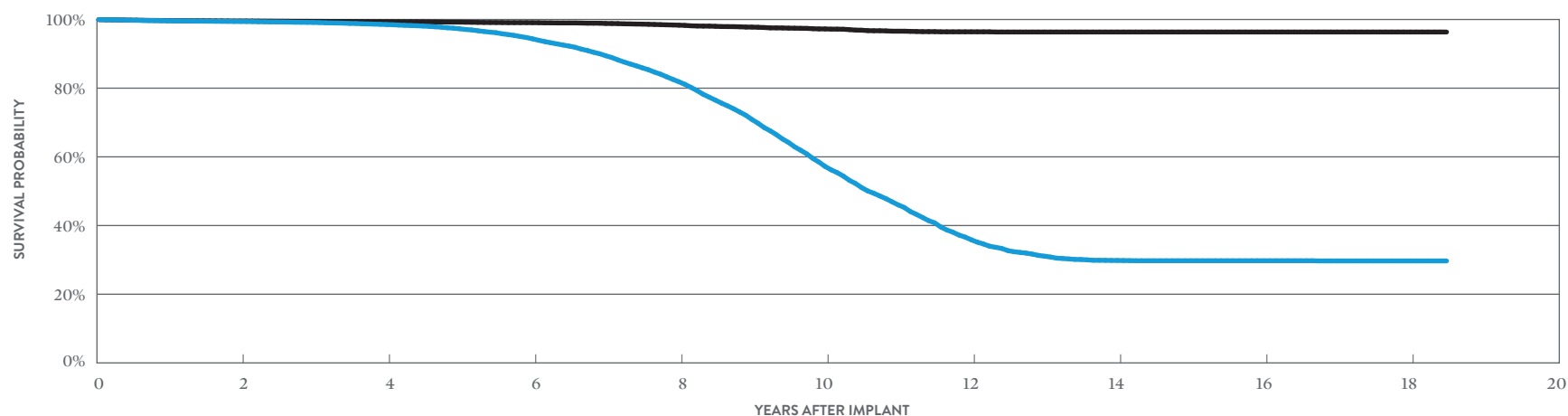
CUSTOMER REPORTED PERFORMANCE DATA

Affinity™ DR MODELS 5330 & 5331

Affinity™ DC MODEL 5230

US Regulatory Approval	(5330) January 1999 (5230/5331) June 1999
Registered US Implants	65,717
Estimated Active US Implants	1,990
Estimated Longevity	6.3 Years
Normal Battery Depletion	4,557
Number of US Advisories (see pg. 340)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	5	<0.01%	283	0.43%
Electrical Interconnect	9	0.01%	13	0.02%
Battery	0	0.00%	6	<0.01%
Software/Firmware	0	0.00%	2	<0.01%
Mechanical	0	0.00%	5	<0.01%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	1	<0.01%	5	<0.01%
Total	15	0.02%	315	0.48%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	18	AT 222 MONTHS
SURVIVAL PROBABILITY	99.41%	98.53%	94.36%	81.86%	57.21%	35.84%	29.83%	29.75%	29.70%	29.70%
± 1 STANDARD ERROR	0.03%	0.05%	0.11%	0.23%	0.37%	0.43%	0.43%	0.43%	0.43%	0.43%
SAMPLE SIZE	54,830	43,850	32,640	20,190	9,440	4,150	2,450	1,780	870	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	18	AT 222 MONTHS
SURVIVAL PROBABILITY	99.56%	99.35%	99.07%	98.34%	97.25%	96.41%	96.35%	96.35%	96.35%	96.35%
± 1 STANDARD ERROR	0.03%	0.03%	0.04%	0.07%	0.12%	0.17%	0.17%	0.17%	0.17%	0.17%

SUMMARY INFORMATION
Dual-Chamber
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Dual-Chamber Pacemakers

Survival Probability Summary

INCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
PM2272	Assurity MRI ⁺ *										
PM2160	Endurity [™] DR	99.84%	99.78%	99.73%							
PM2240	Assurity [™] DR RF	99.95%	99.92%	99.90%							
PM2210	Accent [™] DR RF	99.92%	99.86%	99.78%	99.62%	99.35%	98.75%	97.21%	97.18%		
PM2110	Accent [™] DR	99.96%	99.91%	99.83%	99.64%	99.42%	98.85%	97.47%	97.42%		
5820	Zephyr [™] DR	99.84%	99.75%	99.04%	93.70%	81.97%	78.54%	77.66%	77.29%	77.21%	77.04%
5810	Victory [™] DR	99.87%	99.75%	98.68%	89.39%	68.01%	52.59%	46.38%	45.60%	45.37%	45.21%
5826	Zephyr [™] XL DR	99.91%	99.84%	99.74%	99.48%	98.81%	98.13%	97.67%	96.93%	96.25%	95.96%
5816	Victory [™] XL DR	99.91%	99.83%	99.66%	99.32%	98.04%	94.17%	89.02%	87.66%	86.94%	86.61%
5356/5357/5256	Verity ADx [™] XL DR/ DR(M/S) / DC	99.89%	99.83%	99.69%	99.47%	98.85%	96.87%	94.65%	92.10%	90.17%	89.64%
5366	Integrity ADx [™] XL DR	100.00%	99.94%	99.57%	99.44%	98.65%	95.47%	90.81%	84.17%	78.58%	76.18%
5380	Identity ADx [™] DR	99.77%	99.45%	98.23%	92.12%	70.42%	49.13%	35.64%	30.92%	30.37%	30.27%
5386/5286	Identity ADx [™] XL DR/DC	99.88%	99.78%	99.57%	99.22%	98.33%	94.75%	88.65%	78.70%	70.93%	68.53%
5342/5346	Integrity AFx [™] DR	99.87%	99.73%	99.48%	99.12%	98.13%	95.56%	90.73%	82.53%	67.78%	47.64%
5370	Identity [™]	99.75%	99.37%	97.95%	89.11%	66.70%	49.24%	37.56%	32.37%	30.75%	30.04%
5376	Identity [™] XL	99.79%	99.64%	99.38%	98.92%	97.70%	94.08%	87.11%	70.66%	50.83%	38.09%
5326/5226	Entity [™] DR/DC	99.79%	99.66%	99.40%	98.73%	97.70%	94.64%	90.44%	82.14%	70.03%	54.02%
5330/5331/5230	Affinity [™] DR/DC	99.63%	99.41%	99.14%	98.53%	97.30%	94.36%	89.44%	81.86%	70.87%	57.21%

*No survival probability is stated at one year due to the device not meeting the required minimum sample size of 200 U.S. implants with 12 consecutive months of data. Please refer to the individual graphs for data up to one year.

Dual-Chamber Pacemakers

Survival Probability Summary

EXCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
PM2272	Assurity MRI™*										
PM2160	Endurity™ DR	99.84%	99.81%	99.81%							
PM2240	Assurity™ DR RF	99.95%	99.94%	99.93%							
PM2210	Accent™ DR RF	99.95%	99.90%	99.84%	99.80%	99.77%	99.74%	99.69%	99.68%		
PM2110	Accent™ DR	99.97%	99.95%	99.93%	99.93%	99.92%	99.88%	99.88%	99.88%		
5820	Zephyr™ DR	99.97%	99.96%	99.93%	99.64%	99.30%	98.97%	98.68%	98.56%	98.46%	98.24%
5810	Victory™ DR	99.98%	99.93%	99.69%	99.19%	97.76%	97.35%	96.81%	96.38%	96.16%	95.96%
5826	Zephyr™ XL DR	99.96%	99.93%	99.91%	99.89%	99.83%	99.75%	99.57%	99.26%	99.07%	98.99%
5816	Victory™ XL DR	99.97%	99.95%	99.91%	99.86%	99.81%	99.74%	99.47%	99.19%	98.82%	98.70%
5356/5357/5256	Verity ADx™ XL DR/ DR(M/S) / DC	99.96%	99.95%	99.93%	99.91%	99.89%	99.82%	99.82%	99.79%	99.75%	99.65%
5366	Integrity ADx™ XL DR	100.00%	100.00%	99.96%	99.96%	99.96%	99.91%	99.69%	99.62%	99.36%	98.68%
5380	Identity ADx™ DR	99.96%	99.93%	99.75%	99.26%	97.74%	96.92%	96.79%	96.41%	96.30%	96.17%
5386/5286	Identity ADx™ XL DR/DC	99.92%	99.90%	99.87%	99.85%	99.78%	99.70%	99.55%	99.02%	98.11%	97.11%
5342/5346	Integrity AFx™ DR	99.96%	99.92%	99.86%	99.81%	99.72%	99.70%	99.56%	99.33%	99.08%	98.77%
5370	Identity™	99.93%	99.87%	99.71%	98.91%	96.81%	95.67%	95.00%	94.86%	94.86%	94.86%
5376	Identity™ XL	99.90%	99.80%	99.76%	99.71%	99.55%	99.35%	98.86%	97.21%	95.72%	94.35%
5326/5226	Entity™ DR/DC	99.91%	99.85%	99.79%	99.74%	99.67%	99.60%	99.32%	98.68%	98.23%	97.68%
5330/5331/5230	Affinity™ DR/DC	99.69%	99.56%	99.46%	99.35%	99.23%	99.07%	98.84%	98.34%	97.78%	97.25%

*No survival probability is stated at one year due to the device not meeting the required minimum sample size of 200 U.S. implants with 12 consecutive months of data. Please refer to the individual graphs for data up to one year.

Dual-Chamber Pacemakers

US Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/ FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM2272	Assurity MRI™	36,177	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM2160	Endurity™ DR	8,929	0.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM2240	Assurity™ DR RF	168,347	0.20%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	2	<0.01%	0	0.00%	5	<0.01%
PM2210	Accent™ DR RF	243,042	2.70%	15	<0.01%	7	<0.01%	0	0.00%	0	0.00%	0	0.00%	7	<0.01%	5	<0.01%	34	0.01%
PM2110	Accent™ DR	48,906	2.70%	2	<0.01%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	<0.01%
5820	Zephyr™ DR	53,978	8.00%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	<0.01%
5810	Victory™ DR	26,311	16.80%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
5826	Zephyr™ XL DR	112,224	5.90%	1	<0.01%	4	<0.01%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	7	<0.01%
5816	Victory™ XL DR	62,690	11.50%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	3	<0.01%
5356/5357/5256	Verity ADx™ XL DR/ DR(M/S) / DC	17,366	6.60%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
5366	Integrity ADx™ XL DR	8,083	10.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5380	Identity ADx™ DR	54,047	15.60%	4	<0.01%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	5	<0.01%
5386/5286	Identity ADx™ XL DR/DC	67,384	13.10%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	<0.01%
5342/5346	Integrity AFx™ DR	47,442	14.20%	2	<0.01%	3	<0.01%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	6	0.01%
5370	Identity™	58,366	13.70%	3	<0.01%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	5	<0.01%
5376	Identity™ XL	51,521	17.50%	2	<0.01%	4	<0.01%	0	0.00%	0	0.00%	2	<0.01%	0	0.00%	0	0.00%	8	0.02%
5326/5226	Entity™ DR/DC	21,828	11.10%	1	<0.01%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	3	0.01%
5330/5331/5230	Affinity™ DR/DC	65,717	10.90%	5	<0.01%	9	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	15	0.02%

Definitions of malfunction categories can be found on [pages 5-6](#).

Dual-Chamber Pacemakers

US Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM2272	Assurity MRI	36,177	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	1	<0.01%
PM2160	Endurity DR	8,929	0.30%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	5	0.06%	0	0.00%	2	0.02%	7	0.08%
PM2240	Assurity DR RF	168,347	0.20%	3	<0.01%	0	0.00%	0	0.00%	1	<0.01%	22	0.01%	0	0.00%	7	<0.01%	33	0.02%
PM2210	Accent DR RF	243,042	2.70%	44	0.02%	31	0.01%	0	0.00%	4	<0.01%	20	<0.01%	22	<0.01%	37	0.02%	158	0.07%
PM2110	Accent DR	48,906	2.70%	3	<0.01%	0	0.00%	0	0.00%	3	<0.01%	5	0.01%	2	<0.01%	0	0.00%	13	0.03%
5820	Zephyr DR	53,978	8.00%	34	0.06%	0	0.00%	0	0.00%	9	0.02%	2	<0.01%	1	<0.01%	73	0.14%	119	0.22%
5810	Victory DR	26,311	16.80%	89	0.34%	0	0.00%	0	0.00%	8	0.03%	2	<0.01%	17	0.06%	37	0.14%	153	0.58%
5826	Zephyr XL DR	112,224	5.90%	19	0.02%	0	0.00%	0	0.00%	14	0.01%	9	<0.01%	3	<0.01%	122	0.11%	167	0.15%
5816	Victory XL DR	62,690	11.50%	25	0.04%	0	0.00%	0	0.00%	8	0.01%	9	0.01%	5	<0.01%	82	0.13%	129	0.21%
5356/5357/5256	Verity ADx XL DR/DR(M/S) / DC	17,366	6.60%	10	0.06%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	1	<0.01%	10	0.06%	22	0.13%
5366	Integrity ADx XL DR	8,083	10.90%	9	0.11%	0	0.00%	0	0.00%	2	0.02%	1	0.01%	1	0.01%	14	0.17%	27	0.33%
5380	Identity ADx DR	54,047	15.60%	262	0.48%	0	0.00%	0	0.00%	2	<0.01%	6	0.01%	11	0.02%	17	0.03%	298	0.55%
5386/5286	Identity ADx XL DR/DC	67,384	13.10%	132	0.20%	2	<0.01%	0	0.00%	7	0.01%	10	0.01%	6	<0.01%	106	0.16%	263	0.39%
5342/5346	Integrity AFx DR	47,442	14.20%	92	0.19%	1	<0.01%	2	<0.01%	0	0.00%	3	<0.01%	0	0.00%	6	0.01%	104	0.22%
5370	Identity	58,366	13.70%	398	0.68%	2	<0.01%	0	0.00%	1	<0.01%	5	<0.01%	12	0.02%	12	0.02%	430	0.74%
5376	Identity XL	51,521	17.50%	311	0.60%	2	<0.01%	0	0.00%	12	0.02%	6	0.01%	5	<0.01%	95	0.18%	431	0.84%
5326/5226	Entity DR/DC	21,828	11.10%	65	0.30%	2	<0.01%	1	<0.01%	1	<0.01%	1	<0.01%	1	<0.01%	3	0.01%	74	0.34%
5330/5331/5230	Affinity DR/DC	65,717	10.90%	283	0.43%	13	0.02%	6	<0.01%	2	<0.01%	5	<0.01%	1	<0.01%	5	<0.01%	315	0.48%

Definitions of malfunction categories can be found on [pages 5-6](#).

Dual-Chamber Pacemakers

Worldwide Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM2272	Assurity MRI [™]	147,286	0.18%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
PM2160	Endurity [™] DR	55,296	0.63%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM2240	Assurity [™] DR RF	183,118	1.79%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	2	<0.01%	0	0.00%	5	<0.01%	5	<0.01%
PM2210	Accent [™] DR RF	246,767	5.21%	15	<0.01%	7	<0.01%	0	0.00%	0	0.00%	0	0.00%	6	<0.01%	5	<0.01%	33	0.01%	33	0.01%
PM2110	Accent [™] DR	49,732	5.06%	2	<0.01%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	4	<0.01%	4	<0.01%

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM2272	Assurity MRI [™]	147,286	0.18%	1	<0.01%	0	0.00%	0	0.00%	1	<0.01%	3	<0.01%	1	<0.01%	0	0.00%	6	<0.01%	6	<0.01%
PM2160	Endurity [™] DR	55,296	0.63%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	8	0.01%	0	0.00%	3	<0.01%	12	0.02%	12	0.02%
PM2240	Assurity [™] DR RF	183,118	1.79%	5	<0.01%	0	0.00%	0	0.00%	1	<0.01%	25	0.01%	0	0.00%	6	<0.01%	37	0.02%	37	0.02%
PM2210	Accent [™] DR RF	246,767	5.21%	47	0.02%	32	0.01%	0	0.00%	4	<0.01%	20	<0.01%	22	<0.01%	36	0.01%	161	0.07%	161	0.07%
PM2110	Accent [™] DR	49,732	5.06%	3	<0.01%	0	0.00%	0	0.00%	3	<0.01%	5	0.01%	2	<0.01%	0	0.00%	13	0.03%	13	0.03%

Dual-Chamber Pacemakers

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	LOSS OF TELEMETRY		PERICARDIAL EFFUSION		PREMATURE BATTERY DEPLETION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM2210	1,774	323	55,167	0	0.00%	0	0.00%	1	0.06%	0	0.00%	1	0.06%
PM2110	227	61	9,045	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5820	283	8	7,892	0	0.00%	0	0.00%	0	0.00%	1	0.35%	1	0.35%
5826	1,516	14	47,883	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5816	332	0	10,627	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5386	102	0	3,251	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

A list of of complications can be found on [page 12](#).

Dual-Chamber Pacemakers

Actively Monitored Study Data Summary

MALFUNCTIONS WITH COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/ FIRM-WARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM2210	Fortify™ VR	1,774	4.20%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM2110	Current™ + VR	227	3.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5820	Current™ VR RF	283	17.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5826	Current™ + VR	1,516	6.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5816	Current™ + VR	332	5.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5386	Current™ VR RF	102	2.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

MALFUNCTIONS WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/ FIRM-WARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM2210	Fortify™ VR	1,774	4.20%	1	0.06%	1	0.06%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.11%
PM2110	Current™ + VR	227	3.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5820	Current™ VR RF	283	17.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5826	Current™ + VR	1,516	6.90%	1	0.07%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.07%
5816	Current™ + VR	332	5.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5386	Current™ VR RF	102	2.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

Definitions of malfunction categories can be found on [pages 5-6](#).

Single-Chamber Pacemakers

Single-Chamber Pacemakers

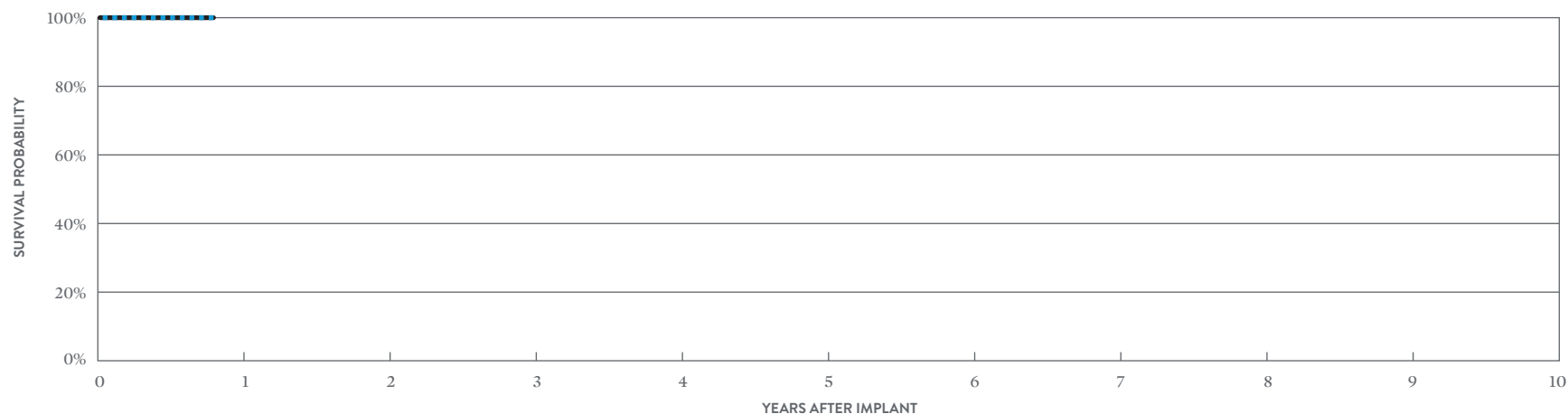
CUSTOMER REPORTED PERFORMANCE DATA

Assurity MRI™

MODEL PM1272

US Regulatory Approval	January 2017
Registered US Implants	4,415
Estimated Active US Implants	4,178
Estimated Longevity	13.7 Years
Normal Battery Depletion	0
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	AT 10 MONTHS
SURVIVAL PROBABILITY	100.00%
± 1 STANDARD ERROR	0.00%
SAMPLE SIZE	460

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	AT 10 MONTHS
SURVIVAL PROBABILITY	100.00%
± 1 STANDARD ERROR	0.00%

Single-Chamber Pacemakers

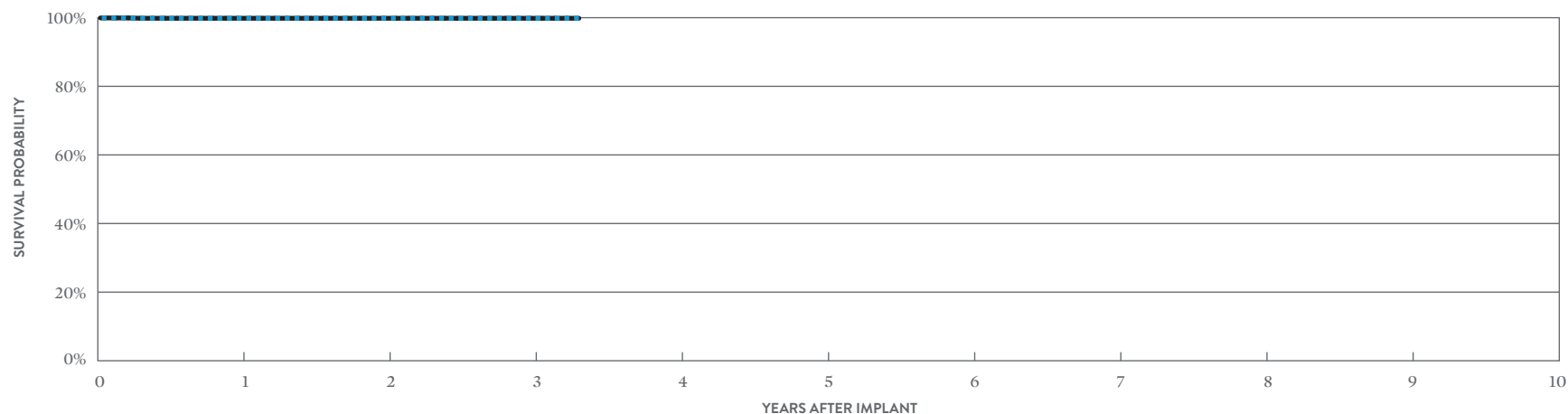
CUSTOMER REPORTED PERFORMANCE DATA

Endurity™ VR

MODEL PM1160

US Regulatory Approval	March 2014
Registered US Implants	2,455
Estimated Active US Implants	1,876
Estimated Longevity	14.6 Years
Normal Battery Depletion	0
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	0.04%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	1	0.04%
Total	0	0.00%	2	0.08%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 40 MONTHS
SURVIVAL PROBABILITY	99.83%	99.83%	99.83%	99.83%
± 1 STANDARD ERROR	0.09%	0.09%	0.09%	0.09%
SAMPLE SIZE	2,210	1,700	960	280

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 40 MONTHS
SURVIVAL PROBABILITY	99.83%	99.83%	99.83%	99.83%
± 1 STANDARD ERROR	0.09%	0.09%	0.09%	0.09%

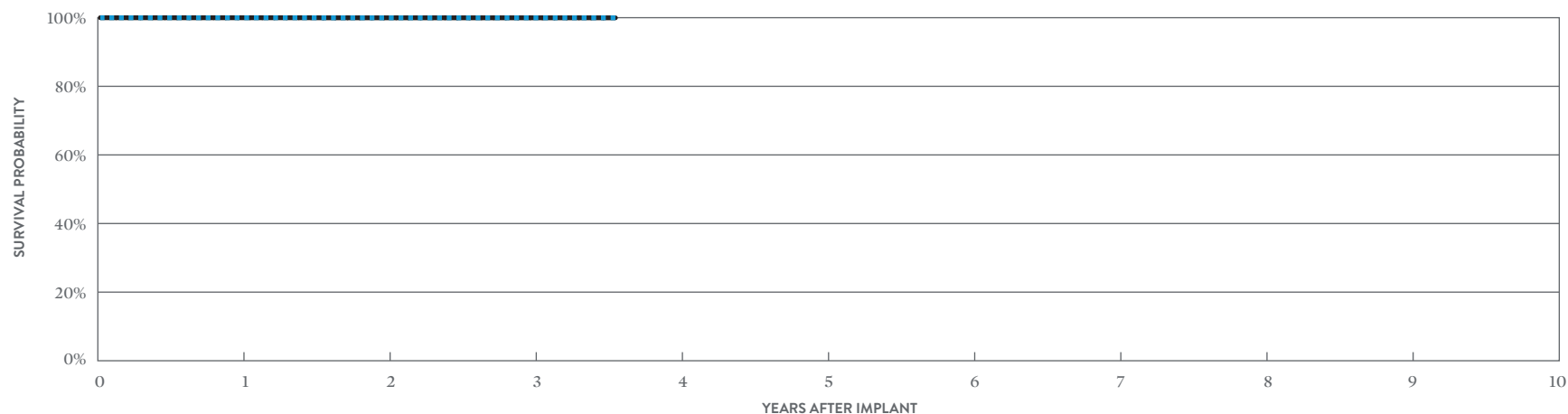
Single-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

Assurity™ VR MODEL PM1240

US Regulatory Approval	March 2014
Registered US Implants	25,978
Estimated Active US Implants	20,522
Estimated Longevity	14.1 Years
Normal Battery Depletion	0
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	2	<0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	2	<0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	4	0.02%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 43 MONTHS
SURVIVAL PROBABILITY	99.98%	99.96%	99.96%	99.96%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.01%
SAMPLE SIZE	22,310	14,180	6,080	230

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	AT 43 MONTHS
SURVIVAL PROBABILITY	99.98%	99.96%	99.96%	99.96%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.01%

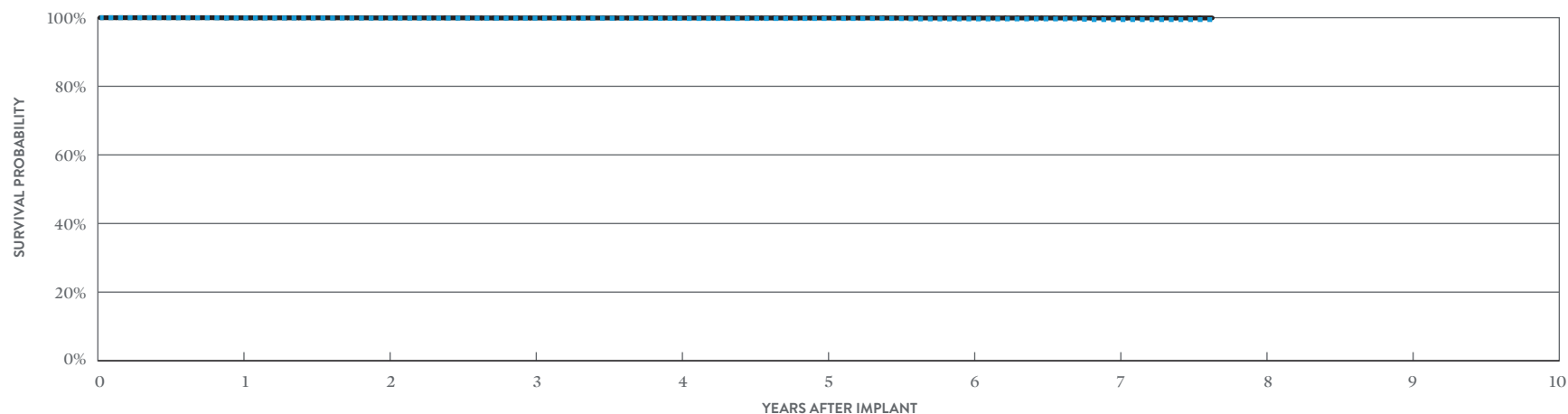
Single-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

Accent™ SR MODEL PM1110

US Regulatory Approval	July 2009
Registered US Implants	13,590
Estimated Active US Implants	7,427
Estimated Longevity	12.9 Years
Normal Battery Depletion	10
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	2	0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	0	0.00%	0	0.00%
Total	0	0.00%	4	0.03%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 92 MONTHS
SURVIVAL PROBABILITY	99.92%	99.87%	99.85%	99.79%	99.76%	99.64%	99.45%	99.45%
± 1 STANDARD ERROR	0.03%	0.03%	0.04%	0.05%	0.05%	0.08%	0.16%	0.16%
SAMPLE SIZE	12,520	10,780	9,470	7,820	5,640	3,450	1,600	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	AT 92 MONTHS
SURVIVAL PROBABILITY	99.96%	99.94%	99.92%	99.92%	99.92%	99.92%	99.92%	99.92%
± 1 STANDARD ERROR	0.02%	0.02%	0.03%	0.03%	0.03%	0.03%	0.03%	0.03%

Single-Chamber Pacemakers

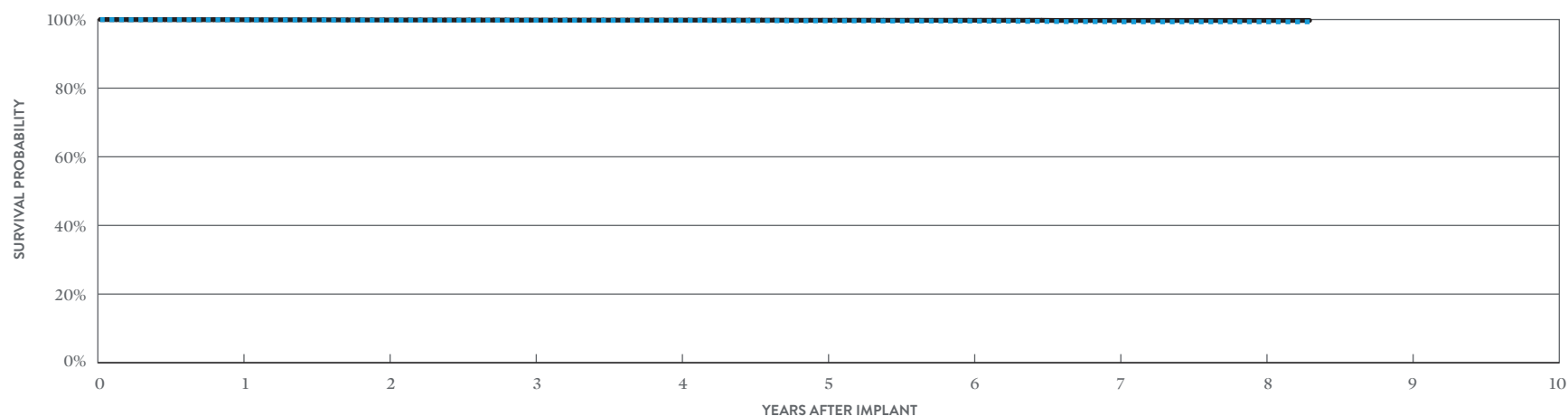
CUSTOMER REPORTED PERFORMANCE DATA

Accent™ SR RF

MODEL PM1210

US Regulatory Approval	July 2009
Registered US Implants	39,812
Estimated Active US Implants	21,126
Estimated Longevity	10.9 Years
Normal Battery Depletion	25
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	2	<0.01%	8	0.02%
Electrical Interconnect	1	<0.01%	3	<0.01%
Battery	0	0.00%	1	<0.01%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	4	0.01%
Possible Early Battery Depletion	2	<0.01%	2	<0.01%
Other	0	0.00%	7	0.02%
Total	5	0.01%	26	0.07%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 100 MONTHS
SURVIVAL PROBABILITY	99.89%	99.80%	99.76%	99.74%	99.61%	99.54%	99.33%	99.33%	99.33%
± 1 STANDARD ERROR	0.02%	0.02%	0.03%	0.03%	0.04%	0.05%	0.07%	0.08%	0.08%
SAMPLE SIZE	36,670	31,520	27,780	23,190	16,920	10,740	5,880	2,250	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	AT 100 MONTHS
SURVIVAL PROBABILITY	99.93%	99.87%	99.83%	99.82%	99.78%	99.77%	99.73%	99.73%	99.73%
± 1 STANDARD ERROR	0.01%	0.02%	0.02%	0.02%	0.03%	0.03%	0.04%	0.04%	0.04%

Single-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

Accent™ SR RF

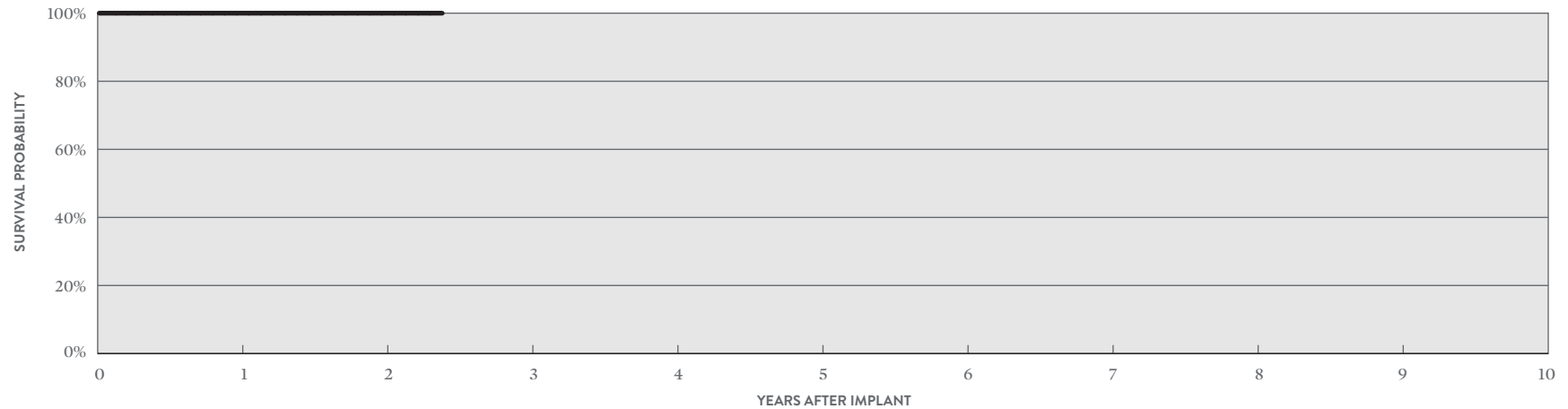
MODEL PM1210

US Regulatory Approval	July 2009
Number of Devices Enrolled in Study	236
Active Devices Enrolled in Study	26
Cumulative Months of Follow-up	5,678
Estimated Longevity	10.9 Years

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	AT 29 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%
SAMPLE SIZE	200	120	50

Single-Chamber Pacemakers

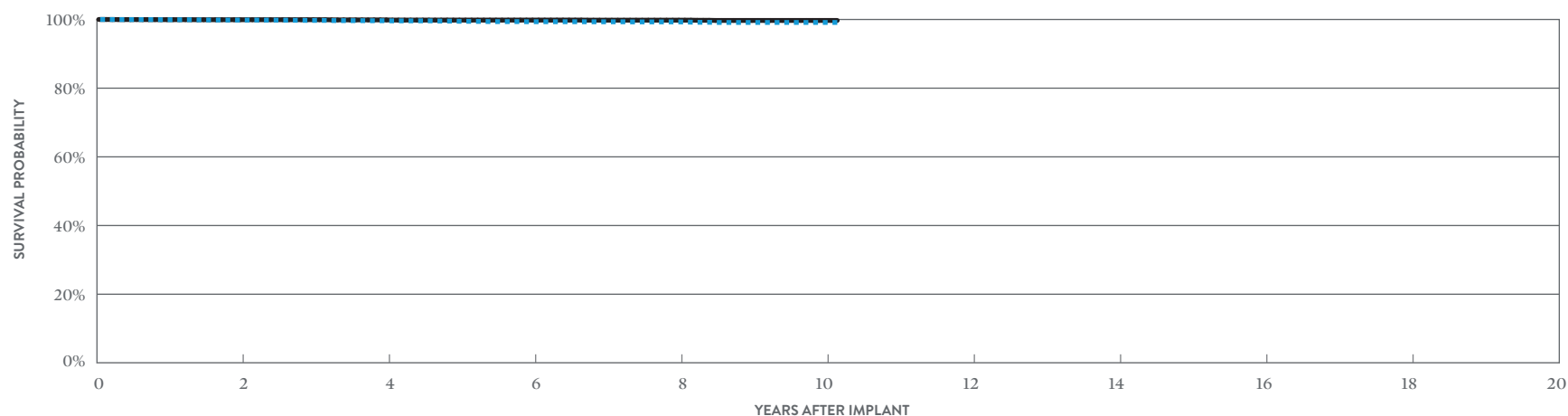
CUSTOMER REPORTED PERFORMANCE DATA

Zephyr™ XL SR

MODEL 5626

US Regulatory Approval	May 2007
Registered US Implants	20,647
Estimated Active US Implants	7,699
Estimated Longevity	15.8 Years
Normal Battery Depletion	29
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	4	0.02%
Electrical Interconnect	1	<0.01%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	1	<0.01%	8	0.04%
Total	2	<0.01%	12	0.06%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 122 MONTHS
SURVIVAL PROBABILITY	99.83%	99.64%	99.35%	99.30%	99.13%	99.13%
± 1 STANDARD ERROR	0.03%	0.05%	0.07%	0.08%	0.10%	0.10%
SAMPLE SIZE	15,780	11,860	8,920	6,240	1,370	280

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 122 MONTHS
SURVIVAL PROBABILITY	99.93%	99.88%	99.83%	99.80%	99.72%	99.72%
± 1 STANDARD ERROR	0.02%	0.03%	0.04%	0.04%	0.06%	0.06%

Single-Chamber Pacemakers

ACTIVELY MONITORED STUDY DATA

Zephyr™ XL SR

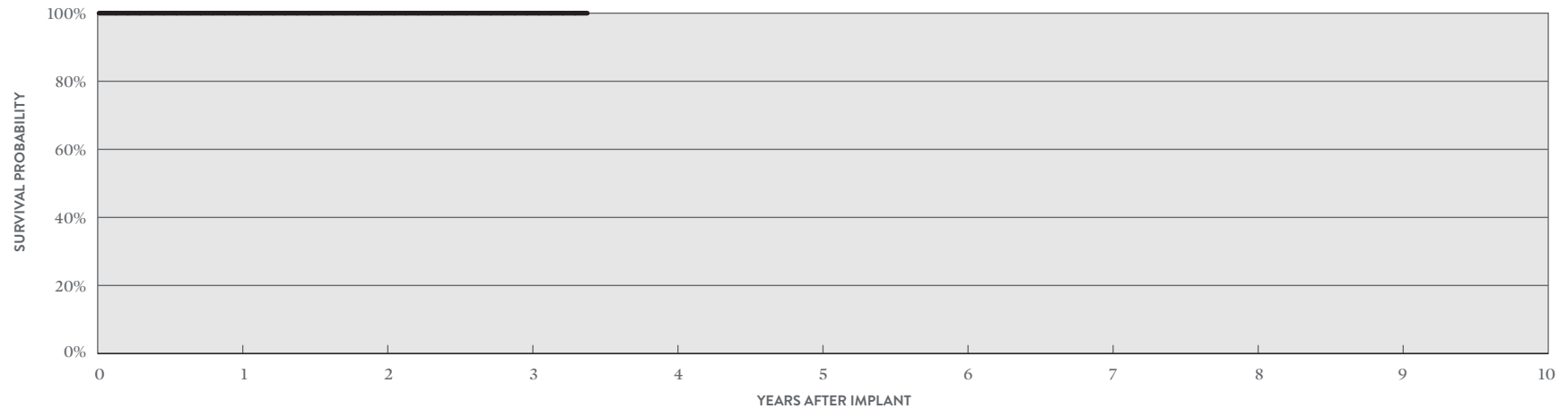
MODEL 5626

US Regulatory Approval	May 2007
Number of Devices Enrolled in Study	230
Active Devices Enrolled in Study	2
Cumulative Months of Follow-up	6,564
Estimated Longevity	15.8 Years

QUALIFYING COMPLICATIONS

None Reported

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	0	0.00%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	0	0.00%
Total	0	0.00%	0	0.00%



ACTIVELY MONITORED STUDY DATA

YEAR	1	2	3	AT 41 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	100.00%	100.00%
± 1 STANDARD ERROR	0.00%	0.00%	0.00%	0.00%
SAMPLE SIZE	220	180	120	50

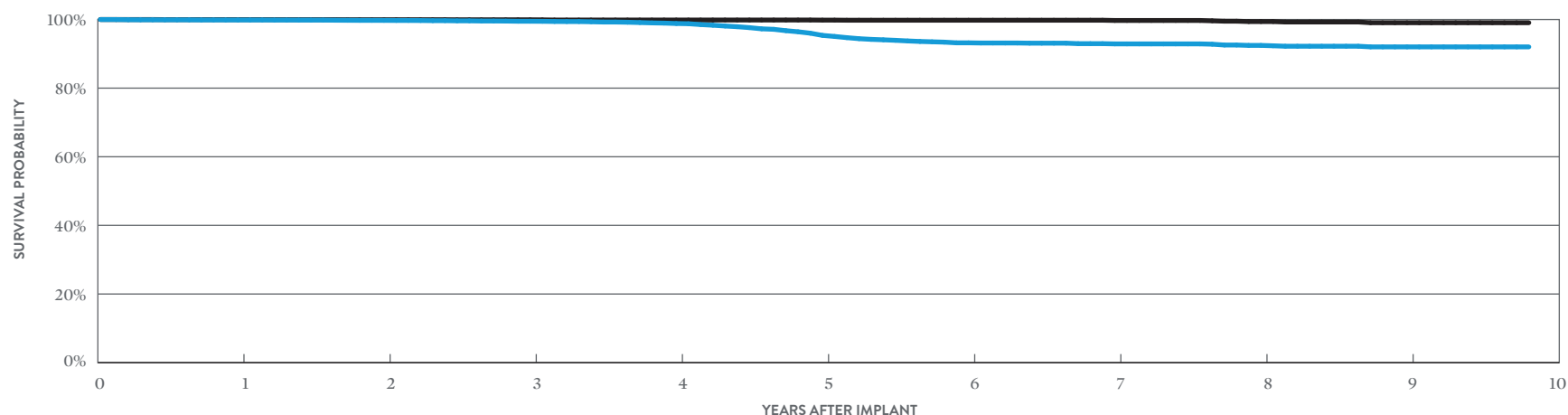
Single-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

Zephyr™ SR MODEL 5620

US Regulatory Approval	March 2007
Registered US Implants	17,349
Estimated Active US Implants	6,649
Estimated Longevity	8.8 Years
Normal Battery Depletion	196
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	4	0.02%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	2	0.01%
Mechanical	1	<0.01%	0	0.00%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	9	0.05%
Total	1	<0.01%	15	0.09%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 118 MONTHS
SURVIVAL PROBABILITY	99.86%	99.74%	99.49%	98.83%	95.37%	93.20%	92.89%	92.45%	92.02%	92.02%
± 1 STANDARD ERROR	0.03%	0.04%	0.06%	0.10%	0.22%	0.30%	0.31%	0.35%	0.39%	0.39%
SAMPLE SIZE	15,500	12,560	10,630	8,930	7,150	5,260	3,560	2,200	1,140	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 118 MONTHS
SURVIVAL PROBABILITY	99.97%	99.94%	99.92%	99.85%	99.82%	99.79%	99.72%	99.40%	99.06%	99.06%
± 1 STANDARD ERROR	0.01%	0.02%	0.02%	0.04%	0.04%	0.05%	0.05%	0.15%	0.23%	0.23%

Single-Chamber Pacemakers

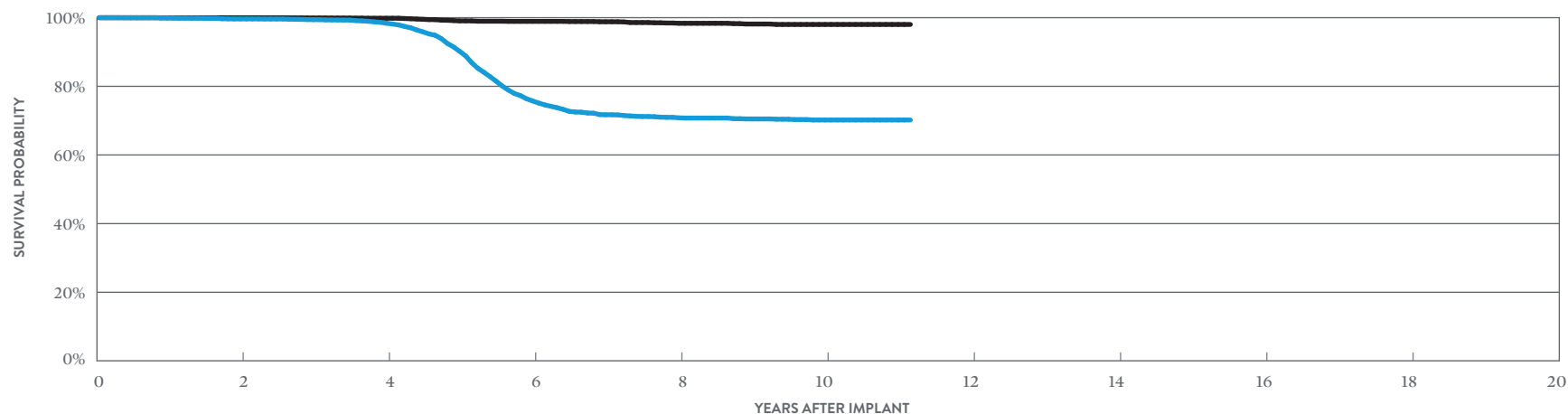
CUSTOMER REPORTED PERFORMANCE DATA

Victory™ SR

MODEL 5610

US Regulatory Approval	December 2005
Registered US Implants	13,688
Estimated Active US Implants	2,208
Estimated Longevity	8.8 Years
Normal Battery Depletion	669
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	23	0.17%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	1	<0.01%	14	0.10%
Total	1	<0.01%	39	0.28%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 134 MONTHS
SURVIVAL PROBABILITY	99.63%	98.38%	75.74%	70.82%	70.19%	70.19%
± 1 STANDARD ERROR	0.06%	0.13%	0.54%	0.60%	0.62%	0.62%
SAMPLE SIZE	10,140	7,280	4,870	2,870	1,420	260

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 134 MONTHS
SURVIVAL PROBABILITY	99.96%	99.83%	98.92%	98.36%	98.02%	98.02%
± 1 STANDARD ERROR	0.02%	0.05%	0.13%	0.18%	0.23%	0.23%

Single-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

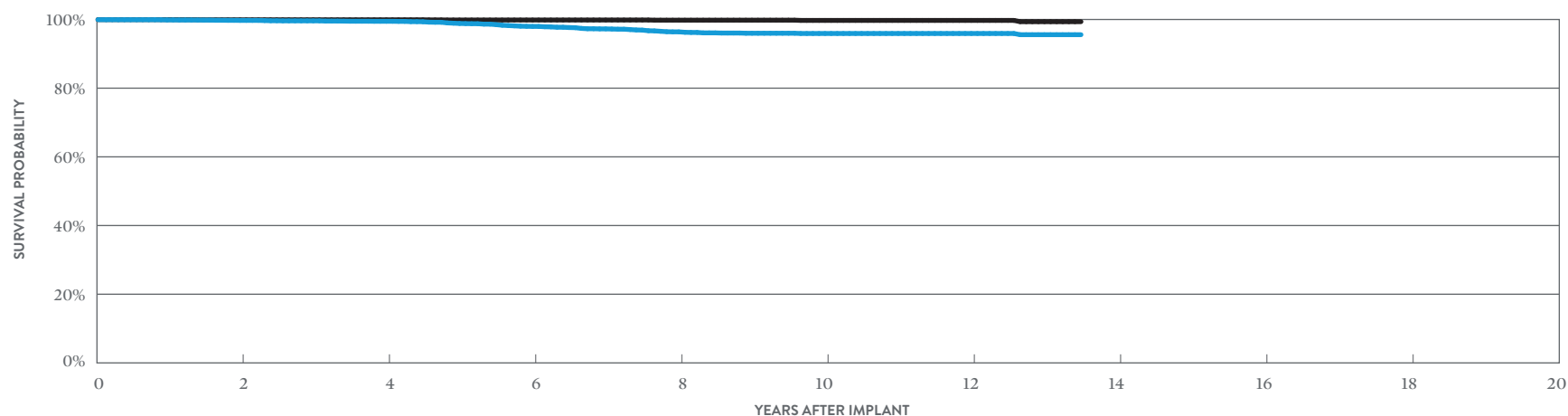
Verity ADx™ XL SR MODEL 5156

Verity ADx™ XL SR M/S MODEL 5157M/S

Verity ADx™ XL SC MODEL 5056

US Regulatory Approval	May 2003
Registered US Implants	14,503
Estimated Active US Implants	3,606
Estimated Longevity	10.2 Years
Normal Battery Depletion	91
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	<0.01%	4	0.03%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	0	0.00%	3	0.02%
Total	1	<0.01%	9	0.06%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 162 MONTHS
SURVIVAL PROBABILITY	99.73%	99.47%	97.98%	96.40%	95.92%	95.92%	95.57%
± 1 STANDARD ERROR	0.05%	0.07%	0.17%	0.25%	0.27%	0.27%	0.37%
SAMPLE SIZE	10,910	7,850	5,660	4,270	2,960	1,280	210

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 162 MONTHS
SURVIVAL PROBABILITY	99.91%	99.91%	99.85%	99.80%	99.73%	99.73%	99.36%
± 1 STANDARD ERROR	0.03%	0.03%	0.04%	0.05%	0.07%	0.07%	0.27%

Single-Chamber Pacemakers

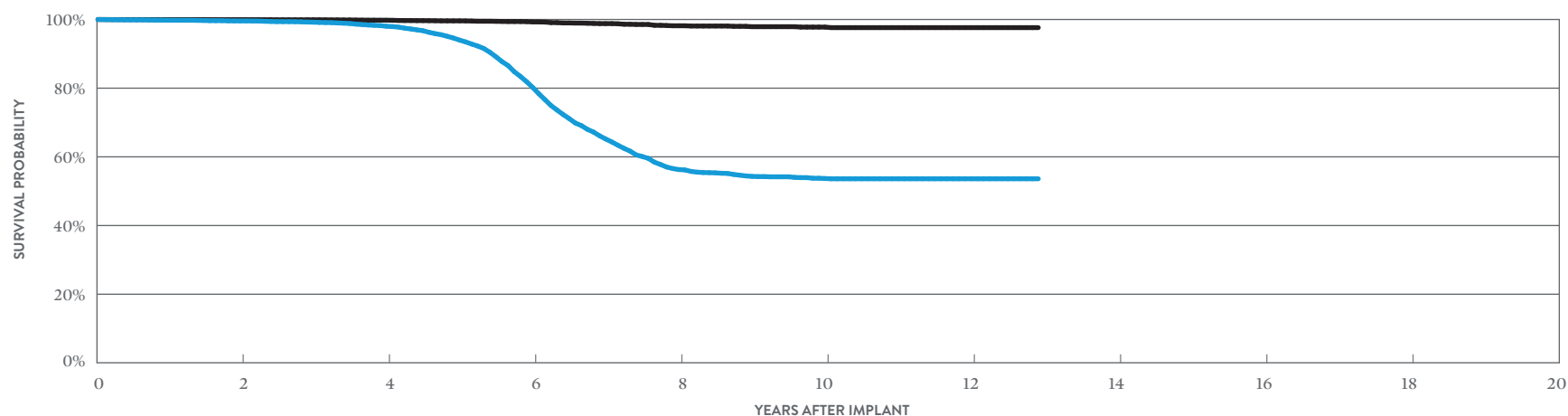
CUSTOMER REPORTED PERFORMANCE DATA

Identity ADx™ SR

MODEL 5180

US Regulatory Approval	May 2003
Registered US Implants	20,868
Estimated Active US Implants	2,104
Estimated Longevity	5.7 Years
Normal Battery Depletion	1,243
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	35	0.17%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	6	0.03%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	0	0.00%	8	0.04%
Other	0	0.00%	8	0.04%
Total	0	0.00%	58	0.28%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 155 MONTHS
SURVIVAL PROBABILITY	99.57%	98.03%	80.25%	56.29%	53.69%	53.61%	53.61%
± 1 STANDARD ERROR	0.05%	0.12%	0.43%	0.61%	0.64%	0.64%	0.64%
SAMPLE SIZE	15,440	10,930	6,880	3,300	1,660	700	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	AT 155 MONTHS
SURVIVAL PROBABILITY	99.94%	99.79%	99.28%	98.17%	97.78%	97.63%	97.63%
± 1 STANDARD ERROR	0.02%	0.04%	0.08%	0.19%	0.24%	0.26%	0.26%

Single-Chamber Pacemakers

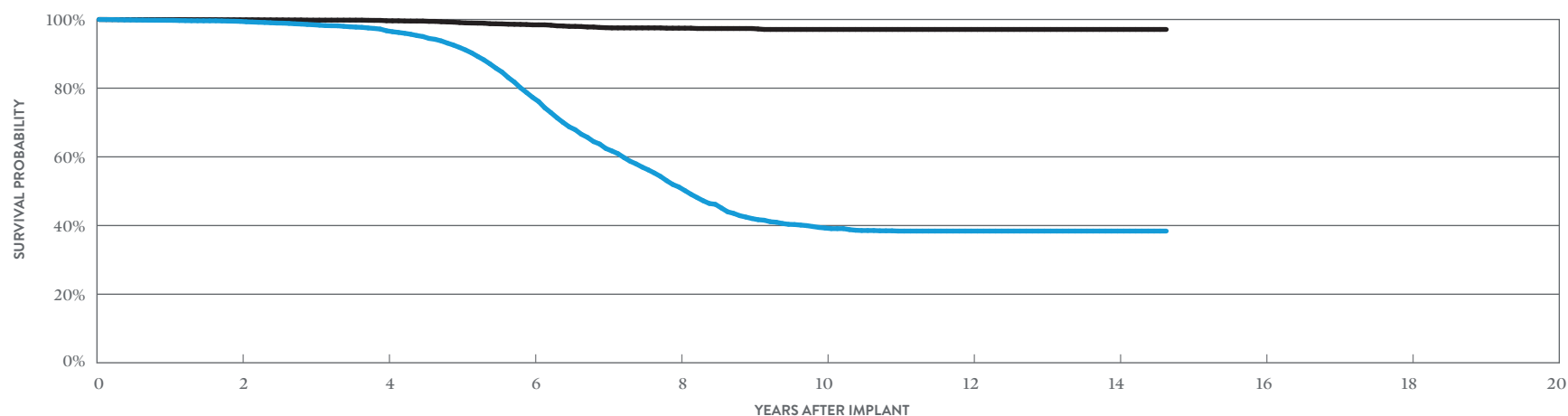
CUSTOMER REPORTED PERFORMANCE DATA

Identity™ SR

MODEL 5172

US Regulatory Approval	November 2001
Registered US Implants	21,884
Estimated Active US Implants	1,006
Estimated Longevity	7.8 Years
Normal Battery Depletion	1,473
Number of US Advisories (see pg. 334)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	1	<0.01%	64	0.29%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	1	<0.01%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	8	0.04%
Other	0	0.00%	6	0.03%
Total	1	<0.01%	79	0.36%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	AT 176 MONTHS
SURVIVAL PROBABILITY	99.45%	96.72%	77.32%	51.22%	39.27%	38.39%	38.39%	38.39%
± 1 STANDARD ERROR	0.05%	0.14%	0.45%	0.65%	0.70%	0.71%	0.71%	0.71%
SAMPLE SIZE	16,210	11,390	6,580	2,760	1,330	820	400	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	AT 176 MONTHS
SURVIVAL PROBABILITY	99.92%	99.63%	98.44%	97.47%	97.10%	97.10%	97.10%	97.10%
± 1 STANDARD ERROR	0.02%	0.04%	0.13%	0.21%	0.26%	0.26%	0.26%	0.26%

Single-Chamber Pacemakers

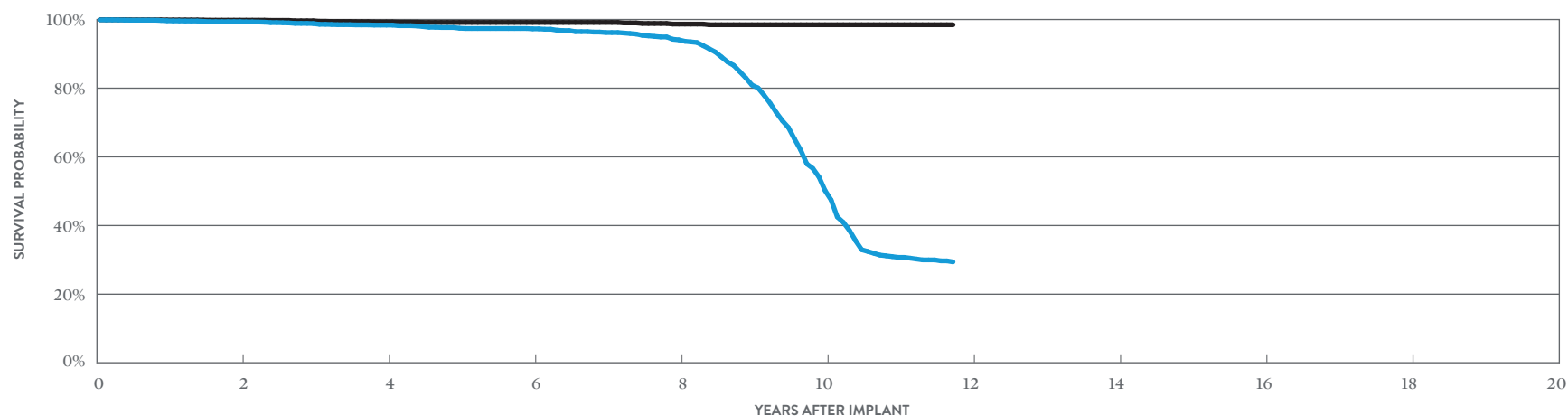
CUSTOMER REPORTED PERFORMANCE DATA

Microny™

MODELS 2425T, 2525T & 2535K

US Regulatory Approval	April 2001
Registered US Implants	7,781
Estimated Active US Implants	1,421
Estimated Longevity	7.5 Years
Normal Battery Depletion	308
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	1	0.01%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	0	0.00%
Possible Early Battery Depletion	0	0.00%	1	0.01%
Other	0	0.00%	0	0.00%
Total	0	0.00%	2	0.03%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 141 MONTHS
SURVIVAL PROBABILITY	99.36%	98.37%	97.28%	94.08%	50.17%	29.41%
± 1 STANDARD ERROR	0.11%	0.19%	0.27%	0.53%	1.61%	1.50%
SAMPLE SIZE	5,050	3,320	2,090	1,260	620	200

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	AT 141 MONTHS
SURVIVAL PROBABILITY	99.79%	99.29%	99.15%	98.67%	98.47%	98.47%
± 1 STANDARD ERROR	0.06%	0.13%	0.15%	0.24%	0.28%	0.28%

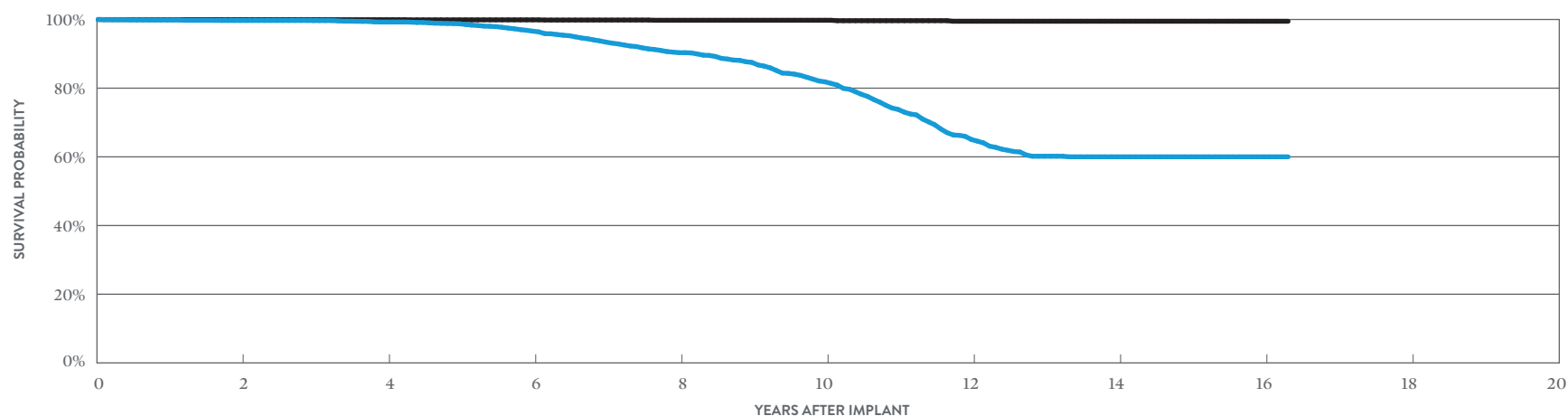
Single-Chamber Pacemakers

CUSTOMER REPORTED PERFORMANCE DATA

Integrity™ SR MODEL 5142

US Regulatory Approval	April 2000
Registered US Implants	10,491
Estimated Active US Implants	587
Estimated Longevity	8.6 Years
Normal Battery Depletion	386
Number of US Advisories	None

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	5	0.05%
Electrical Interconnect	0	0.00%	0	0.00%
Battery	0	0.00%	0	0.00%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	0	0.00%	1	<0.01%
Other	1	<0.01%	0	0.00%
Total	1	<0.01%	7	0.07%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 196 MONTHS
SURVIVAL PROBABILITY	99.71%	99.26%	96.60%	90.33%	81.84%	65.06%	60.01%	60.01%	60.01%
± 1 STANDARD ERROR	0.06%	0.10%	0.25%	0.48%	0.71%	1.01%	1.10%	1.10%	1.10%
SAMPLE SIZE	8,050	5,870	4,210	2,910	1,950	1,240	660	320	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 196 MONTHS
SURVIVAL PROBABILITY	99.93%	99.93%	99.89%	99.77%	99.77%	99.48%	99.48%	99.48%	99.48%
± 1 STANDARD ERROR	0.03%	0.03%	0.04%	0.07%	0.07%	0.16%	0.16%	0.16%	0.16%

Single-Chamber Pacemakers

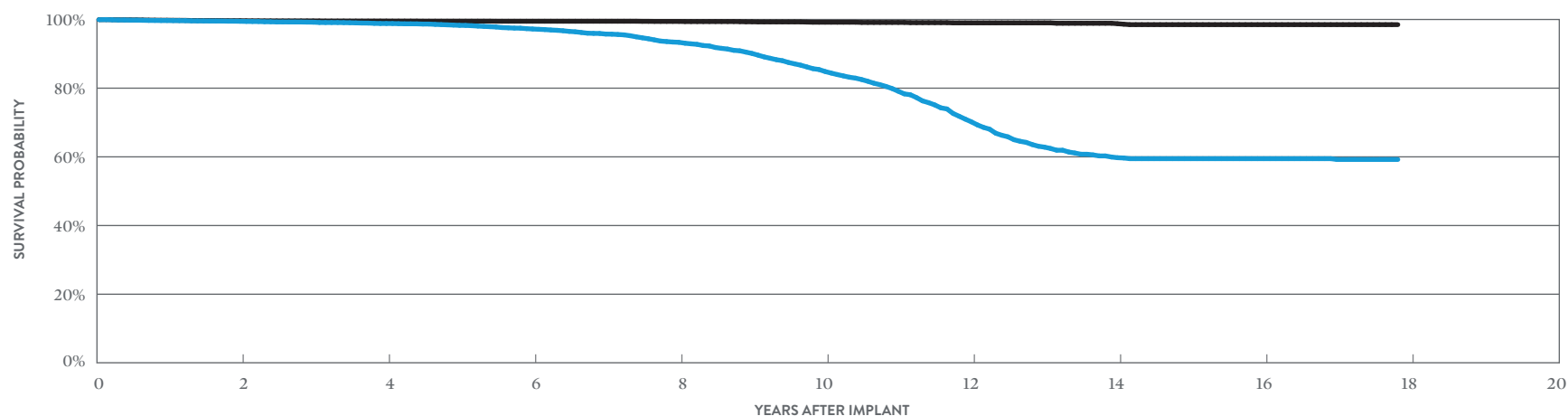
CUSTOMER REPORTED PERFORMANCE DATA

Affinity™ SR

MODELS 5130 & 5131

US Regulatory Approval	(5130) January 1999 (5131) June 1999
Registered US Implants	28,800
Estimated Active US Implants	1,268
Estimated Longevity	8.6 Years
Normal Battery Depletion	793
Number of US Advisories (see pg. 340)	One

	MALFUNCTIONS W/ COMPROMISED THERAPY		MALFUNCTIONS W/O COMPROMISED THERAPY	
	QTY	RATE	QTY	RATE
Electrical Component	0	0.00%	46	0.16%
Electrical Interconnect	3	0.01%	2	<0.01%
Battery	0	0.00%	3	0.01%
Software/Firmware	0	0.00%	0	0.00%
Mechanical	0	0.00%	1	<0.01%
Possible Early Battery Depletion	0	0.00%	0	0.00%
Other	1	<0.01%	7	0.02%
Total	4	0.01%	59	0.20%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 214 MONTHS
SURVIVAL PROBABILITY	99.47%	98.83%	97.23%	93.35%	84.85%	70.20%	59.75%	59.47%	59.23%
± 1 STANDARD ERROR	0.05%	0.08%	0.14%	0.25%	0.43%	0.65%	0.77%	0.78%	0.79%
SAMPLE SIZE	21,440	15,220	10,660	7,160	4,570	2,870	1,700	970	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	2	4	6	8	10	12	14	16	AT 214 MONTHS
SURVIVAL PROBABILITY	99.64%	99.54%	99.49%	99.44%	99.20%	99.01%	98.77%	98.50%	98.50%
± 1 STANDARD ERROR	0.04%	0.05%	0.05%	0.06%	0.09%	0.12%	0.14%	0.22%	0.22%

SUMMARY INFORMATION
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Single-Chamber Pacemakers

Survival Probability Summary

INCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
PM1272	AssurityMRI™*										
PM1160	Endurity™ SR	99.83%	99.83%	99.83%							
PM1240	Assurity™ SR	99.98%	99.96%	99.96%							
PM1110	Accent™ SR	99.92%	99.87%	99.85%	99.79%	99.76%	99.64%	99.45%			
PM1210	Accent™ SR RF	99.89%	99.80%	99.76%	99.74%	99.61%	99.54%	99.33%	99.33%		
5626	Zephyr™ XL SR	99.92%	99.83%	99.73%	99.64%	99.47%	99.35%	99.33%	99.30%	99.13%	99.13%
5620	Zephyr™ SR	99.86%	99.74%	99.49%	98.83%	95.37%	93.20%	92.89%	92.45%	92.02%	
5610	Victory™ SR	99.92%	99.63%	99.42%	98.38%	90.22%	75.74%	71.71%	70.82%	70.50%	70.19%
5156/5157/5056	Verity ADx™ XL SR/SR(M/S)/SC	99.87%	99.73%	99.60%	99.47%	98.84%	97.98%	97.26%	96.40%	95.99%	95.92%
5180	Identity ADx™ SR	99.79%	99.57%	99.21%	98.03%	94.00%	80.25%	65.19%	56.29%	54.31%	53.69%
5172	Identity™ SR	99.76%	99.45%	98.46%	96.72%	91.86%	77.32%	62.45%	51.22%	42.04%	39.27%
2425T/2525T/2535T	Microny™	99.63%	99.36%	98.82%	98.37%	97.46%	97.28%	96.20%	94.08%	80.90%	50.17%
5142	Integrity™ SR	99.86%	99.71%	99.68%	99.26%	98.75%	96.60%	93.42%	90.33%	87.49%	81.84%
5130/5131	Affinity™ SR	99.69%	99.47%	99.21%	98.83%	98.29%	97.23%	95.75%	93.35%	90.11%	84.85%

*No survival probability is stated at one year due to the device not meeting the required minimum sample size of 200 U.S. implants with 12 consecutive months of data. Please refer to the individual graphs for data up to one year.

Single-Chamber Pacemakers Survival Probability Summary

EXCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
PM1272	AssurityMRI™*										
PM1160	Endurity™ SR	99.83%	99.83%	99.83%							
PM1240	Assurity™ SR	99.98%	99.96%	99.96%							
PM1110	Accent™ SR	99.96%	99.94%	99.92%	99.92%	99.92%	99.92%	99.92%			
PM1210	Accent™ SR RF	99.93%	99.87%	99.83%	99.82%	99.78%	99.77%	99.73%	99.73%		
5626	Zephyr™ XL SR	99.95%	99.93%	99.93%	99.88%	99.83%	99.83%	99.80%	99.80%	99.72%	99.72%
5620	Zephyr™ SR	99.97%	99.94%	99.92%	99.85%	99.82%	99.79%	99.72%	99.40%	99.06%	
5610	Victory™ SR	99.98%	99.96%	99.91%	99.83%	99.06%	98.92%	98.81%	98.36%	98.15%	98.02%
5156/5157/5056	Verity ADx™ XL SR/SR(M/S)/SC	99.97%	99.91%	99.91%	99.91%	99.85%	99.85%	99.85%	99.80%	99.80%	99.73%
5180	Identity ADx™ SR	99.96%	99.94%	99.91%	99.79%	99.60%	99.28%	98.80%	98.17%	97.90%	97.78%
5172	Identity™ SR	99.97%	99.92%	99.81%	99.63%	99.10%	98.44%	97.61%	97.47%	97.37%	97.10%
2425T/2525T/2535T	Microny™	99.87%	99.79%	99.61%	99.29%	99.15%	99.15%	99.15%	98.67%	98.47%	98.47%
5142	Integrity™ SR	99.98%	99.93%	99.93%	99.93%	99.89%	99.89%	99.84%	99.77%	99.77%	99.77%
5130/5131	Affinity™ SR	99.78%	99.64%	99.58%	99.54%	99.51%	99.49%	99.49%	99.44%	99.33%	99.20%

*No survival probability is stated at one year due to the device not meeting the required minimum sample size of 200 U.S. implants with 12 consecutive months of data. Please refer to the individual graphs for data up to one year.

Single-Chamber Pacemakers

US Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM1272	Assurity MRI™	4,415	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM1160	Endurity™ SR	2,455	0.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM1240	Assurity™ SR	25,978	0.10%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM1110	Accent™ SR	13,590	3.60%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM1210	Accent™ SR RF	39,812	3.60%	2	<0.01%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	2	<0.01%	0	0.00%	5	0.01%
5626	Zephyr™ XL SR	20,647	5.40%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	2	<0.01%
5620	Zephyr™ SR	17,349	5.70%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	1	<0.01%
5610	Victory™ SR	13,688	12.70%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%
5156/5157/5056	Verity ADx™ XL SR/SR(M/S)/SC	14,503	5.80%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
5180	Identity ADx™ SR	20,868	11.70%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5172	Identity™ SR	21,884	11.30%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
2425T/2525T/2535T	Microny™	7,781	6.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5142	Integrity™ SR	10,491	8.60%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%
5130/5131	Affinity™ SR	28,800	7.00%	0	0.00%	3	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	4	0.01%

Definitions of malfunction categories can be found on [pages 5-6](#).

Single-Chamber Pacemakers

US Malfunction Summary

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM1272	Assurity MRI™	4,415	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM1160	Endurity™ SR	2,455	0.50%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	1	0.04%	2	0.08%
PM1240	Assurity™ SR	25,978	0.10%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	2	<0.01%	0	0.00%	0	0.00%	4	0.02%
PM1110	Accent™ SR	13,590	3.60%	2	0.01%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	0	0.00%	4	0.03%
PM1210	Accent™ SR RF	39,812	3.60%	8	0.02%	3	<0.01%	1	<0.01%	1	<0.01%	4	0.01%	2	<0.01%	7	0.02%	26	0.07%
5626	Zephyr™ XL SR	20,647	5.40%	4	0.02%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	8	0.04%	12	0.06%
5620	Zephyr™ SR	17,349	5.70%	4	0.02%	0	0.00%	0	0.00%	2	0.01%	0	0.00%	0	0.00%	9	0.05%	15	0.09%
5610	Victory™ SR	13,688	12.70%	23	0.17%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	1	<0.01%	14	0.10%	39	0.28%
5156/5157/5056	Verity ADx™ XL SR/SR(M/S)/SC	14,503	5.80%	4	0.03%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%	0	0.00%	3	0.02%	9	0.06%
5180	Identity ADx™ SR	20,868	11.70%	35	0.17%	0	0.00%	0	0.00%	6	0.03%	1	<0.01%	8	0.04%	8	0.04%	58	0.28%
5172	Identity™ SR	21,884	11.30%	64	0.29%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	8	0.04%	6	0.03%	79	0.36%
2425T/2525T/2535T	Microny™	7,781	6.50%	1	0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.01%	0	0.00%	2	0.03%
5142	Integrity™ SR	10,491	8.60%	5	0.05%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	1	<0.01%	0	0.00%	7	0.07%
5130/5131	Affinity™ SR	28,800	7.00%	46	0.16%	2	<0.01%	3	0.01%	0	0.00%	1	<0.01%	0	0.00%	7	0.02%	59	0.20%

Definitions of malfunction categories can be found on [pages 5-6](#).

Single-Chamber Pacemakers

Worldwide Malfunction Summary

WITH COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM1272	Assurity MRI™	33,282	0.27%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
PM1160	Endurity™ SR	25,409	0.52%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%
PM1240	Assurity™ SR	29,452	2.48%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM1110	Accent™ SR	54,050	1.84%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
PM1210	Accent™ SR RF	48,071	4.92%	4	<0.01%	1	<0.01%	0	0.00%	0	0.00%	0	0.00%	2	<0.01%	0	0.00%	7	0.01%

WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM1272	Assurity MRI™	33,282	0.27%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	<0.01%	0	0.00%	0	0.00%	1	<0.01%
PM1160	Endurity™ SR	25,409	0.52%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	3	0.01%	0	0.00%	1	<0.01%	4	0.02%
PM1240	Assurity™ SR	29,452	2.48%	2	<0.01%	0	0.00%	0	0.00%	0	0.00%	3	0.01%	0	0.00%	0	0.00%	5	0.02%
PM1110	Accent™ SR	54,050	1.84%	3	<0.01%	0	0.00%	0	0.00%	2	<0.01%	0	0.00%	1	<0.01%	2	<0.01%	8	0.01%
PM1210	Accent™ SR RF	48,071	4.92%	11	0.02%	3	<0.01%	1	<0.01%	1	<0.01%	4	<0.01%	2	<0.01%	8	0.02%	30	0.06%

Definitions of malfunction categories can be found on [pages 5-6](#).

Single-Chamber Pacemakers

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	LOSS OF TELEMETRY		PERICARDIAL EFFUSION		PREMATURE BATTERY DEPLETION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM1210	236	23	5,678	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5626	230	2	6,564	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

MALFUNCTIONS WITH COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/ FIRM-WARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM1210	Accent™ VR	236	3.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5626	Zephyr™ XL SR	230	3.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

MALFUNCTIONS WITHOUT COMPROMISED THERAPY

MODELS	FAMILY	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/ FIRM-WARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
PM1210	Accent™ VR	236	3.40%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
5626	Zephyr™ XL SR	230	3.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%

A list of complications can be found on [page 12](#).

Definitions of malfunction categories can be found on [pages 5-6](#).

Pacing Leads

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

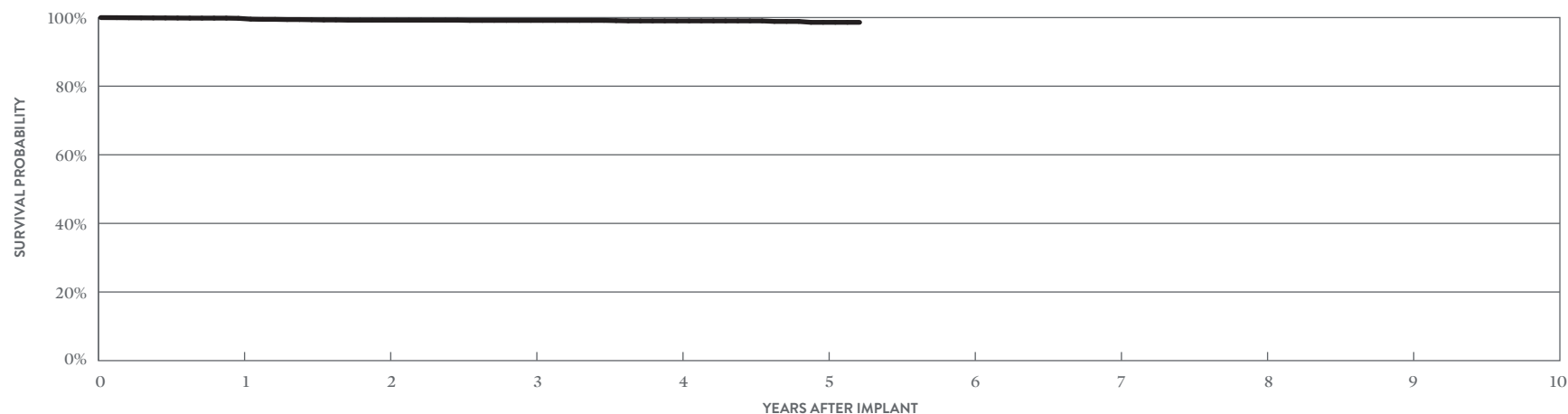
Tendril MRI™

MODEL LPA1200M

US Regulatory Approval	January 2017
Registered US Implants	57,116
Estimated Active US Implants	48,436
Insulation	Optim™*
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	12	0.02%	3	<0.01%
Conductor Fracture	0	0.00%	0	0.00%
Lead Dislodgement	95	0.17%	49	0.09%
Failure to Capture	6	0.01%	24	0.04%
Oversensing	1	<0.01%	22	0.04%
Failure to Sense	6	0.01%	7	0.01%
Insulation Breach	0	0.00%	0	0.00%
Abnormal Pacing Impedance	0	0.00%	2	<0.01%
Extracardiac Stimulation	2	<0.01%	2	<0.01%
Other	9	0.02%	5	<0.01%
Total	131	0.23%	114	0.20%
Total Returned for Analysis	29		25	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	3	<0.01%
Insulation Breach	5	<0.01%
Crimps, Welds & Bonds	0	0.00%
Other	2	<0.01%
Extrinsic Factors	29	0.05%
Total	39	0.07%



YEAR	1	2	3	4	5	AT 63 MONTHS
SURVIVAL PROBABILITY	99.78%	99.24%	99.17%	99.03%	98.61%	98.61%
± 1 STANDARD ERROR	0.03%	0.19%	0.20%	0.23%	0.31%	0.31%
SAMPLE SIZE	29,400	1,620	1,500	1,410	920	210

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

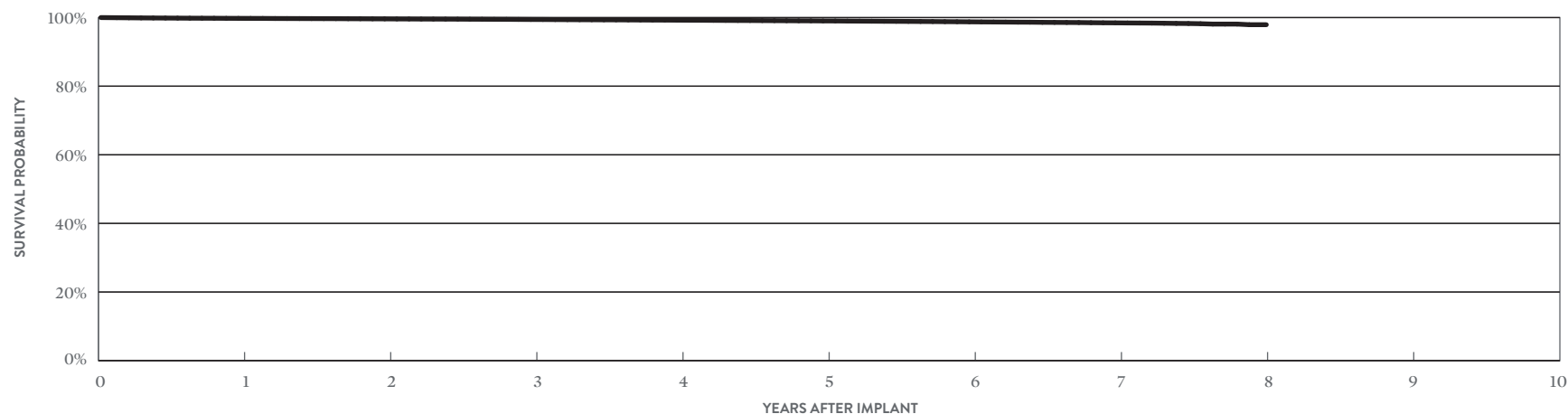
Tendril™ STS

MODEL 2088TC

US Regulatory Approval	May 2009
Registered US Implants	585,050
Estimated Active US Implants	382,081
Insulation	Optim™*
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	83	0.01%	52	<0.01%
Conductor Fracture	6	<0.01%	192	0.03%
Lead Dislodgement	582	0.10%	826	0.14%
Failure to Capture	148	0.03%	635	0.11%
Oversensing	42	<0.01%	1726	0.30%
Failure to Sense	20	<0.01%	95	0.02%
Insulation Breach	12	<0.01%	169	0.03%
Abnormal Pacing Impedance	31	<0.01%	128	0.02%
Extracardiac Stimulation	4	<0.01%	29	<0.01%
Other	99	0.02%	129	0.02%
Total	1027	0.18%	3981	0.68%
Total Returned for Analysis	460		1387	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	43	<0.01%
Insulation Breach	570	0.10%
Crimps, Welds & Bonds	0	0.00%
Other	27	<0.01%
Extrinsic Factors	995	0.17%
Total	1635	0.28%



YEAR	1	2	3	4	5	6	7	8
SURVIVAL PROBABILITY	99.77%	99.62%	99.44%	99.24%	99.02%	98.76%	98.46%	97.92%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.02%	0.02%	0.03%	0.04%	0.20%
SAMPLE SIZE	517,680	392,900	290,530	208,350	139,830	83,290	38,580	280

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

ACTIVELY MONITORED STUDY DATA

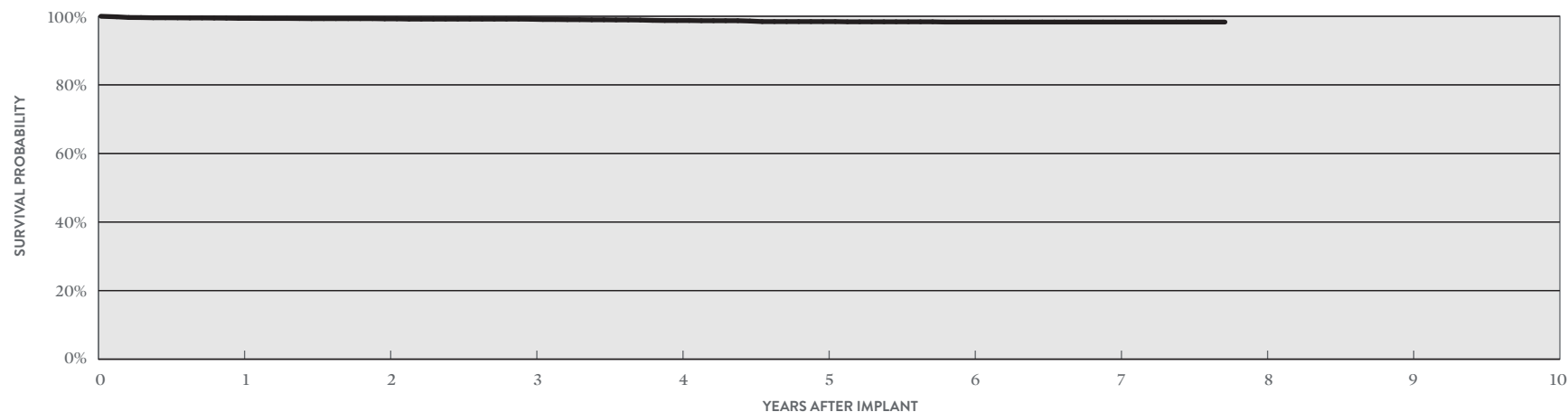
Tendril™ STS

MODEL 2088TC

US Regulatory Approval	May 2009
Number of Devices Enrolled in Study	3,841
Active Devices Enrolled in Study	1,795
Cumulative Months of Follow-up	194,241
Insulation	Optim™*
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Pacing Impedance	1	0.03%
Cardiac Perforation	1	0.03%
Conductor Fracture	2	0.05%
Extracardiac Stimulation	1	0.03%
Failure to Capture	6	0.16%
Failure to Sense	2	0.05%
Insulation Breach	6	0.16%
Lead Dislodgement	15	0.39%
Oversensing	12	0.31%
Pericardial Effusion	1	0.03%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	1	0.03%
Insulation Breach	12	0.31%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	14	0.36%
Total	27	0.70%



YEAR	1	2	3	4	5	6	7	AT 93 MONTHS
SURVIVAL PROBABILITY	99.40%	99.25%	99.14%	98.76%	98.45%	98.32%	98.32%	98.32%
± 1 STANDARD ERROR	0.12%	0.14%	0.15%	0.20%	0.24%	0.26%	0.26%	0.26%
SAMPLE SIZE	3,640	3,220	2,770	2,350	1,900	1,480	880	60

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

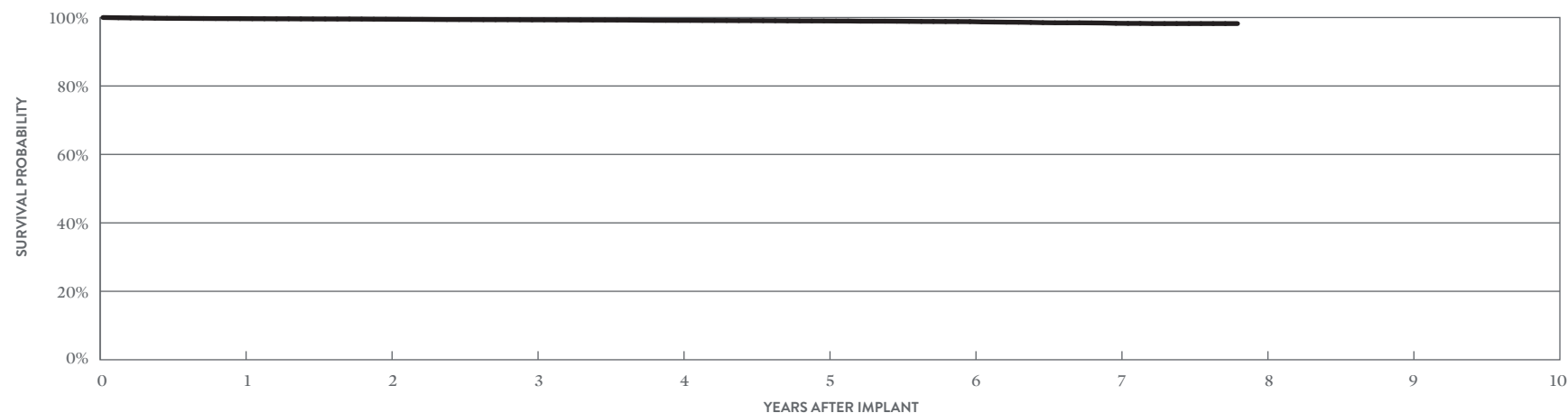
OptiSense™

MODEL 1999

US Regulatory Approval	May 2007
Registered US Implants	46,528
Estimated Active US Implants	28,723
Insulation	Optim™*
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	4	<0.01%	0	0.00%
Conductor Fracture	0	0.00%	7	0.02%
Lead Dislodgement	64	0.14%	154	0.33%
Failure to Capture	8	0.02%	51	0.11%
Oversensing	6	0.01%	149	0.32%
Failure to Sense	3	<0.01%	21	0.05%
Insulation Breach	1	<0.01%	25	0.05%
Abnormal Pacing Impedance	0	0.00%	11	0.02%
Extracardiac Stimulation	0	0.00%	1	<0.01%
Other	12	0.03%	16	0.03%
Total	98	0.21%	435	0.93%
Total Returned for Analysis	52		171	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	5	0.01%
Insulation Breach	43	0.09%
Crimps, Welds & Bonds	0	0.00%
Other	6	0.01%
Extrinsic Factors	144	0.31%
Total	198	0.43%



YEAR	1	2	3	4	5	6	7	AT 94 MONTHS
SURVIVAL PROBABILITY	99.67%	99.51%	99.32%	99.16%	99.00%	98.80%	98.31%	98.23%
± 1 STANDARD ERROR	0.03%	0.03%	0.04%	0.05%	0.06%	0.07%	0.11%	0.14%
SAMPLE SIZE	42,890	35,700	28,480	21,510	15,360	10,010	5,270	330

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

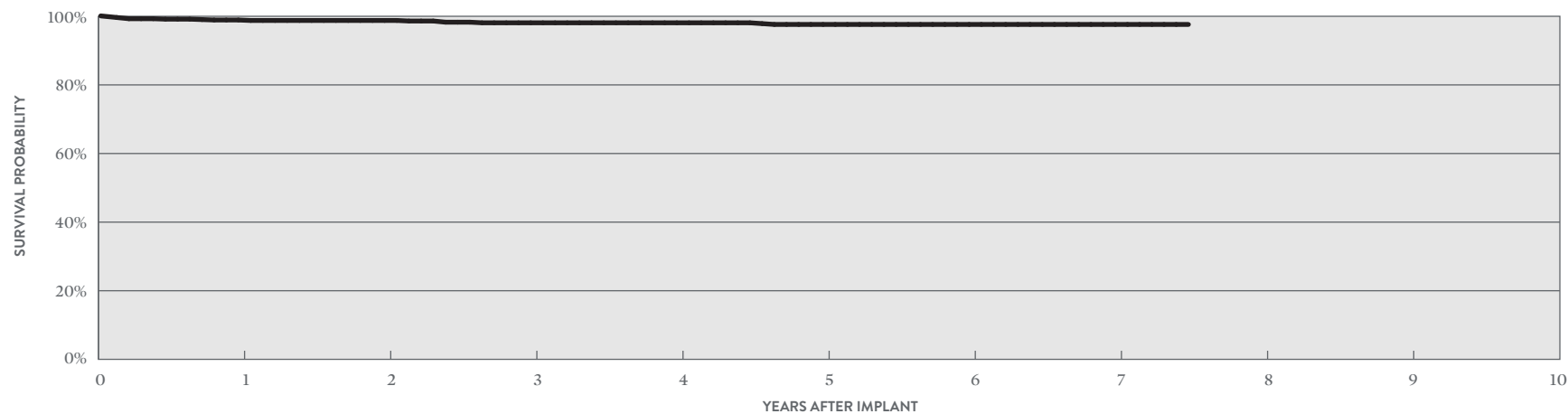
Pacing Leads

ACTIVELY MONITORED STUDY DATA

OptiSense™

MODEL 1999

		QUALIFYING COMPLICATIONS			MALFUNCTIONS		
			QTY	RATE		QTY	RATE
US Regulatory Approval	May 2007	Abnormal Pacing Impedance	1	0.12%	Conductor Fracture	0	0.00%
Number of Devices Enrolled in Study	869	Conductor Fracture	1	0.12%	Insulation Breach	5	0.58%
Active Devices Enrolled in Study	404	Failure to Sense	2	0.23%	Crimps, Welds & Bonds	0	0.00%
Cumulative Months of Follow-up	41,920	Insulation Breach	1	0.12%	Other	0	0.00%
Insulation	Optim™*	Lead Dislodgement	10	1.15%	Extrinsic Factors	8	0.92%
Type and/or Fixation	Active	Oversensing	1	0.12%	Total	13	1.50%
Polarity	Bipolar						
Steroid	Yes						



YEAR	1	2	3	4	5	6	7	AT 90 MONTHS
SURVIVAL PROBABILITY	98.92%	98.78%	98.10%	98.10%	97.62%	97.62%	97.62%	97.62%
± 1 STANDARD ERROR	0.36%	0.38%	0.51%	0.51%	0.61%	0.61%	0.61%	0.61%
SAMPLE SIZE	810	690	580	500	420	320	190	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

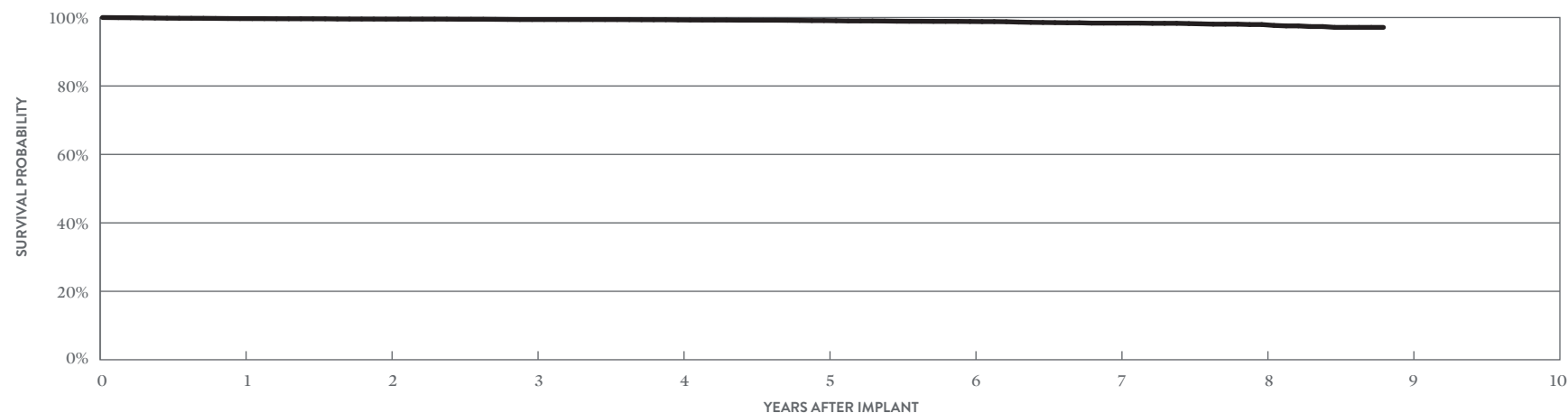
Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

IsoFlex™ Optim™

MODEL 1944

		ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)		MALFUNCTIONS		QTY	RATE
		QTY	RATE	QTY	RATE			QTY	RATE
US Regulatory Approval	March 2008	Cardiac Perforation	0	0.00%	1	<0.01%	Conductor Fracture	0	0.00%
Registered US Implants	16,578	Conductor Fracture	0	0.00%	6	0.04%	Insulation Breach	8	0.05%
Estimated Active US Implants	9,638	Lead Dislodgement	67	0.40%	46	0.28%	Crimps, Welds & Bonds	0	0.00%
Insulation	Optim™*	Failure to Capture	8	0.05%	29	0.17%	Other	1	<0.01%
Type and/or Fixation	Passive	Oversensing	0	0.00%	49	0.30%	Extrinsic Factors	23	0.14%
Polarity	Bipolar	Failure to Sense	2	0.01%	6	0.04%	Total	32	0.19%
Steroid	Yes	Insulation Breach	0	0.00%	5	0.03%			
Number of US Advisories	None	Abnormal Pacing Impedance	0	0.00%	1	<0.01%			
		Extracardiac Stimulation	2	0.01%	1	<0.01%			
		Other	1	<0.01%	2	0.01%			
		Total	80	0.48%	146	0.88%			
		Total Returned for Analysis	45		27				



YEAR	1	2	3	4	5	6	7	8	AT 106 MONTHS
SURVIVAL PROBABILITY	99.69%	99.54%	99.41%	99.30%	99.11%	98.84%	98.37%	97.95%	97.12%
± 1 STANDARD ERROR	0.04%	0.06%	0.07%	0.07%	0.10%	0.12%	0.19%	0.27%	0.46%
SAMPLE SIZE	14,870	12,030	9,790	7,650	5,650	3,920	2,470	1,300	230

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

ACTIVELY MONITORED STUDY DATA

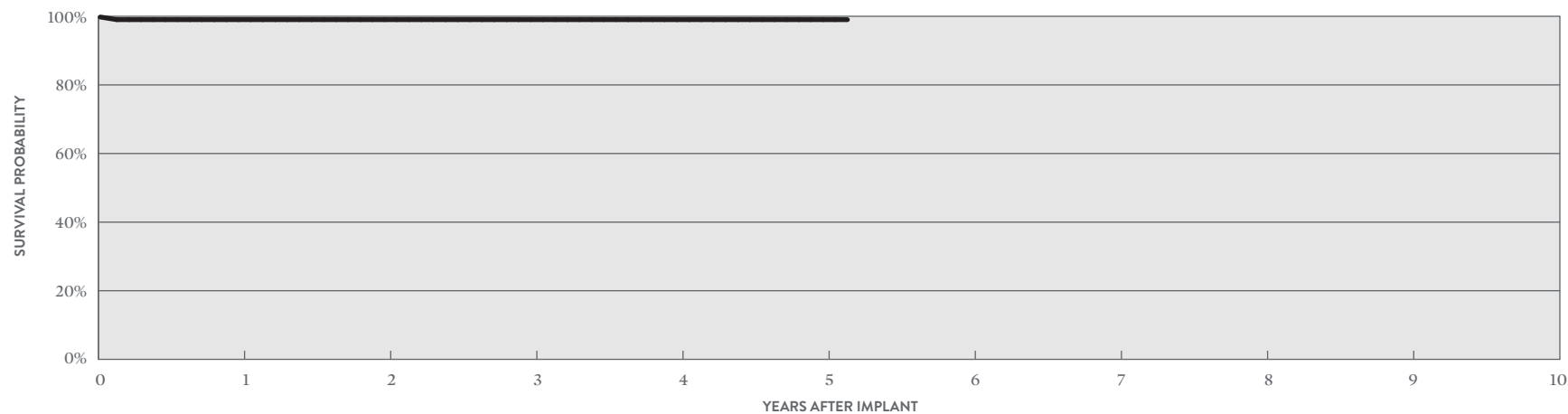
IsoFlex™ Optim™

MODEL 1944

US Regulatory Approval	March 2008
Number of Devices Enrolled in Study	104
Active Devices Enrolled in Study	31
Cumulative Months of Follow-up	5,742
Insulation	Optim™*
Type and/or Fixation	Passive
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Lead Dislodgement	1	0.96%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Insulation Breach	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	0	0.00%
Total	0	0.00%



YEAR	1	2	3	5	4	AT 62 MONTHS
SURVIVAL PROBABILITY	99.02%	99.02%	99.02%	99.02%	99.02%	99.02%
± 1 STANDARD ERROR	0.97%	0.97%	0.97%	0.97%	0.97%	0.97%
SAMPLE SIZE	100	80	70	60	60	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

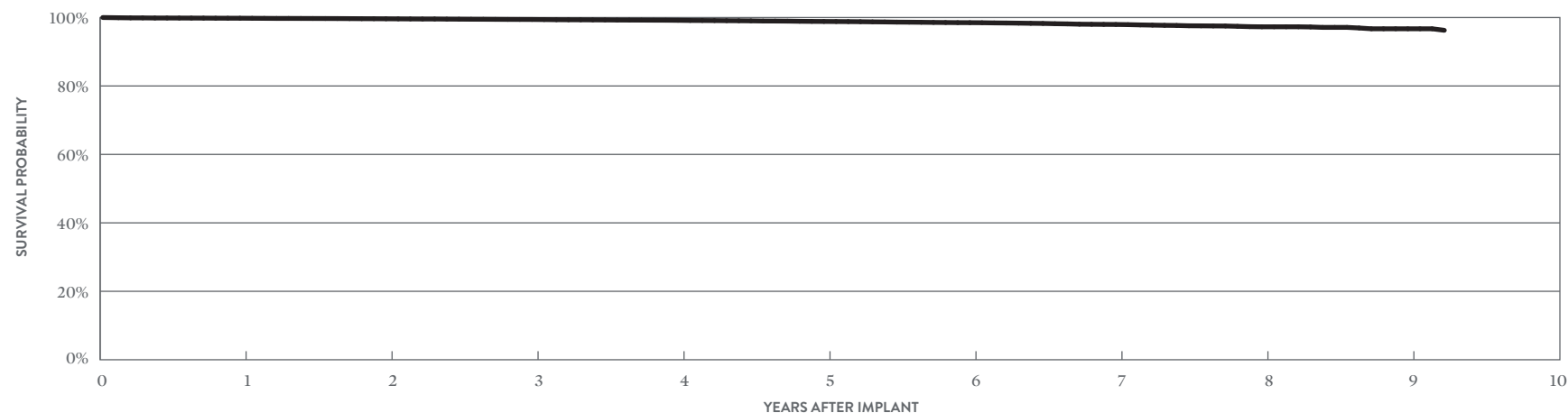
IsoFlex™ Optim™

MODEL 1948

US Regulatory Approval	March 2008
Registered US Implants	63,721
Estimated Active US Implants	37,190
Insulation	Optim™*
Type and/or Fixation	Passive
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	3	<0.01%	10	0.02%
Conductor Fracture	0	0.00%	69	0.11%
Lead Dislodgement	51	0.08%	63	0.10%
Failure to Capture	25	0.04%	135	0.21%
Oversensing	1	<0.01%	189	0.30%
Failure to Sense	2	<0.01%	2	<0.01%
Insulation Breach	4	<0.01%	43	0.07%
Abnormal Pacing Impedance	1	<0.01%	30	0.05%
Extracardiac Stimulation	2	<0.01%	5	<0.01%
Other	6	<0.01%	12	0.02%
Total	95	0.15%	558	0.88%
Total Returned for Analysis	51		110	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	10	0.02%
Insulation Breach	75	0.12%
Crimps, Welds & Bonds	0	0.00%
Other	1	<0.01%
Extrinsic Factors	74	0.12%
Total	160	0.25%



YEAR	1	2	3	4	5	6	7	8	9	AT 111 MONTHS
SURVIVAL PROBABILITY	99.80%	99.63%	99.43%	99.18%	98.85%	98.50%	97.97%	97.29%	96.71%	96.27%
± 1 STANDARD ERROR	0.02%	0.03%	0.04%	0.05%	0.06%	0.08%	0.11%	0.17%	0.28%	0.28%
SAMPLE SIZE	57,090	45,600	36,380	27,960	20,140	13,460	8,310	4,370	1,530	260

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

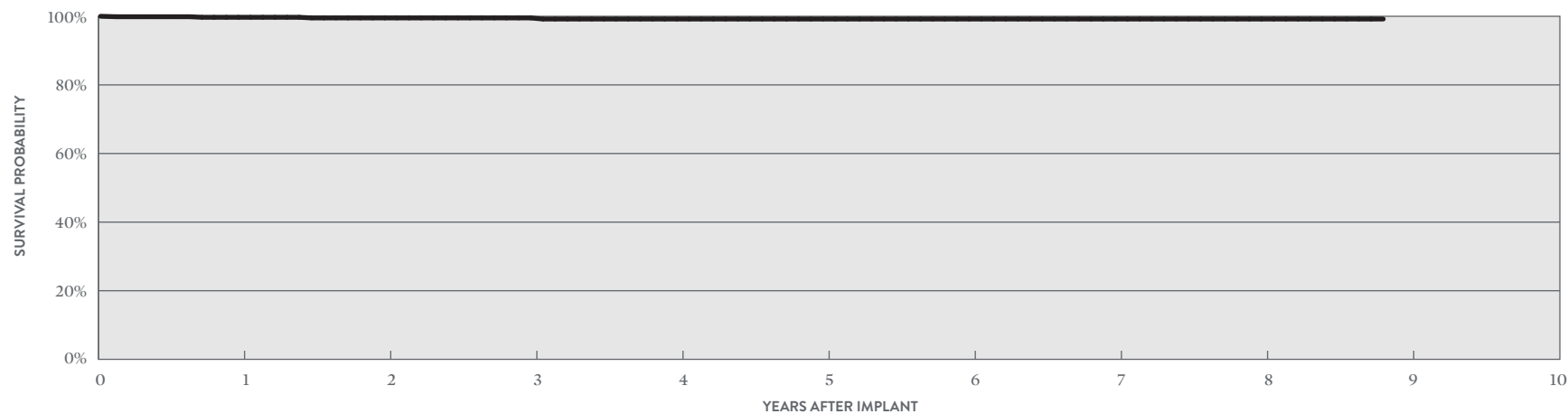
Pacing Leads

ACTIVELY MONITORED STUDY DATA

IsoFlex™ Optim™

MODEL 1948

		QUALIFYING COMPLICATIONS		MALFUNCTIONS			
		QTY	RATE	QTY	RATE		
US Regulatory Approval	March 2008	Failure to Capture	1	0.13%	Conductor Fracture	0	0.00%
Number of Devices Enrolled in Study	765	Insulation Breach	1	0.13%	Insulation Breach	4	0.52%
Active Devices Enrolled in Study	197	Lead Dislodgement	2	0.26%	Crimps, Welds & Bonds	0	0.00%
Cumulative Months of Follow-up	33,414				Other	0	0.00%
Insulation	Optim™*				Extrinsic Factors	1	0.13%
Type and/or Fixation	Passive				Total	5	0.65%
Polarity	Bipolar						
Steroid	Yes						



YEAR	1	2	3	4	5	6	7	8	AT 106 MONTHS
SURVIVAL PROBABILITY	99.71%	99.52%	99.52%	99.20%	99.20%	99.20%	99.20%	99.20%	99.20%
± 1 STANDARD ERROR	0.20%	0.28%	0.28%	0.43%	0.43%	0.43%	0.43%	0.43%	0.43%
SAMPLE SIZE	680	520	380	300	260	220	190	180	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

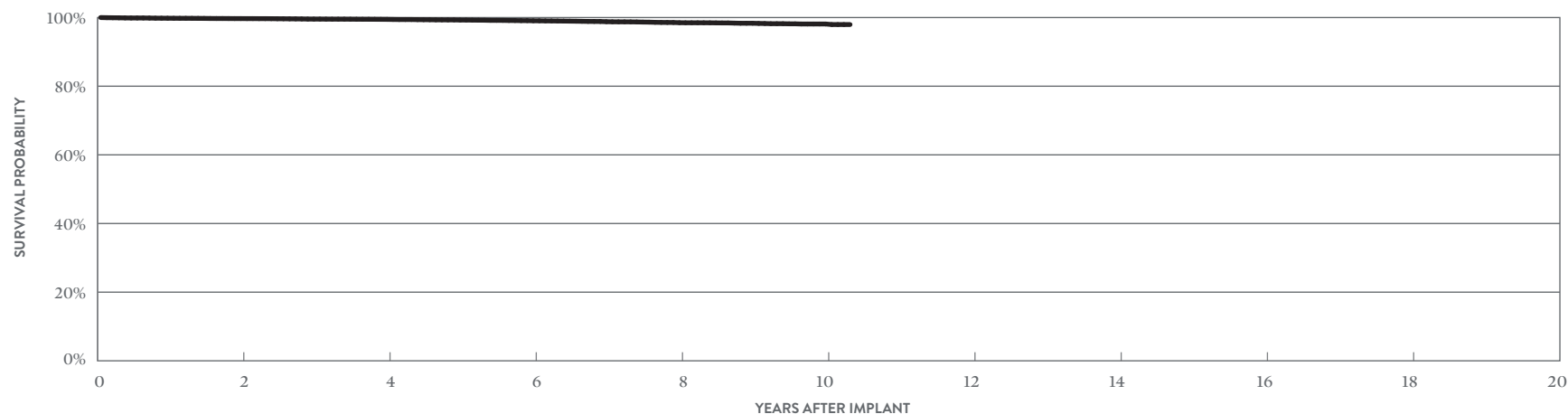
OptiSense™

MODELS 1699T & 1699TC

US Regulatory Approval	May 2007
Registered US Implants	22,878
Estimated Active US Implants	9,693
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	1	<0.01%	0	0.00%
Conductor Fracture	0	0.00%	16	0.07%
Lead Dislodgement	4	0.02%	48	0.21%
Failure to Capture	3	0.01%	43	0.19%
Oversensing	2	<0.01%	87	0.38%
Failure to Sense	8	0.03%	24	0.10%
Insulation Breach	0	0.00%	6	0.03%
Abnormal Pacing Impedance	0	0.00%	19	0.08%
Extracardiac Stimulation	0	0.00%	3	0.01%
Other	2	<0.01%	3	0.01%
Total	20	0.09%	249	1.09%
Total Returned for Analysis	16		73	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	13	0.06%
Insulation Breach	31	0.14%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	53	0.23%
Total	97	0.42%



YEAR	2	4	6	8	10	AT 124 MONTHS
SURVIVAL PROBABILITY	99.70%	99.47%	99.06%	98.50%	98.12%	97.96%
± 1 STANDARD ERROR	0.04%	0.05%	0.08%	0.10%	0.15%	0.21%
SAMPLE SIZE	18,670	15,160	12,620	9,520	2,440	250

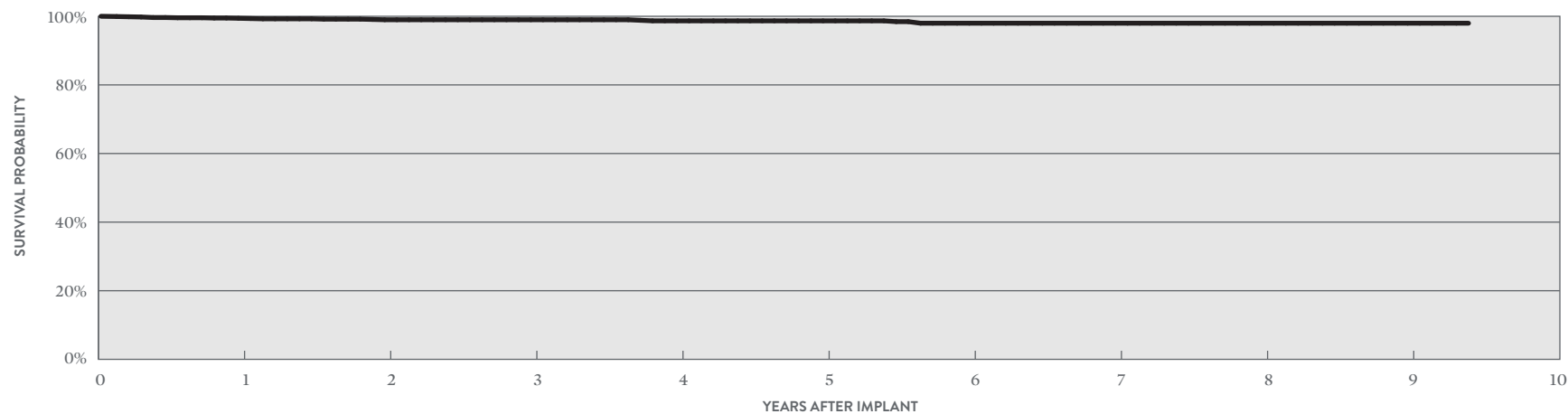
Pacing Leads

ACTIVELY MONITORED STUDY DATA

OptiSense™

MODELS 1699T & 1699TC

		QUALIFYING COMPLICATIONS			MALFUNCTIONS		
			QTY	RATE		QTY	RATE
US Regulatory Approval	May 2007	Abnormal Pacing Impedance	1	0.07%	Conductor Fracture	0	0.00%
Number of Devices Enrolled in Study	1,451	Conductor Fracture	3	0.21%	Insulation Breach	2	0.14%
Active Devices Enrolled in Study	337	Failure to Capture	4	0.28%	Crimps, Welds & Bonds	0	0.00%
Cumulative Months of Follow-up	71,422	Insulation Breach	1	0.07%	Other	0	0.00%
Insulation	Silicone	Lead Dislodgement	8	0.55%	Extrinsic Factors	6	0.41%
Type and/or Fixation	Active	Oversensing	1	0.07%	Total	8	0.55%
Polarity	Bipolar						
Steroid	Yes						



YEAR	1	2	3	4	5	6	7	8	9	AT 113 MONTHS
SURVIVAL PROBABILITY	99.42%	98.99%	98.99%	98.67%	98.67%	97.98%	97.98%	97.98%	97.98%	97.98%
± 1 STANDARD ERROR	0.19%	0.27%	0.28%	0.36%	0.36%	0.53%	0.53%	0.53%	0.53%	0.53%
SAMPLE SIZE	1,360	1,160	940	680	510	430	380	310	180	60

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

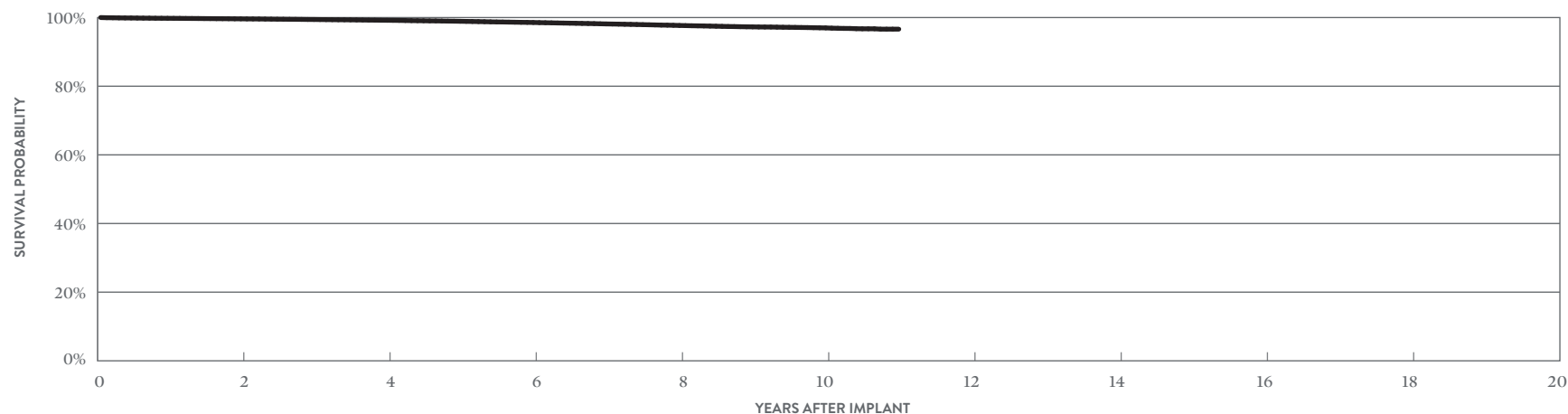
Tendril™ ST Optim™

MODELS 1888T & 1888TC

US Regulatory Approval	June 2006
Registered US Implants	301,676
Estimated Active US Implants	137,588
Insulation	Optim™*
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	39	0.01%	40	0.01%
Conductor Fracture	7	<0.01%	222	0.07%
Lead Dislodgement	156	0.05%	540	0.18%
Failure to Capture	85	0.03%	749	0.25%
Oversensing	16	<0.01%	1776	0.59%
Failure to Sense	14	<0.01%	106	0.04%
Insulation Breach	7	<0.01%	300	0.10%
Abnormal Pacing Impedance	9	<0.01%	211	0.07%
Extracardiac Stimulation	5	<0.01%	36	0.01%
Other	40	0.01%	114	0.04%
Total	378	0.13%	4094	1.36%
Total Returned for Analysis	200		1184	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	40	0.01%
Insulation Breach	760	0.25%
Crimps, Welds & Bonds	1	<0.01%
Other	12	<0.01%
Extrinsic Factors	781	0.26%
Total	1594	0.53%



YEAR	2	4	6	8	10	AT 132 MONTHS
SURVIVAL PROBABILITY	99.62%	99.20%	98.56%	97.69%	96.98%	96.62%
± 1 STANDARD ERROR	0.01%	0.02%	0.03%	0.04%	0.07%	0.14%
SAMPLE SIZE	242,080	181,770	126,570	73,650	18,020	260

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

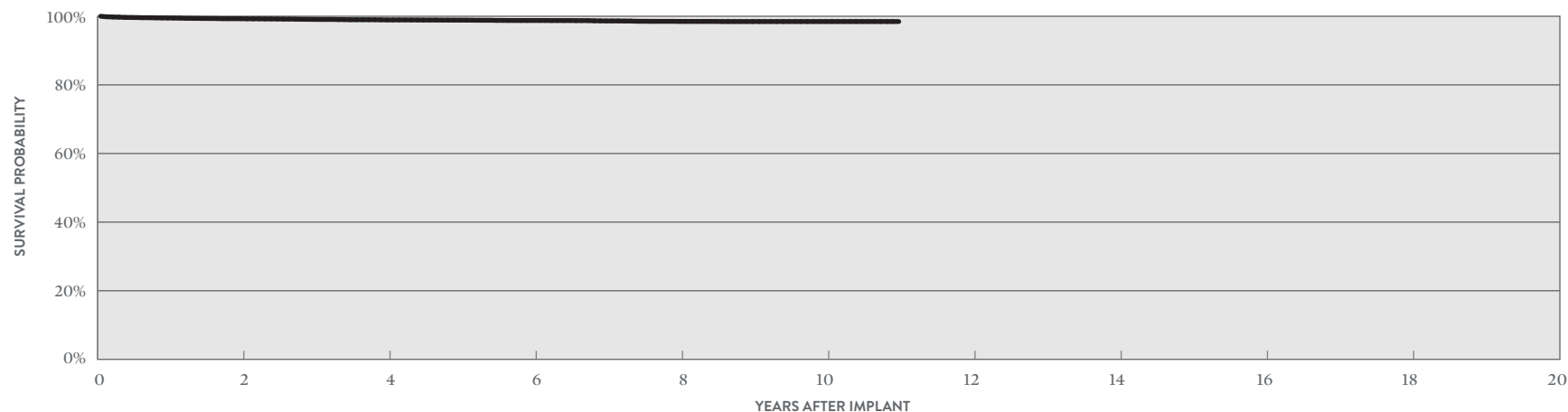
Pacing Leads

ACTIVELY MONITORED STUDY DATA

Tendril™ ST Optim™

MODELS 1888T & 1888TC

		QUALIFYING COMPLICATIONS			MALFUNCTIONS		
			QTY	RATE		QTY	RATE
US Regulatory Approval	June 2006	Abnormal Pacing Impedance	6	0.04%	Conductor Fracture	3	0.02%
Number of Devices Enrolled in Study	14,503	Cardiac Perforation	2	0.01%	Insulation Breach	24	0.17%
Active Devices Enrolled in Study	4,601	Conductor Fracture	8	0.06%	Crimps, Welds & Bonds	0	0.00%
Cumulative Months of Follow-up	822,216	Extracardiac Stimulation	4	0.03%	Other	0	0.00%
Insulation	Optim™*	Failure to Capture	19	0.13%	Extrinsic Factors	35	0.24%
Type and/or Fixation	Active	Failure to Sense	4	0.03%	Total	62	0.43%
Polarity	Bipolar	Insulation Breach	26	0.18%			
Steroid	Yes	Lead Dislodgement	58	0.40%			
		Oversensing	22	0.15%			
		Skin Erosion	1	<0.01%			



YEAR	2	4	6	8	10	AT 132 MONTHS
SURVIVAL PROBABILITY	99.29%	98.93%	98.77%	98.49%	98.46%	98.46%
± 1 STANDARD ERROR	0.07%	0.09%	0.11%	0.13%	0.14%	0.14%
SAMPLE SIZE	11,880	7,610	5,260	4,370	1,860	80

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

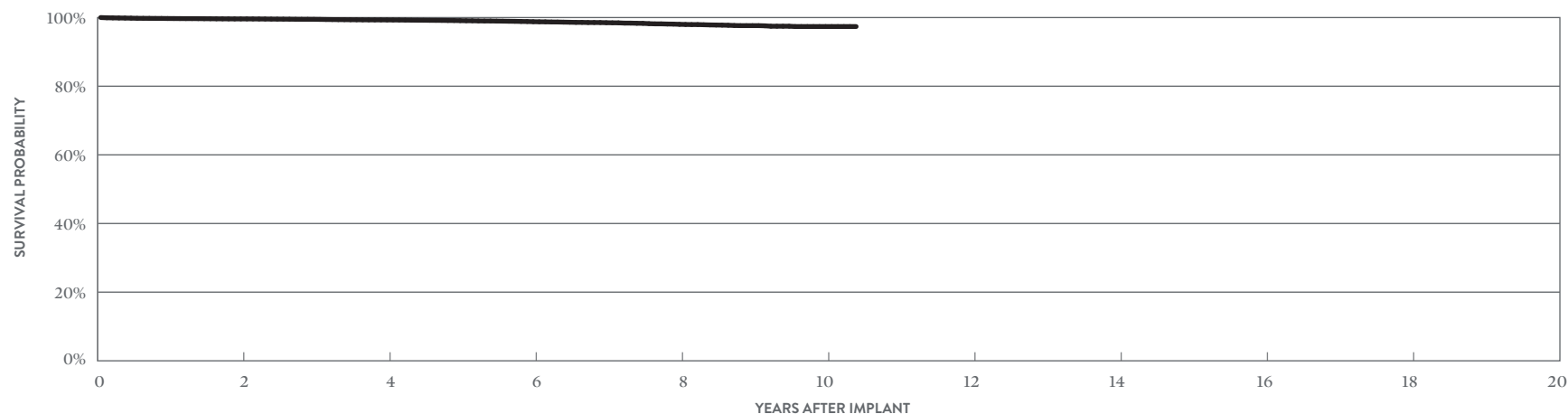
Tendril™ ST Optim™

MODELS 1882T & 1882TC

US Regulatory Approval	June 2006
Registered US Implants	48,680
Estimated Active US Implants	26,686
Insulation	Optim™*
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	3	<0.01%	3	<0.01%
Conductor Fracture	0	0.00%	12	0.02%
Lead Dislodgement	44	0.09%	129	0.26%
Failure to Capture	10	0.02%	68	0.14%
Oversensing	5	0.01%	159	0.33%
Failure to Sense	4	<0.01%	20	0.04%
Insulation Breach	0	0.00%	33	0.07%
Abnormal Pacing Impedance	1	<0.01%	11	0.02%
Extracardiac Stimulation	0	0.00%	3	<0.01%
Other	13	0.03%	22	0.05%
Total	80	0.16%	460	0.94%
Total Returned for Analysis	45		149	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	2	<0.01%
Insulation Breach	57	0.12%
Crimps, Welds & Bonds	0	0.00%
Other	3	<0.01%
Extrinsic Factors	124	0.25%
Total	186	0.38%



YEAR	2	4	6	8	10	AT 125 MONTHS
SURVIVAL PROBABILITY	99.60%	99.30%	98.79%	98.02%	97.38%	97.38%
± 1 STANDARD ERROR	0.03%	0.04%	0.07%	0.12%	0.21%	0.21%
SAMPLE SIZE	36,770	24,320	13,890	6,150	1,280	240

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

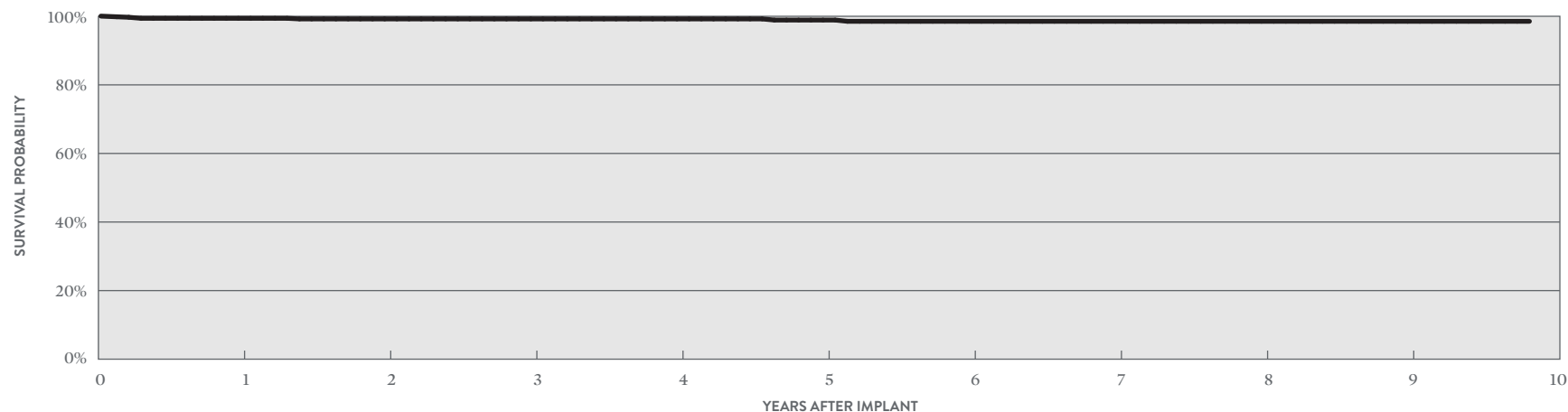
Pacing Leads

ACTIVELY MONITORED STUDY DATA

Tendril™ ST Optim™

MODELS 1882T & 1882TC

		QUALIFYING COMPLICATIONS			MALFUNCTIONS		
			QTY	RATE		QTY	RATE
US Regulatory Approval	June 2006	Extracardiac Stimulation	1	0.14%	Conductor Fracture	0	0.00%
Number of Devices Enrolled in Study	690	Failure to Capture	1	0.14%	Insulation Breach	2	0.29%
Active Devices Enrolled in Study	205	Lead Dislodgement	2	0.29%	Crimps, Welds & Bonds	0	0.00%
Cumulative Months of Follow-up	38,104	Oversensing	2	0.29%	Other	0	0.00%
Insulation	Optim™*	Skin Erosion	1	0.14%	Extrinsic Factors	0	0.00%
Type and/or Fixation	Active				Total	2	0.29%
Polarity	Bipolar						
Steroid	Yes						



YEAR	1	2	3	4	5	6	7	8	9	AT 118 MONTHS
SURVIVAL PROBABILITY	99.41%	99.23%	99.23%	99.23%	98.90%	98.53%	98.53%	98.53%	98.53%	98.53%
± 1 STANDARD ERROR	0.30%	0.34%	0.34%	0.34%	0.47%	0.60%	0.60%	0.60%	0.60%	0.60%
SAMPLE SIZE	650	560	460	380	310	250	220	180	130	50

*Optim™ lead insulation is a copolymer of silicone and polyurethane.

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

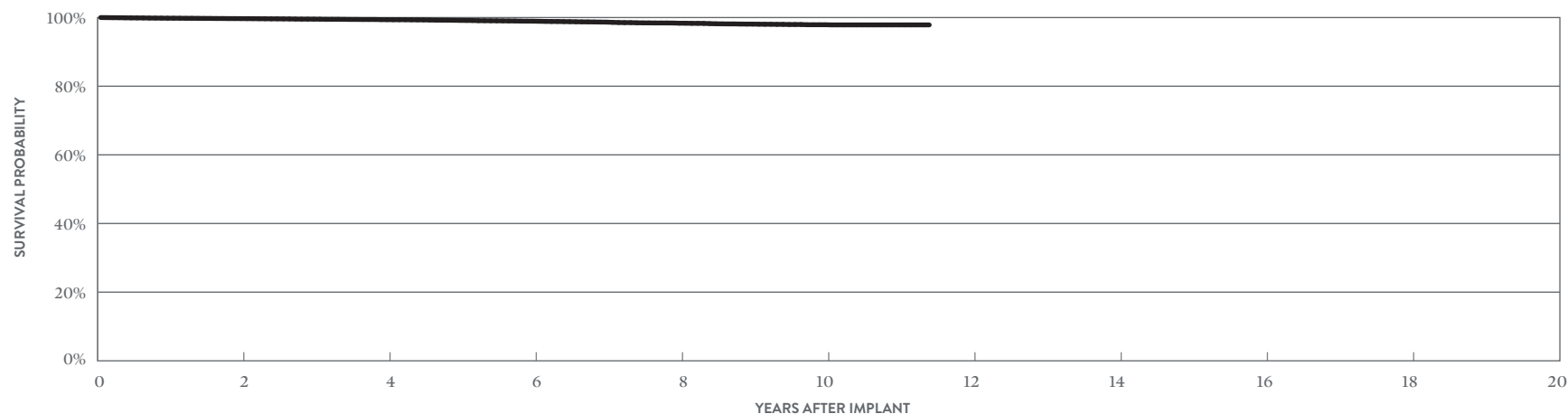
Tendril™

MODELS 1782T & 1782TC

US Regulatory Approval	February 2006
Registered US Implants	16,404
Estimated Active US Implants	6,479
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	6	0.04%	0	0.00%
Conductor Fracture	0	0.00%	5	0.03%
Lead Dislodgement	13	0.08%	49	0.30%
Failure to Capture	5	0.03%	42	0.26%
Oversensing	0	0.00%	46	0.28%
Failure to Sense	0	0.00%	7	0.04%
Insulation Breach	0	0.00%	4	0.02%
Abnormal Pacing Impedance	2	0.01%	15	0.09%
Extracardiac Stimulation	1	<0.01%	1	<0.01%
Other	2	0.01%	3	0.02%
Total	29	0.18%	172	1.05%
Total Returned for Analysis	16		61	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	1	<0.01%
Insulation Breach	34	0.21%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	49	0.30%
Total	84	0.51%



YEAR	2	4	6	8	10	AT 137 MONTHS
SURVIVAL PROBABILITY	99.69%	99.38%	98.94%	98.35%	97.90%	97.86%
± 1 STANDARD ERROR	0.05%	0.07%	0.10%	0.13%	0.17%	0.18%
SAMPLE SIZE	13,340	10,720	8,510	5,950	3,010	220

Pacing Leads

ACTIVELY MONITORED STUDY DATA

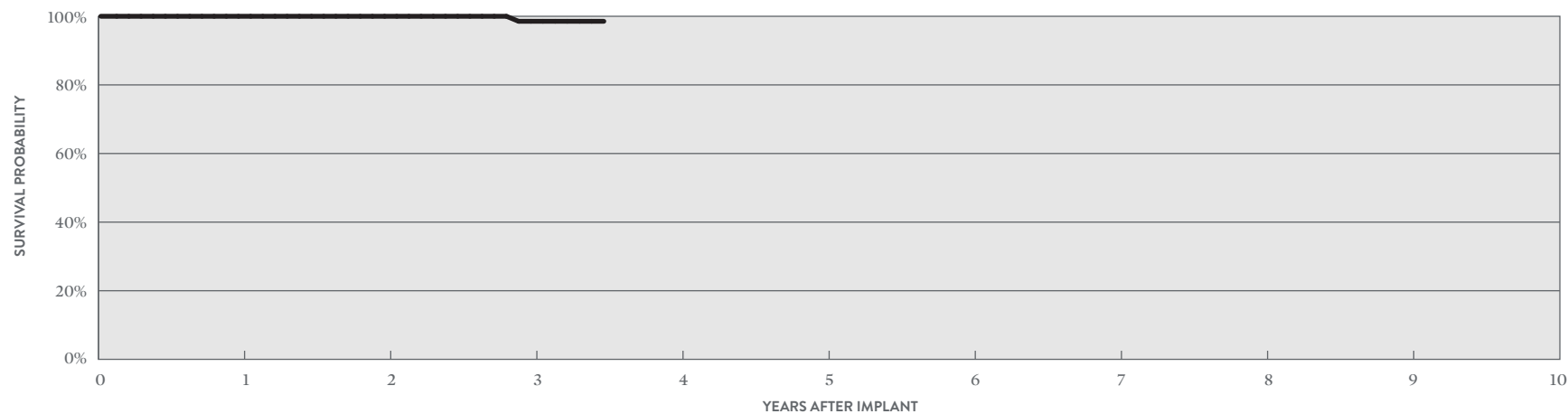
Tendril™

MODELS 1782T & 1782TC

US Regulatory Approval	February 2006
Number of Devices Enrolled in Study	165
Active Devices Enrolled in Study	15
Cumulative Months of Follow-up	5,762
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Oversensing	1	0.61%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Insulation Breach	1	0.61%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	0	0.00%
Total	1	0.61%



YEAR	1	2	3	AT 42 MONTHS
SURVIVAL PROBABILITY	100.00%	100.00%	98.54%	98.54%
± 1 STANDARD ERROR	0.00%	0.00%	1.45%	1.45%
SAMPLE SIZE	150	120	80	60

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

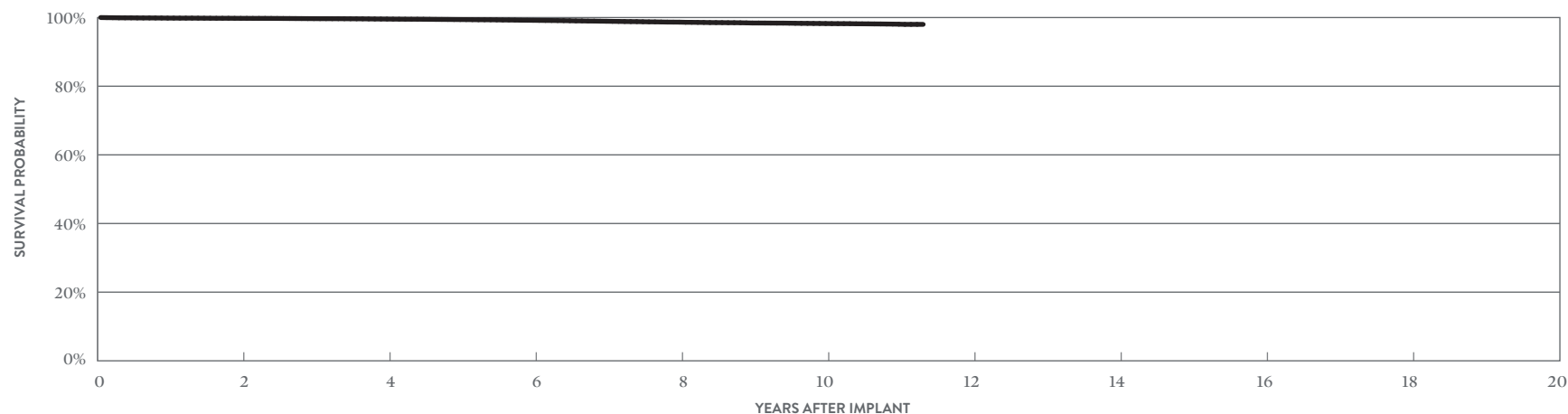
Tendril™

MODELS 1788T & 1788TC

US Regulatory Approval	February 2006
Registered US Implants	65,235
Estimated Active US Implants	24,261
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	12	0.02%	7	0.01%
Conductor Fracture	1	<0.01%	28	0.04%
Lead Dislodgement	31	0.05%	77	0.12%
Failure to Capture	30	0.05%	154	0.24%
Oversensing	2	<0.01%	176	0.27%
Failure to Sense	2	<0.01%	24	0.04%
Insulation Breach	1	<0.01%	32	0.05%
Abnormal Pacing Impedance	9	0.01%	45	0.07%
Extracardiac Stimulation	2	<0.01%	7	0.01%
Other	20	0.03%	25	0.04%
Total	110	0.17%	575	0.88%
Total Returned for Analysis	47		147	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	10	0.02%
Insulation Breach	110	0.17%
Crimps, Welds & Bonds	1	<0.01%
Other	1	<0.01%
Extrinsic Factors	101	0.15%
Total	223	0.34%



YEAR	2	4	6	8	10	AT 136 MONTHS
SURVIVAL PROBABILITY	99.77%	99.56%	99.19%	98.66%	98.25%	98.01%
± 1 STANDARD ERROR	0.02%	0.03%	0.04%	0.06%	0.08%	0.10%
SAMPLE SIZE	52,690	41,710	33,740	26,700	16,300	430

Pacing Leads

ACTIVELY MONITORED STUDY DATA

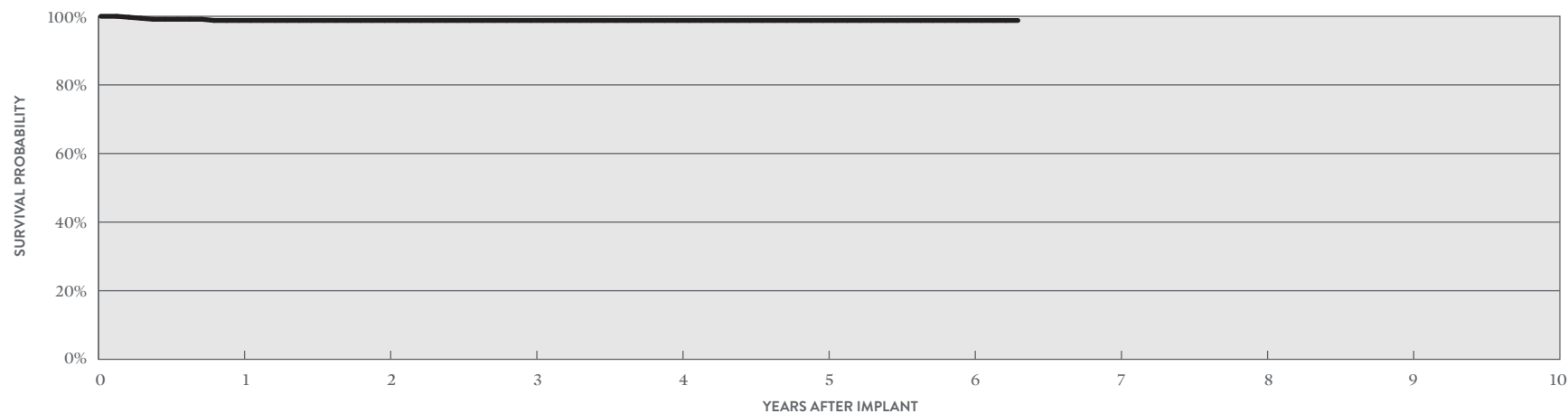
Tendril™

MODELS 1788T & 1788TC

US Regulatory Approval	February 2006
Number of Devices Enrolled in Study	363
Active Devices Enrolled in Study	41
Cumulative Months of Follow-up	12,360
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Extracardiac Stimulation	1	0.28%
Lead Dislodgement	3	0.83%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Insulation Breach	1	0.28%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	0	0.00%
Total	1	0.28%



YEAR	1	2	3	4	5	6	AT 76 MONTHS
SURVIVAL PROBABILITY	98.78%	98.78%	98.78%	98.78%	98.78%	98.78%	98.78%
± 1 STANDARD ERROR	0.61%	0.61%	0.61%	0.61%	0.61%	0.61%	0.61%
SAMPLE SIZE	310	240	170	100	70	60	50

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

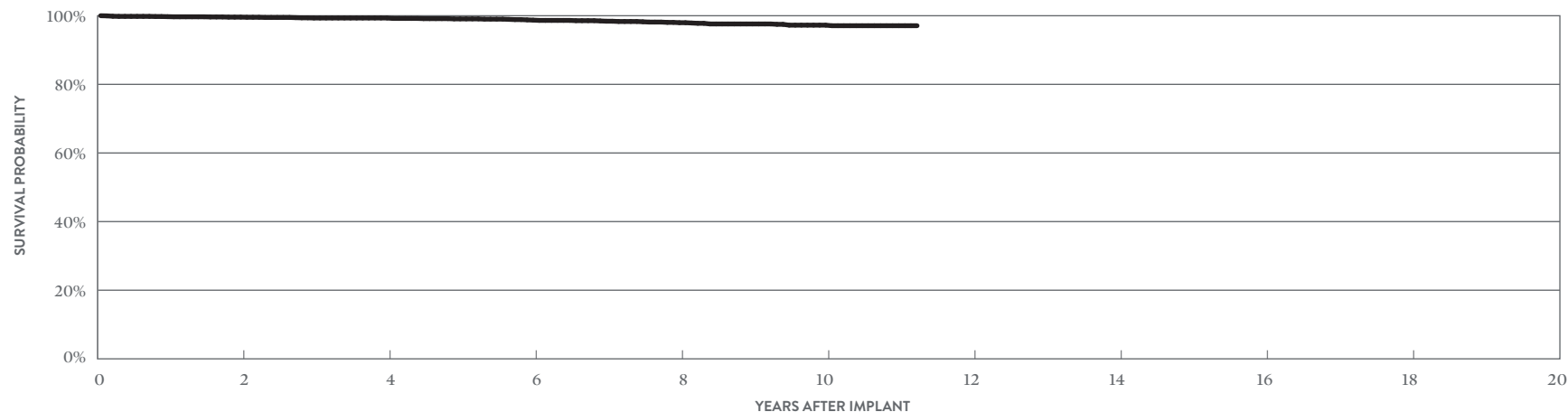
IsoFlex™ P

MODEL 1648T

US Regulatory Approval	April 2005
Registered US Implants	2,834
Estimated Active US Implants	996
Insulation	Polyurethane
Type and/or Fixation	Passive
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	0	0.00%
Conductor Fracture	0	0.00%	5	0.18%
Lead Dislodgement	2	0.07%	2	0.07%
Failure to Capture	2	0.07%	9	0.32%
Oversensing	0	0.00%	2	0.07%
Failure to Sense	1	0.04%	1	0.04%
Insulation Breach	0	0.00%	12	0.42%
Abnormal Pacing Impedance	0	0.00%	3	0.11%
Extracardiac Stimulation	1	0.04%	0	0.00%
Other	0	0.00%	5	0.18%
Total	6	0.21%	39	1.38%
Total Returned for Analysis	1		6	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Insulation Breach	11	0.39%
Crimps, Welds & Bonds	0	0.00%
Other	2	0.07%
Extrinsic Factors	5	0.18%
Total	18	0.64%



YEAR	2	4	6	8	10	AT 135 MONTHS
SURVIVAL PROBABILITY	99.59%	99.34%	98.72%	97.97%	97.25%	97.09%
± 1 STANDARD ERROR	0.13%	0.17%	0.26%	0.35%	0.45%	0.48%
SAMPLE SIZE	2,210	1,740	1,390	1,150	780	200

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

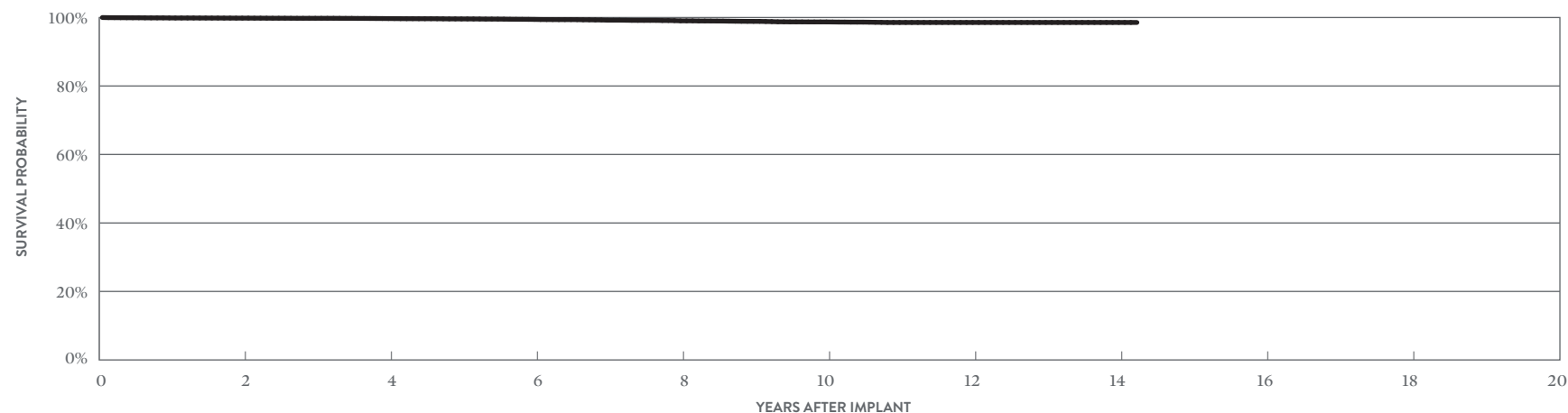
IsoFlex™ S

MODEL 1642T

US Regulatory Approval	May 2002
Registered US Implants	27,127
Estimated Active US Implants	9,232
Insulation	Silicone
Type and/or Fixation	Passive
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	0	0.00%	0	0.00%
Conductor Fracture	0	0.00%	7	0.03%
Lead Dislodgement	49	0.18%	42	0.15%
Failure to Capture	6	0.02%	59	0.22%
Oversensing	0	0.00%	36	0.13%
Failure to Sense	3	0.01%	16	0.06%
Insulation Breach	0	0.00%	6	0.02%
Abnormal Pacing Impedance	3	0.01%	11	0.04%
Extracardiac Stimulation	1	<0.01%	2	<0.01%
Other	0	0.00%	2	<0.01%
Total	62	0.23%	181	0.67%
Total Returned for Analysis	39		28	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Insulation Breach	21	0.08%
Crimps, Welds & Bonds	1	<0.01%
Other	2	<0.01%
Extrinsic Factors	20	0.07%
Total	44	0.16%



YEAR	2	4	6	8	10	12	14	AT 171 MONTHS
SURVIVAL PROBABILITY	99.82%	99.69%	99.44%	99.03%	98.71%	98.55%	98.55%	98.55%
± 1 STANDARD ERROR	0.03%	0.04%	0.06%	0.08%	0.11%	0.12%	0.12%	0.12%
SAMPLE SIZE	22,070	17,740	13,880	10,260	6,350	2,900	730	220

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

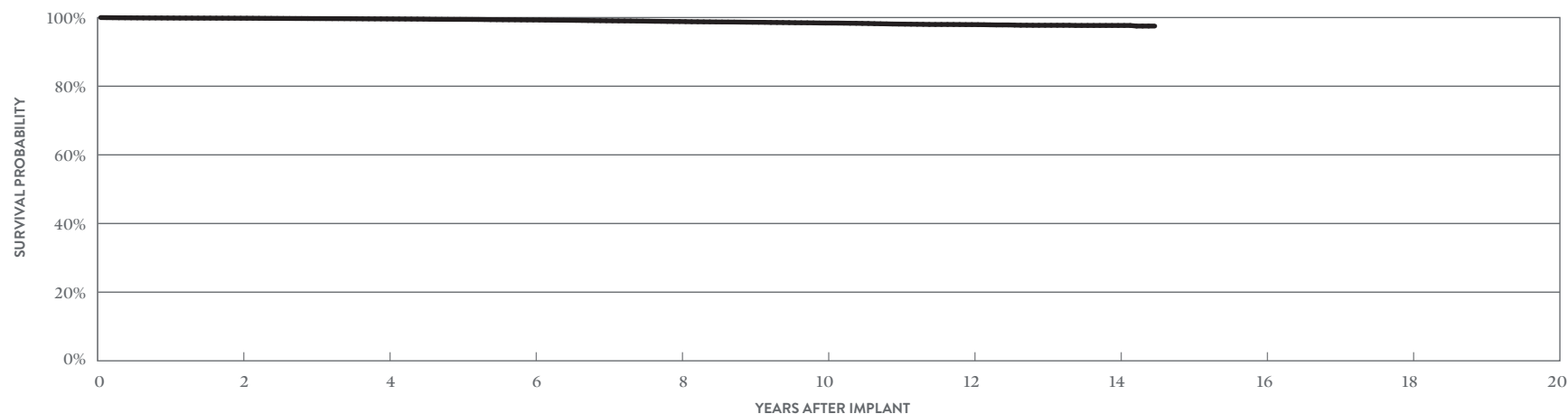
IsoFlex™ S

MODEL 1646T

US Regulatory Approval	May 2002
Registered US Implants	90,360
Estimated Active US Implants	29,805
Insulation	Silicone
Type and/or Fixation	Passive
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	4	<0.01%	2	<0.01%
Conductor Fracture	2	<0.01%	106	0.12%
Lead Dislodgement	37	0.04%	36	0.04%
Failure to Capture	33	0.04%	303	0.34%
Oversensing	0	0.00%	124	0.14%
Failure to Sense	2	<0.01%	12	0.01%
Insulation Breach	2	<0.01%	42	0.05%
Abnormal Pacing Impedance	6	<0.01%	112	0.12%
Extracardiac Stimulation	0	0.00%	7	<0.01%
Other	2	<0.01%	21	0.02%
Total	88	0.10%	765	0.85%
Total Returned for Analysis	38		102	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	21	0.02%
Insulation Breach	64	0.07%
Crimps, Welds & Bonds	0	0.00%
Other	6	<0.01%
Extrinsic Factors	64	0.07%
Total	155	0.17%



YEAR	2	4	6	8	10	12	14	AT 174 MONTHS
SURVIVAL PROBABILITY	99.80%	99.60%	99.30%	98.84%	98.40%	97.94%	97.69%	97.52%
± 1 STANDARD ERROR	0.02%	0.02%	0.03%	0.05%	0.07%	0.09%	0.12%	0.21%
SAMPLE SIZE	71,950	56,350	43,710	31,720	19,640	8,960	2,110	230

Pacing Leads

ACTIVELY MONITORED STUDY DATA

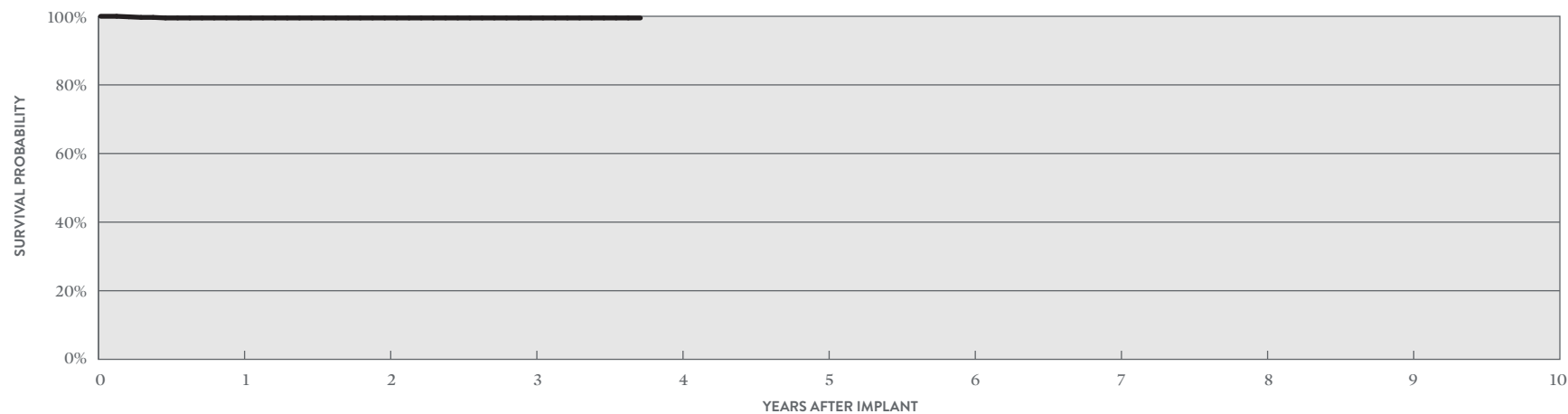
IsoFlex™ S

MODEL 1646T

US Regulatory Approval	May 2002
Number of Devices Enrolled in Study	641
Active Devices Enrolled in Study	3
Cumulative Months of Follow-up	15,791
Insulation	Silicone
Type and/or Fixation	Passive
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Failure to Capture	2	0.31%
Lead Dislodgement	1	0.16%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Insulation Breach	0	0.00%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	0	0.00%
Total	0	0.00%



YEAR	1	2	3	AT 45 MONTHS
SURVIVAL PROBABILITY	99.51%	99.51%	99.51%	99.51%
± 1 STANDARD ERROR	0.28%	0.28%	0.28%	0.28%
SAMPLE SIZE	570	410	250	60

Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

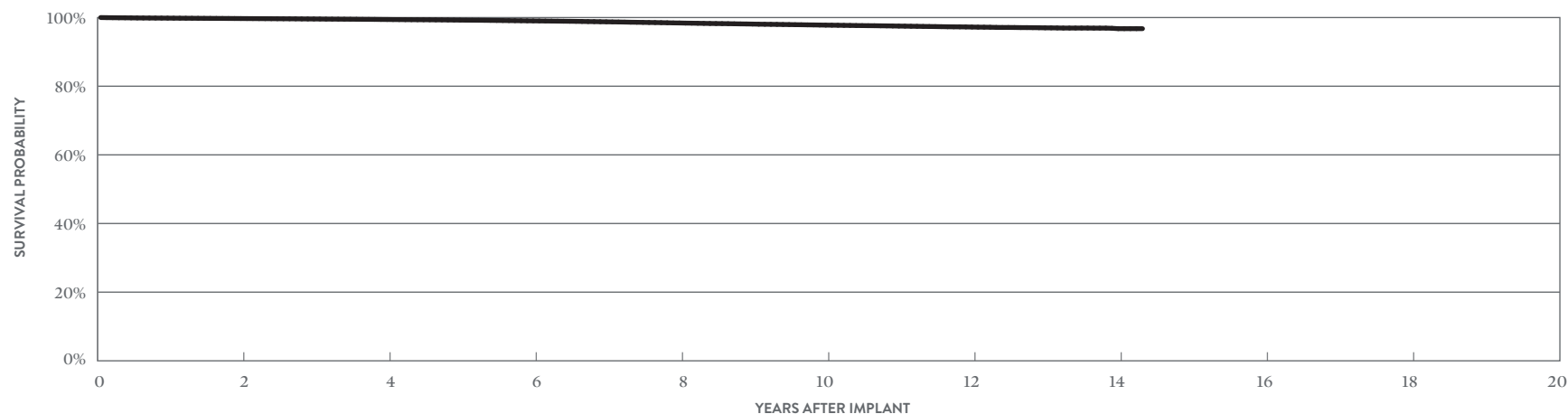
Tendril™ SDX

MODELS 1688T & 1688TC

US Regulatory Approval	June 2003
Registered US Implants	489,893
Estimated Active US Implants	191,315
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

	ACUTE OBSERVATIONS (POST IMPLANT, ≤30 DAYS)		CHRONIC COMPLICATIONS (>30 DAYS)	
	QTY	RATE	QTY	RATE
Cardiac Perforation	81	0.02%	40	<0.01%
Conductor Fracture	5	<0.01%	467	0.10%
Lead Dislodgement	311	0.06%	554	0.11%
Failure to Capture	193	0.04%	1329	0.27%
Oversensing	18	<0.01%	1425	0.29%
Failure to Sense	33	<0.01%	137	0.03%
Insulation Breach	10	<0.01%	213	0.04%
Abnormal Pacing Impedance	28	<0.01%	535	0.11%
Extracardiac Stimulation	7	<0.01%	43	<0.01%
Other	62	0.01%	161	0.03%
Total	748	0.15%	4904	1.00%
Total Returned for Analysis	336		1328	

MALFUNCTIONS	QTY	RATE
Conductor Fracture	203	0.04%
Insulation Breach	849	0.17%
Crimps, Welds & Bonds	2	<0.01%
Other	18	<0.01%
Extrinsic Factors	767	0.16%
Total	1839	0.38%



YEAR	2	4	6	8	10	12	14	AT 172 MONTHS
SURVIVAL PROBABILITY	99.71%	99.43%	99.03%	98.42%	97.79%	97.23%	96.76%	96.76%
± 1 STANDARD ERROR	0.01%	0.01%	0.02%	0.03%	0.04%	0.05%	0.08%	0.12%
SAMPLE SIZE	388,540	284,560	200,530	137,710	86,590	43,100	6,550	390

Pacing Leads

ACTIVELY MONITORED STUDY DATA

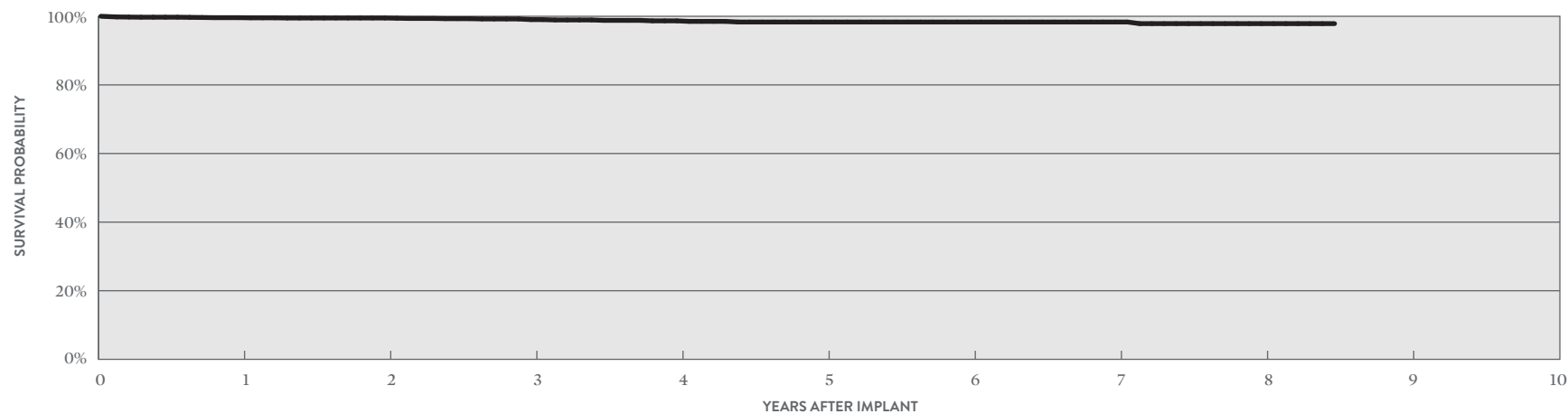
Tendril™ SDX

MODELS 1688T & 1688TC

US Regulatory Approval	June 2003
Number of Devices Enrolled in Study	2,645
Active Devices Enrolled in Study	417
Cumulative Months of Follow-up	92,643
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes

QUALIFYING COMPLICATIONS	QTY	RATE
Abnormal Pacing Impedance	5	0.19%
Conductor Fracture	2	0.08%
Failure to Capture	3	0.11%
Insulation Breach	3	0.11%
Lead Dislodgement	6	0.23%
Oversensing	3	0.11%
Pericardial Effusion	1	0.04%
Skin Erosion	1	0.04%

MALFUNCTIONS	QTY	RATE
Conductor Fracture	0	0.00%
Insulation Breach	4	0.15%
Crimps, Welds & Bonds	0	0.00%
Other	0	0.00%
Extrinsic Factors	5	0.19%
Total	9	0.34%



YEAR	1	2	3	4	5	6	7	8	AT 102 MONTHS
SURVIVAL PROBABILITY	99.60%	99.50%	99.03%	98.67%	98.33%	98.33%	98.33%	97.86%	97.86%
± 1 STANDARD ERROR	0.13%	0.14%	0.20%	0.32%	0.40%	0.40%	0.40%	0.61%	0.61%
SAMPLE SIZE	2,390	1,840	1,300	840	540	410	300	170	50

Pacing Leads

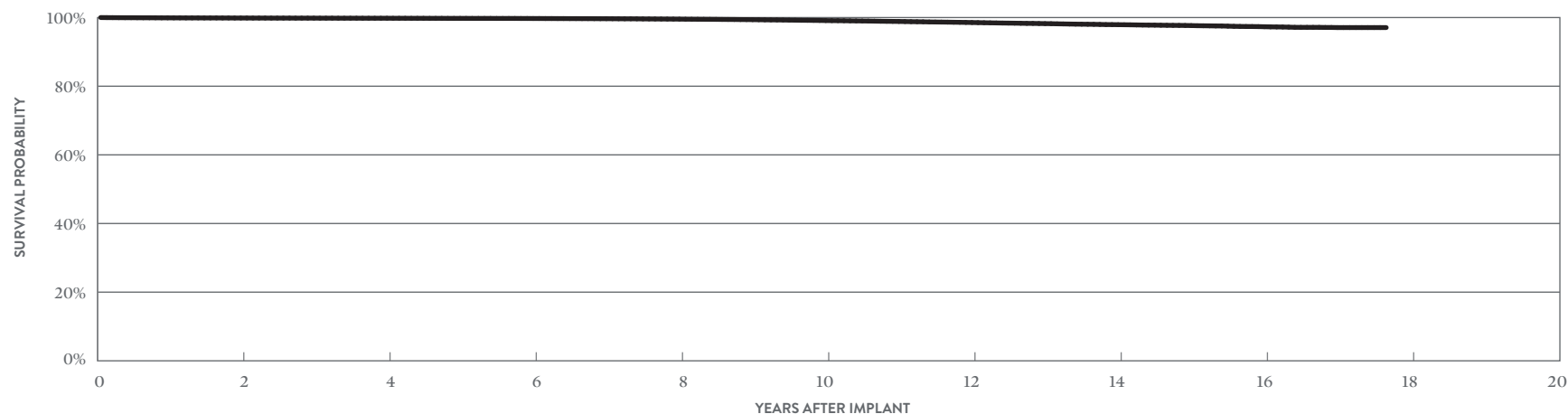
CUSTOMER REPORTED PERFORMANCE DATA

Tendril™ SDX

MODELS 1488T & 1488TC

US Regulatory Approval	March 2000
Registered US Implants	270,810
Estimated Active US Implants	57,182
Insulation	Silicone
Type and/or Fixation	Active
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None

MALFUNCTIONS	QTY	RATE
Conductor Fracture	158	0.06%
Insulation Breach	304	0.11%
Crimps, Welds & Bonds	5	<0.01%
Other	3	<0.01%
Extrinsic Factors	360	0.13%
Total	830	0.31%



YEAR	2	4	6	8	10	12	14	16	AT 212 MONTHS
SURVIVAL PROBABILITY	99.86%	99.80%	99.72%	99.54%	99.12%	98.55%	97.92%	97.32%	97.08%
± 1 STANDARD ERROR	0.01%	0.01%	0.01%	0.02%	0.03%	0.04%	0.05%	0.07%	0.10%
SAMPLE SIZE	221,920	177,430	137,150	104,580	81,200	63,070	42,260	15,920	290

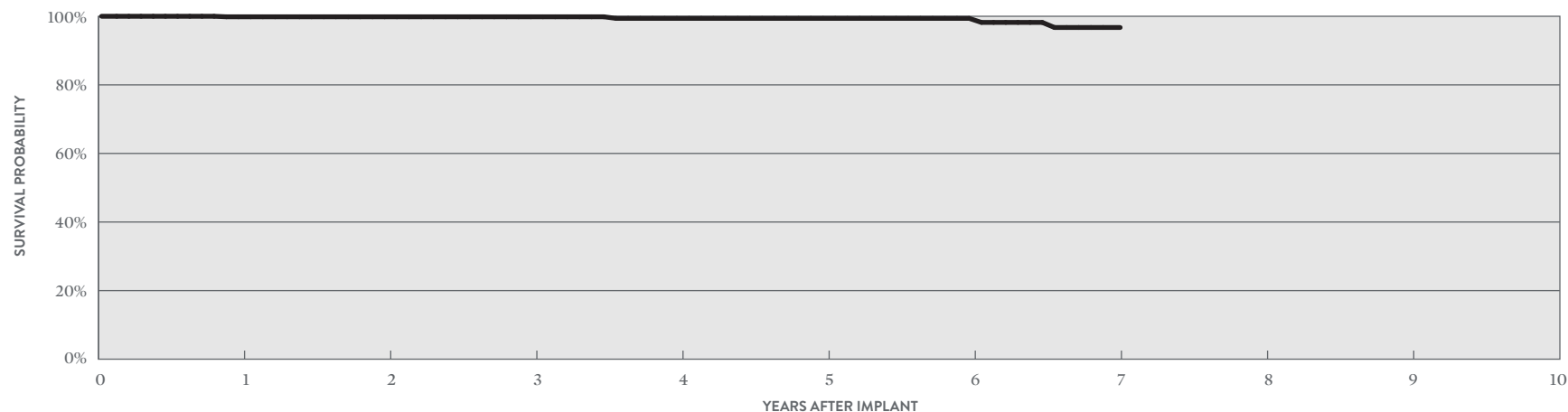
Pacing Leads

ACTIVELY MONITORED STUDY DATA

Tendril™ SDX

MODELS 1488T & 1488TC

		QUALIFYING COMPLICATIONS		MALFUNCTIONS			
			QTY	RATE		QTY	RATE
US Regulatory Approval	March 2000	Failure to Capture	1	0.12%	Conductor Fracture	0	0.00%
Number of Devices Enrolled in Study	803	Insulation Breach	2	0.25%	Insulation Breach	4	0.50%
Active Devices Enrolled in Study	50	Oversensing	1	0.12%	Crimps, Welds & Bonds	0	0.00%
Cumulative Months of Follow-up	26,898				Other	0	0.00%
Insulation	Silicone				Extrinsic Factors	1	0.12%
Type and/or Fixation	Active				Total	5	0.62%
Polarity	Bipolar						
Steroid	Yes						



YEAR	1	2	3	4	5	6	7
SURVIVAL PROBABILITY	99.85%	99.85%	99.85%	99.39%	99.39%	99.39%	96.73%
± 1 STANDARD ERROR	0.15%	0.15%	0.15%	0.48%	0.48%	0.48%	1.93%
SAMPLE SIZE	730	580	400	220	120	90	50

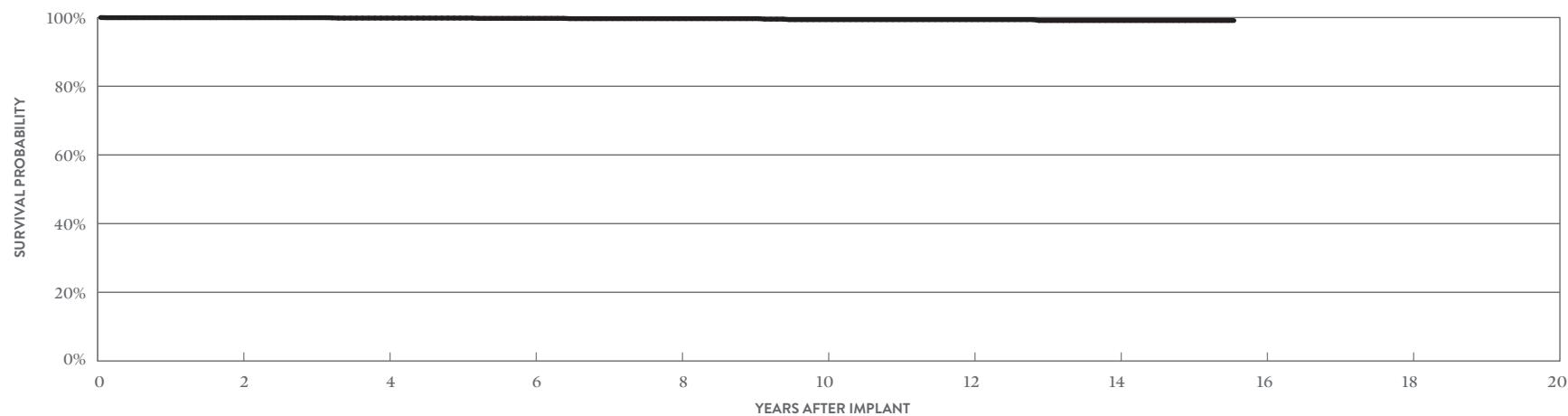
Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

AV Plus™ DX

MODEL 1368

US Regulatory Approval	May 1999
Registered US Implants	2,827
Estimated Active US Implants	820
Insulation	Silicone
Type and/or Fixation	Passive
Atrial Polarity	Bipolar
Ventricular Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None



YEAR	2	4	6	8	10	12	14	AT 187 MONTHS
SURVIVAL PROBABILITY	99.96%	99.85%	99.77%	99.68%	99.39%	99.39%	99.14%	99.14%
± 1 STANDARD ERROR	0.04%	0.09%	0.12%	0.15%	0.25%	0.25%	0.36%	0.36%
SAMPLE SIZE	2,170	1,670	1,230	930	690	490	340	200

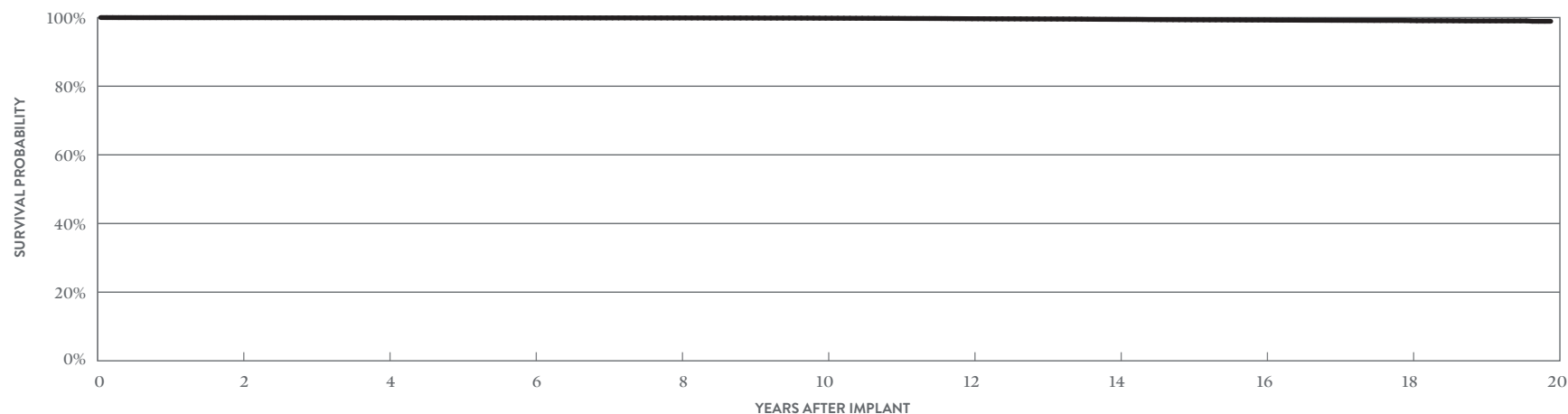
Pacing Leads

CUSTOMER REPORTED PERFORMANCE DATA

Passive Plus™ DX

MODELS 1336T, 1342T & 1346T

US Regulatory Approval	(1336T) August 1999; (1342T, 1346T) January 1998
Registered US Implants	198,482
Estimated Active US Implants	34,287
Insulation	Silicone
Type and/or Fixation	Passive
Polarity	Bipolar
Steroid	Yes
Number of US Advisories	None



YEAR	2	4	6	8	10	12	14	16	18	AT 239 MONTHS
SURVIVAL PROBABILITY	99.96%	99.94%	99.91%	99.89%	99.82%	99.65%	99.46%	99.28%	99.11%	98.91%
± 1 STANDARD ERROR	0.00%	0.01%	0.01%	0.01%	0.01%	0.02%	0.03%	0.04%	0.05%	0.13%
SAMPLE SIZE	160,540	127,420	98,940	75,770	57,450	44,550	34,610	22,210	10,040	270

SUMMARY INFORMATION
Pacing Leads

Pacing Leads

Survival Probability Summary

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
LPA1200M	Tendril MRI™	99.78%	99.24%	99.17%	99.03%	98.61%					
2088TC	Tendril™ STS	99.77%	99.62%	99.44%	99.24%	99.02%	98.76%	98.46%	97.92%		
1999	OptiSense™ Optim™	99.67%	99.51%	99.32%	99.16%	99.00%	98.80%	98.31%			
1944	IsoFlex™ Optim™	99.69%	99.54%	99.41%	99.30%	99.11%	98.84%	98.37%	97.95%		
1948	IsoFlex™ Optim™	99.80%	99.63%	99.43%	99.18%	98.85%	98.50%	97.97%	97.29%	96.71%	
1699T/TC	OptiSense™	99.80%	99.70%	99.55%	99.47%	99.27%	99.06%	98.79%	98.50%	98.34%	98.12%
1888T/TC	Tendril™ ST Optim™	99.78%	99.62%	99.43%	99.20%	98.91%	98.56%	98.14%	97.69%	97.28%	96.98%
1882T/TC	Tendril™ ST Optim™	99.72%	99.60%	99.48%	99.30%	99.10%	98.79%	98.50%	98.02%	97.64%	97.38%
1782T/TC	Tendril™	99.81%	99.69%	99.53%	99.38%	99.17%	98.94%	98.67%	98.35%	98.09%	97.90%
1788T/TC	Tendril™	99.84%	99.77%	99.68%	99.56%	99.41%	99.19%	98.92%	98.66%	98.41%	98.25%
1648T	IsoFlex™ P	99.73%	99.59%	99.34%	99.34%	99.08%	98.72%	98.40%	97.97%	97.60%	97.25%
1642T	IsoFlex™ S	99.87%	99.82%	99.77%	99.69%	99.58%	99.44%	99.22%	99.03%	98.87%	98.71%
1646T	IsoFlex™ S	99.86%	99.80%	99.70%	99.60%	99.47%	99.30%	99.07%	98.84%	98.63%	98.40%
1688T/TC	Tendril™ SDX	99.83%	99.71%	99.58%	99.43%	99.25%	99.03%	98.76%	98.42%	98.10%	97.79%
1488T/TC	Tendril™ SDX	99.91%	99.86%	99.83%	99.80%	99.76%	99.72%	99.66%	99.54%	99.38%	99.12%
1368	AV Plus™ DX	99.96%	99.96%	99.96%	99.85%	99.85%	99.77%	99.68%	99.68%	99.68%	99.39%
1336T, 1342T, 1346T	Passive Plus™ DX	99.97%	99.96%	99.95%	99.94%	99.93%	99.91%	99.90%	99.89%	99.86%	99.82%

Pacing Leads

Acute Observation Summary

POST IMPLANT ≤30 DAYS

MODELS	US REGULATORY APPROVAL	REGISTERED US IMPLANTS	ESTIMATED ACTIVE US IMPLANTS	CARDIAC PERFORATION		CONDUCTOR FRACTURE		LEAD DISLODGEMENT		FAILURE TO CAPTURE		OVERSENSING		FAILURE TO SENSE		INSULATION BREACH		ABNORMAL PACING IMPEDANCE		EXTRACARDIAC STIMULATION		OTHER		TOTAL		TOTAL RETURNED FOR ANALYSIS
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	
LPA1200M	Jan-17	57,116	48,436	12	0.02%	0	0.00%	95	0.17%	6	0.01%	1	<0.01%	6	0.01%	0	0.00%	0	0.00%	2	<0.01%	9	0.02%	131	0.23%	29
2088TC	May-09	585,050	382,081	83	0.01%	6	<0.01%	582	0.10%	148	0.03%	42	<0.01%	20	<0.01%	12	<0.01%	31	<0.01%	4	<0.01%	99	0.02%	1027	0.18%	460
1999	May-07	46,528	28,723	4	<0.01%	0	0.00%	64	0.14%	8	0.02%	6	0.01%	3	<0.01%	1	<0.01%	0	0.00%	0	0.00%	12	0.03%	98	0.21%	52
1944	Mar-08	16,578	9,638	0	0.00%	0	0.00%	67	0.40%	8	0.05%	0	0.00%	2	0.01%	0	0.00%	0	0.00%	2	0.01%	1	<0.01%	80	0.48%	45
1948	Mar-08	63,721	37,190	3	<0.01%	0	0.00%	51	0.08%	25	0.04%	1	<0.01%	2	<0.01%	4	<0.01%	1	<0.01%	2	<0.01%	6	<0.01%	95	0.15%	51
1699T/TC	May-07	22,878	9,693	1	<0.01%	0	0.00%	4	0.02%	3	0.01%	2	<0.01%	8	0.03%	0	0.00%	0	0.00%	0	0.00%	2	<0.01%	20	0.09%	16
1888T/TC	Jun-06	301,676	137,588	39	0.01%	7	<0.01%	156	0.05%	85	0.03%	16	<0.01%	14	<0.01%	7	<0.01%	9	<0.01%	5	<0.01%	40	0.01%	378	0.13%	200
1882T/TC	Jun-06	48,680	26,686	3	<0.01%	0	0.00%	44	0.09%	10	0.02%	5	0.01%	4	<0.01%	0	0.00%	1	<0.01%	0	0.00%	13	0.03%	80	0.16%	45
1782T/TC	Feb-06	16,404	6,479	6	0.04%	0	0.00%	13	0.08%	5	0.03%	0	0.00%	0	0.00%	0	0.00%	2	0.01%	1	<0.01%	2	0.01%	29	0.18%	16
1788T/TC	Feb-06	65,235	24,261	12	0.02%	1	<0.01%	31	0.05%	30	0.05%	2	<0.01%	2	<0.01%	1	<0.01%	9	0.01%	2	<0.01%	20	0.03%	110	0.17%	47
1648T	Apr-05	2,834	996	0	0.00%	0	0.00%	2	0.07%	2	0.07%	0	0.00%	1	0.04%	0	0.00%	0	0.00%	1	0.04%	0	0.00%	6	0.21%	1
1642T	May-02	27,127	9,232	0	0.00%	0	0.00%	49	0.18%	6	0.02%	0	0.00%	3	0.01%	0	0.00%	3	0.01%	1	<0.01%	0	0.00%	62	0.23%	39
1646T	May-02	90,360	29,805	4	<0.01%	2	<0.01%	37	0.04%	33	0.04%	0	0.00%	2	<0.01%	2	<0.01%	6	<0.01%	0	0.00%	2	<0.01%	88	0.10%	38
1688T/TC	Jun-03	489,893	191,315	81	0.02%	5	<0.01%	311	0.06%	193	0.04%	18	<0.01%	33	<0.01%	10	<0.01%	28	<0.01%	7	<0.01%	62	0.01%	748	0.15%	336

Chronic Complication Summary

>30 DAYS

MODELS	US REGULATORY APPROVAL	REGISTERED US IMPLANTS	ESTIMATED ACTIVE US IMPLANTS	CARDIAC PERFORATION		CONDUCTOR FRACTURE		LEAD DISLODGEMENT		FAILURE TO CAPTURE		OVERSENSING		FAILURE TO SENSE		INSULATION BREACH		ABNORMAL PACING IMPEDANCE		EXTRACARDIAC STIMULATION		OTHER		TOTAL		TOTAL RETURNED FOR ANALYSIS
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	
LPA1200M	Jan-17	57,116	48,436	3	<0.01%	0	0.00%	49	0.09%	24	0.04%	22	0.04%	7	0.01%	0	0.00%	2	<0.01%	2	<0.01%	5	<0.01%	114	0.20%	25
2088TC	May-09	585,050	382,081	52	<0.01%	192	0.03%	826	0.14%	635	0.11%	1726	0.30%	95	0.02%	169	0.03%	128	0.02%	29	<0.01%	129	0.02%	3981	0.68%	1387
1999	May-07	46,528	28,723	0	0.00%	7	0.02%	154	0.33%	51	0.11%	149	0.32%	21	0.05%	25	0.05%	11	0.02%	1	<0.01%	16	0.03%	435	0.93%	171
1944	Mar-08	16,578	9,638	1	<0.01%	6	0.04%	46	0.28%	29	0.17%	49	0.30%	6	0.04%	5	0.03%	1	<0.01%	1	<0.01%	2	0.01%	146	0.88%	27
1948	Mar-08	63,721	37,190	10	0.02%	69	0.11%	63	0.10%	135	0.21%	189	0.30%	2	<0.01%	43	0.07%	30	0.05%	5	<0.01%	12	0.02%	558	0.88%	110
1699T/TC	May-07	22,878	9,693	0	0.00%	16	0.07%	48	0.21%	43	0.19%	87	0.38%	24	0.10%	6	0.03%	19	0.08%	3	0.01%	3	0.01%	249	1.09%	73
1888T/TC	Jun-06	301,676	137,588	40	0.01%	222	0.07%	540	0.18%	749	0.25%	1776	0.59%	106	0.04%	300	0.10%	211	0.07%	36	0.01%	114	0.04%	4094	1.36%	1184
1882T/TC	Jun-06	48,680	26,686	3	<0.01%	12	0.02%	129	0.26%	68	0.14%	159	0.33%	20	0.04%	33	0.07%	11	0.02%	3	<0.01%	22	0.05%	460	0.94%	149
1782T/TC	Feb-06	16,404	6,479	0	0.00%	5	0.03%	49	0.30%	42	0.26%	46	0.28%	7	0.04%	4	0.02%	15	0.09%	1	<0.01%	3	0.02%	172	1.05%	61
1788T/TC	Feb-06	65,235	24,261	7	0.01%	28	0.04%	77	0.12%	154	0.24%	176	0.27%	24	0.04%	32	0.05%	45	0.07%	7	0.01%	25	0.04%	575	0.88%	147
1648T	Apr-05	2,834	996	0	0.00%	5	0.18%	2	0.07%	9	0.32%	2	0.07%	1	0.04%	12	0.42%	3	0.11%	0	0.00%	5	0.18%	39	1.38%	6
1642T	May-02	27,127	9,232	0	0.00%	7	0.03%	42	0.15%	59	0.22%	36	0.13%	16	0.06%	6	0.02%	11	0.04%	2	<0.01%	2	<0.01%	181	0.67%	28
1646T	May-02	90,360	29,805	2	<0.01%	106	0.12%	36	0.04%	303	0.34%	124	0.14%	12	0.01%	42	0.05%	112	0.12%	7	<0.01%	21	0.02%	765	0.85%	102
1688T/TC	Jun-03	489,893	191,315	40	<0.01%	467	0.10%	554	0.11%	1329	0.27%	1425	0.29%	137	0.03%	213	0.04%	535	0.11%	43	<0.01%	161	0.03%	4904	1.00%	1328

Definitions of observations and complications can be found on [page 7](#).

Pacing Leads U.S. Malfunction Summary

MODELS	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
LPA1200M	57,116	1.50%	3	<0.01%	5	<0.01%	0	0.00%	2	<0.01%	29	0.05%	39	0.07%
2088TC	585,050	3.20%	43	<0.01%	570	0.10%	0	0.00%	27	<0.01%	995	0.17%	1635	0.28%
1999	46,528	3.50%	5	0.01%	43	0.09%	0	0.00%	6	0.01%	144	0.31%	198	0.43%
1944	16,578	4.90%	0	0.00%	8	0.05%	0	0.00%	1	<0.01%	23	0.14%	32	0.19%
1948	63,721	3.20%	10	0.02%	75	0.12%	0	0.00%	1	<0.01%	74	0.12%	160	0.25%
1699T/TC	22,878	4.60%	13	0.06%	31	0.14%	0	0.00%	0	0.00%	53	0.23%	97	0.42%
1888T/TC	301,676	4.00%	40	0.01%	760	0.25%	1	<0.01%	12	<0.01%	781	0.26%	1594	0.53%
1882T/TC	48,680	3.30%	2	<0.01%	57	0.12%	0	0.00%	3	<0.01%	124	0.25%	186	0.38%
1782T/TC	16,404	4.70%	1	<0.01%	34	0.21%	0	0.00%	0	0.00%	49	0.30%	84	0.51%
1788T/TC	65,235	5.00%	10	0.02%	110	0.17%	1	<0.01%	1	<0.01%	101	0.15%	223	0.34%
1648T	2,834	5.60%	0	0.00%	11	0.39%	0	0.00%	2	0.07%	5	0.18%	18	0.64%
1642T	27,127	4.60%	0	0.00%	21	0.08%	1	<0.01%	2	<0.01%	20	0.07%	44	0.16%
1646T	90,360	4.40%	21	0.02%	64	0.07%	0	0.00%	6	<0.01%	64	0.07%	155	0.17%
1688T/TC	489,893	4.40%	203	0.04%	849	0.17%	2	<0.01%	18	<0.01%	767	0.16%	1839	0.38%
1488T/TC	270,810	4.40%	158	0.06%	304	0.11%	5	<0.01%	3	<0.01%	360	0.13%	830	0.31%

Definitions of malfunction categories can be found on [pages 8-9](#).

Pacing Leads

Worldwide Malfunction Summary

MODELS	WORLDWIDE SALES	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
LPA1200M	333,295	0.4%	20	0.01%	29	0.01%	0	0.00%	11	<0.01%	110	0.03%	170	0.05%
2088TC	1,581,830	1.2%	61	<0.01%	682	0.04%	0	0.00%	66	<0.01%	1307	0.08%	2116	0.13%
1888T/TC	1,082,799	1.3%	62	0.01%	930	0.09%	1	<0.01%	31	<0.01%	1128	0.10%	2152	0.20%

Definitions of malfunction categories can be found on [pages 8-9](#).

Pacing Leads

Actively Monitored Study Data Summary

QUALIFYING COMPLICATIONS

MODELS	NUMBER OF DEVICES ENROLLED	ACTIVE DEVICES ENROLLED	CUMULATIVE MONTHS OF FOLLOW-UP	ABNORMAL PACING IMPEDANCE		CARDIAC PERFORATION		CONDUCTOR FRACTURE		EXTRACARDIAC STIMULATION		FAILURE TO CAPTURE		FAILURE TO SENSE		INSULATION BREACH		LEAD DISLODGE MENT		OVERSENSING		PERICARDIAL EFFUSION		SKIN EROSION		TOTAL	
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
2088	3,841	1,795	194,241	1	0.03%	1	0.03%	2	0.05%	1	0.03%	6	0.16%	2	0.05%	6	0.16%	15	0.39%	12	0.31%	1	0.03%	0	0.00%	47	1.22%
1999	869	404	41,920	1	0.12%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	2	0.23%	1	0.12%	10	1.15%	1	0.12%	0	0.00%	0	0.00%	16	1.84%
1944	104	31	5,742	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.96%	0	0.00%	0	0.00%	0	0.00%	1	0.96%
1948	765	197	33,414	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.13%	0	0.00%	1	0.13%	2	0.26%	0	0.00%	0	0.00%	0	0.00%	4	0.52%
1699T/TC	1,451	337	71,422	1	0.07%	0	0.00%	3	0.21%	0	0.00%	4	0.28%	0	0.00%	1	0.07%	8	0.55%	1	0.07%	0	0.00%	0	0.00%	18	1.24%
1888T/TC	14,503	4,601	822,216	6	0.04%	2	0.01%	8	0.06%	4	0.03%	19	0.13%	4	0.03%	26	0.18%	58	0.40%	22	0.15%	0	0.00%	1	<0.01%	150	1.03%
1882T/TC	690	205	38,104	0	0.00%	0	0.00%	0	0.00%	1	0.14%	1	0.14%	0	0.00%	0	0.00%	2	0.29%	2	0.29%	0	0.00%	1	0.14%	7	1.01%
1782T/TC	165	15	5,762	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.61%	0	0.00%	0	0.00%	1	0.61%
1788T/TC	363	41	12,360	0	0.00%	0	0.00%	0	0.00%	1	0.28%	0	0.00%	0	0.00%	0	0.00%	3	0.83%	0	0.00%	0	0.00%	0	0.00%	4	1.10%
1646T	641	3	15,791	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.31%	0	0.00%	0	0.00%	1	0.16%	0	0.00%	0	0.00%	0	0.00%	3	0.47%
1688T/TC	2,645	417	92,643	5	0.19%	0	0.00%	2	0.08%	0	0.00%	3	0.11%	0	0.00%	3	0.11%	6	0.23%	3	0.11%	1	0.04%	1	0.04%	24	0.91%
1488T/TC	803	50	26,898	0	0.00%	0	0.00%	0	0.00%	0	0.00%	1	0.12%	0	0.00%	2	0.25%	0	0.00%	1	0.12%	0	0.00%	0	0.00%	4	0.50%

A list of complications can be found on [page 12](#).

Pacing Leads

Actively Monitored Study Data Summary

MALFUNCTIONS

MODELS	NUMBER OF DEVICES ENROLLED	PERCENT RETURNED FOR ANALYSIS	CONDUCTOR FRACTURE		INSULATION BREACH		CRIMPS, WELDS & BONDS		OTHER		EXTRINSIC FACTORS		TOTAL	
			QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
2088TC	3,841	4.20%	1	0.03%	12	0.31%	0	0.00%	0	0.00%	14	0.36%	27	0.70%
1999	869	6.00%	0	0.00%	5	0.58%	0	0.00%	0	0.00%	8	0.92%	13	1.50%
1944	104	1.90%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
1948	765	4.70%	0	0.00%	4	0.52%	0	0.00%	0	0.00%	1	0.13%	5	0.65%
1699T/TC	1,451	3.10%	0	0.00%	2	0.14%	0	0.00%	0	0.00%	6	0.41%	8	0.55%
1888T/TC	14,503	3.60%	3	0.02%	24	0.17%	0	0.00%	0	0.00%	35	0.24%	62	0.43%
1882T/TC	690	3.90%	0	0.00%	2	0.29%	0	0.00%	0	0.00%	0	0.00%	2	0.29%
1782T/TC	165	3.00%	0	0.00%	1	0.61%	0	0.00%	0	0.00%	0	0.00%	1	0.61%
1788T/TC	363	2.80%	0	0.00%	1	0.28%	0	0.00%	0	0.00%	0	0.00%	1	0.28%
1646T	641	1.60%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
1688T/TC	2,645	5.30%	0	0.00%	4	0.15%	0	0.00%	0	0.00%	5	0.19%	9	0.34%
1488T/TC	803	3.50%	0	0.00%	4	0.50%	0	0.00%	0	0.00%	1	0.12%	5	0.62%

Definitions of malfunction categories can be found on [pages 8-9](#).

Implantable Cardiac Monitors (ICM) Devices

Implantable Cardiac Monitors (ICMs) Devices

CUSTOMER REPORTED PERFORMANCE DATA

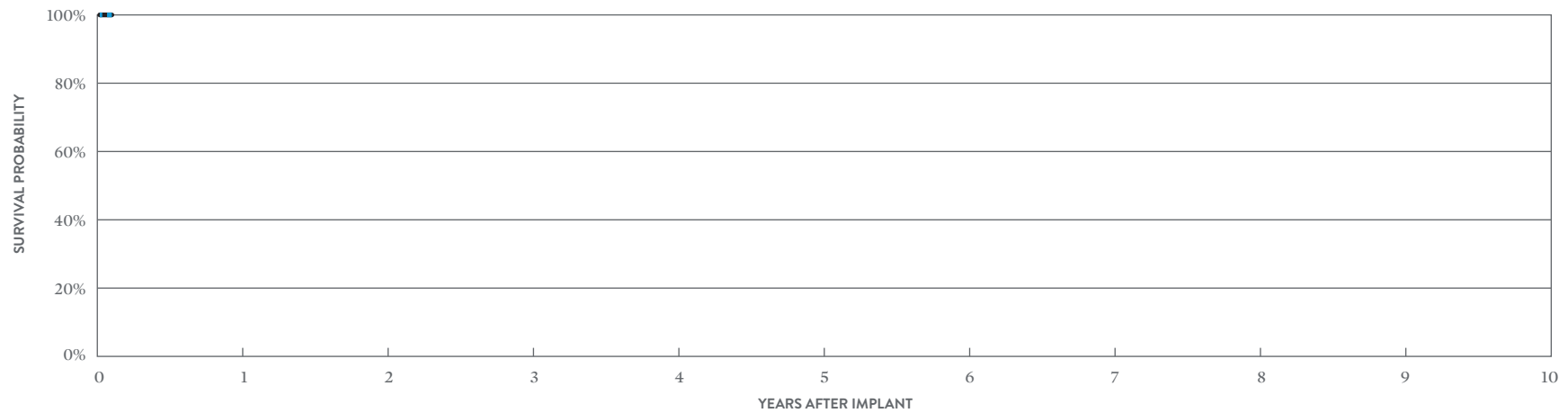
Confirm Rx™ ICM

MODEL DM3500

US Regulatory Approval	September 2017
Registered US Implants	968
Estimated Active US Implants	952
Estimated Longevity	2 Years
Normal Battery Depletion	0
Number of US Advisories	None

MALFUNCTIONS

	QTY	RATE
Electrical Component	0	0.00%
Electrical Interconnect	0	0.00%
Battery	0	0.00%
Software/Firmware	0	0.00%
Mechanical	0	0.00%
Possible Early Battery Depletion	0	0.00%
Other	0	0.00%
Total	0	0.00%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	AT 1 MONTH
SURVIVAL PROBABILITY	100.00%
± 1 STANDARD ERROR	0.00%
SAMPLE SIZE	970

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	AT 1 MONTH
SURVIVAL PROBABILITY	100.00%
± 1 STANDARD ERROR	0.00%

Implantable Cardiac Monitors (ICMs) Devices

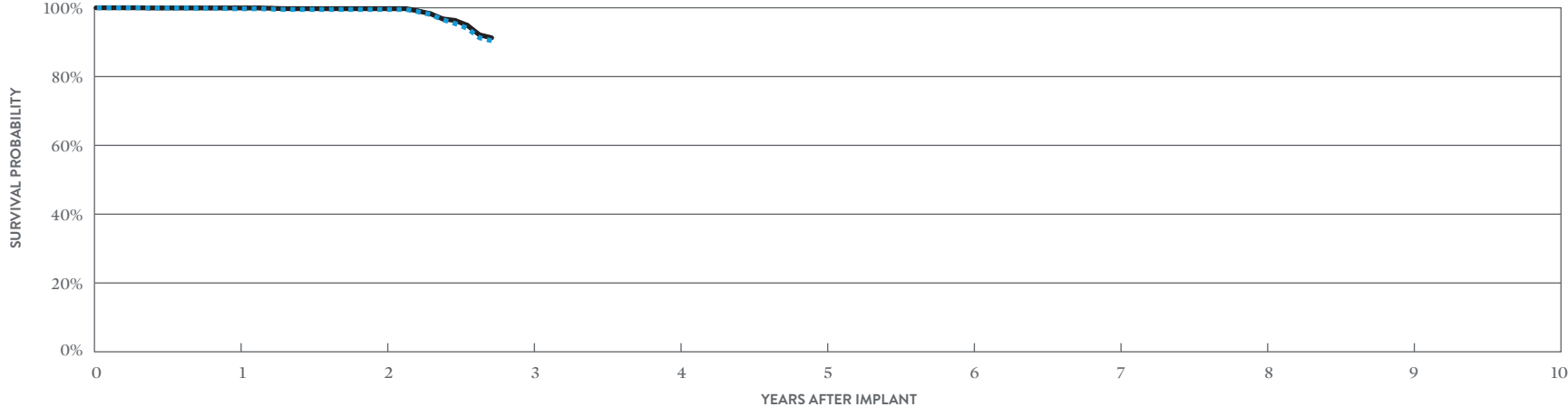
CUSTOMER REPORTED PERFORMANCE DATA

SJM Confirm™ ICM

MODEL DM2102

US Regulatory Approval	May 2014
Registered US Implants	4,282
Estimated Active US Implants	3,226
Estimated Longevity	3 Years
Normal Battery Depletion	11
Number of US Advisories (see pg. 345)	One

MALFUNCTIONS		
	QTY	RATE
Electrical Component	19	0.44%
Electrical Interconnect	0	0.00%
Battery	0	0.00%
Software/Firmware	0	0.00%
Mechanical	0	0.00%
Possible Early Battery Depletion	0	0.00%
Other	2	0.05%
Total	21	0.49%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 33 MONTHS
SURVIVAL PROBABILITY	99.71%	99.50%	90.42%
± 1 STANDARD ERROR	0.09%	0.15%	1.40%
SAMPLE SIZE	3,310	1,650	240

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	AT 33 MONTHS
SURVIVAL PROBABILITY	99.94%	99.74%	91.26%
± 1 STANDARD ERROR	0.04%	0.11%	1.36%

Implantable Cardiac Monitors (ICMs) Devices

CUSTOMER REPORTED PERFORMANCE DATA

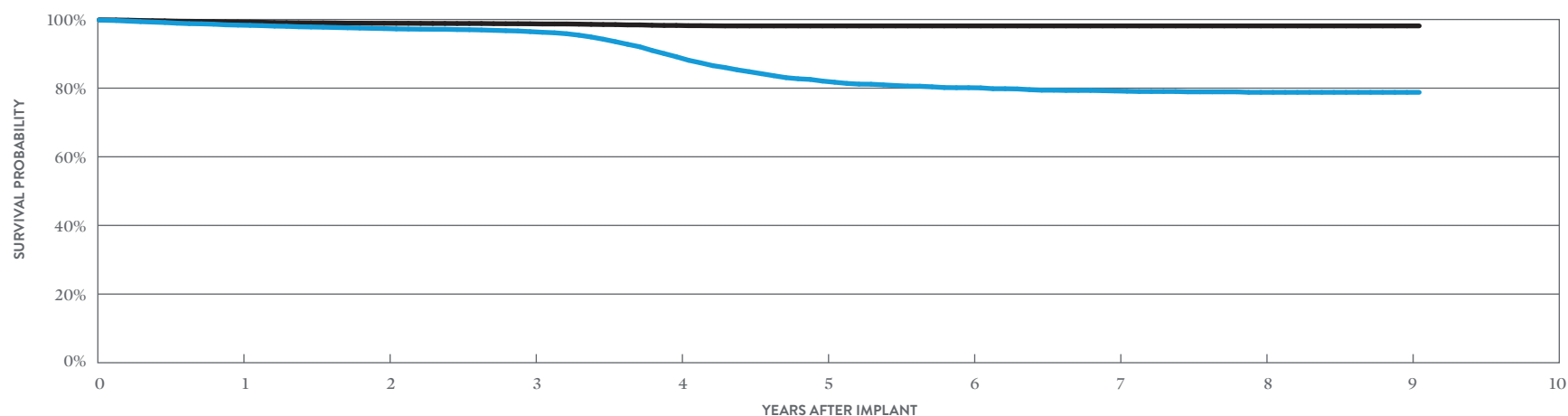
SJM Confirm™ ICM

MODEL DM2100

US Regulatory Approval	August 2008
Registered US Implants	18,686
Estimated Active US Implants	7,334
Estimated Longevity	3 Years
Normal Battery Depletion	645
Number of US Advisories (see pg. 345)	One

MALFUNCTIONS

	QTY	RATE
Electrical Component	15	0.08%
Electrical Interconnect	1	<0.01%
Battery	20	0.11%
Software/Firmware	10	0.05%
Mechanical	0	0.00%
Possible Early Battery Depletion	8	0.04%
Other	41	0.22%
Total	95	0.51%



INCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 109 MONTHS
SURVIVAL PROBABILITY	98.37%	97.36%	96.47%	89.15%	82.06%	80.13%	79.18%	78.78%	78.78%	78.78%
± 1 STANDARD ERROR	0.10%	0.13%	0.15%	0.29%	0.40%	0.44%	0.47%	0.49%	0.49%	0.49%
SAMPLE SIZE	16,340	12,890	10,940	8,820	6,290	4,190	2,680	1,530	630	220

EXCLUDING NORMAL BATTERY DEPLETION

YEAR	1	2	3	4	5	6	7	8	9	AT 109 MONTHS
SURVIVAL PROBABILITY	99.30%	98.90%	98.75%	98.27%	98.14%	98.14%	98.14%	98.14%	98.14%	98.14%
± 1 STANDARD ERROR	0.06%	0.09%	0.09%	0.12%	0.13%	0.13%	0.13%	0.13%	0.13%	0.13%

SUMMARY INFORMATION
**Implantable Cardiac
Monitors (ICMS)**

Implantable Cardiac Monitors (ICMs) Devices

Survival Probability Summary

INCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
DM3500	Confirm Rx™ ICM*										
DM2102	SJM Confirm™ ICM	99.71%	99.50%								
DM2100	SJM Confirm™ ICM	98.37%	97.36%	96.47%	89.15%	82.06%	80.13%	79.18%	78.78%	78.78%	

EXCLUDING NORMAL BATTERY DEPLETION

MODELS	FAMILY	1 YEAR	2 YEAR	3 YEAR	4 YEAR	5 YEAR	6 YEAR	7 YEAR	8 YEAR	9 YEAR	10 YEAR
DM3500	Confirm Rx™ ICM*										
DM2102	SJM Confirm™ ICM	99.94%	99.74%								
DM2100	SJM Confirm™ ICM	99.30%	98.90%	98.75%	98.27%	98.14%	98.14%	98.14%	98.14%	98.14%	

*No survival probability is stated at one year due to the device not meeting the required minimum sample size of 200 U.S. implants with 12 consecutive months of data. Please refer to the individual graphs for data up to one year.

Implantable Cardiac Monitors (ICMs) Devices

US Malfunction Summary

MODELS	FAMILY	REGISTERED US IMPLANTS	PERCENT RETURNED FOR ANALYSIS	ELECTRICAL COMPONENT		ELECTRICAL INTERCONNECT		BATTERY		SOFTWARE/ FIRMWARE		MECHANICAL		POSSIBLE EARLY BATTERY DEPLETION		OTHER		TOTAL			
				QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE	QTY	RATE
DM3500	Confirm Rx™ ICM*	968	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%
DM2102	SJM Confirm™ ICM	4,282	4.30%	19	0.44%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	0	0.00%	2	0.05%	21	0.49%	21	0.49%
DM2100	SJM Confirm™ ICM	18,686	16.30%	15	0.08%	1	<0.01%	20	0.11%	10	0.05%	0	0.00%	8	0.04%	41	0.22%	95	0.51%	95	0.51%

Definitions of malfunction categories can be found on [pages 5-6](#).

Focus on Clinical Performance

ICD Premature Battery Depletion Advisory Update – April 2018

Since the original October 11, 2016 communication Abbott and our Medical Advisory Board have continued to analyze and review performance data from the affected device population. The rates reported below summarize performance data through Feb 28, 2018.

Importantly, the information contained in this notice has not altered our previously communicated patient management recommendations. This information is designed to keep you informed of ongoing analysis of products returned to the company and does not change what we have previously communicated around this issue or how you have approached the management of your patients impacted by our field advisory.

RATES

The table below summarizes the updated worldwide experience for the affected devices that were returned for product analysis due to premature battery depletion. We have included both confirmed and unconfirmed shorts in the rate table below to help you assess the risk to your patients. The table includes both the updated data and data from the original communication.

UPDATED (THROUGH FEBRUARY 28, 2018)

WORLDWIDE PATIENT IMPACT	NUMBER / RATE ORIGINAL OCTOBER 11, 2016	NUMBER / RATE THROUGH NOVEMBER 30, 2017	NUMBER / RATE THROUGH FEBRUARY 28, 2018
No Impact Reported/Additional Surgery Only*	792/0.20%	1332/0.33%	1461/0.37%
Loss of Pacing – Minor (Dizziness)	37/<0.01%	49/0.01%	50/0.01%
Loss of Pacing – Major (Syncope)	10/<0.01%	13/0.01%	13/<0.01%
Loss of Defibrillation – Emergency	0/0%	2/0.01%	3/<0.01%
Loss of Defibrillation – Death	2/<0.01%	2/0.01%	2/<0.01%
Grand Total	841/0.21%	1398/0.35%	1529/0.38%
Total Units Sold	398,740		

**All impacts in this table were related to a replacement surgery, as the data is from units explanted and returned for analysis. The category “ No Impact Reported/Additional Surgery Only” means there was no associated report of patient symptoms in addition to the replacement of the unit with a depleted battery.*

Elective Replacement Interval (ERI) and End of Life (EOL) Analysis Update

WORLDWIDE ERI TO EOL IMPACT TABLE:

The following table represents the analysis results of the time interval between ERI and EOL for devices returned to Abbott for analysis where premature battery depletion was determined, no electronic or other assignable cause was found, battery testing found lithium clusters and the battery voltage was sufficient to allow retrieval of device diagnostics data. Of the 1529 units returned to Abbott as of the date of analysis, 820 units met the above criteria.

ERI TO EOL DURATION (FOR RETURNED UNITS WITH LITHIUM CLUSTER PBD AND DEVICE RETRIEVABLE DATA)**	NUMBER OF UNITS
ERI detected, patient alert delivered	806/98.29%
<= 1 day; patient notifier alert was triggered	135
>1 and <= 10 days patient notifier alert was triggered	149
>10 and <= 30 days patient notifier alert was triggered	90
>30 days; patient notifier alert was triggered	55
Above EOL, therefore ERI to EOL duration not applicable; patient notifier alert was triggered if ERI was reached	377
ERI not detected, patient alert was not delivered, but below ERI threshold of 2.59V	14/1.71%
Total Number of Units	820
Total Units Sold	398,740

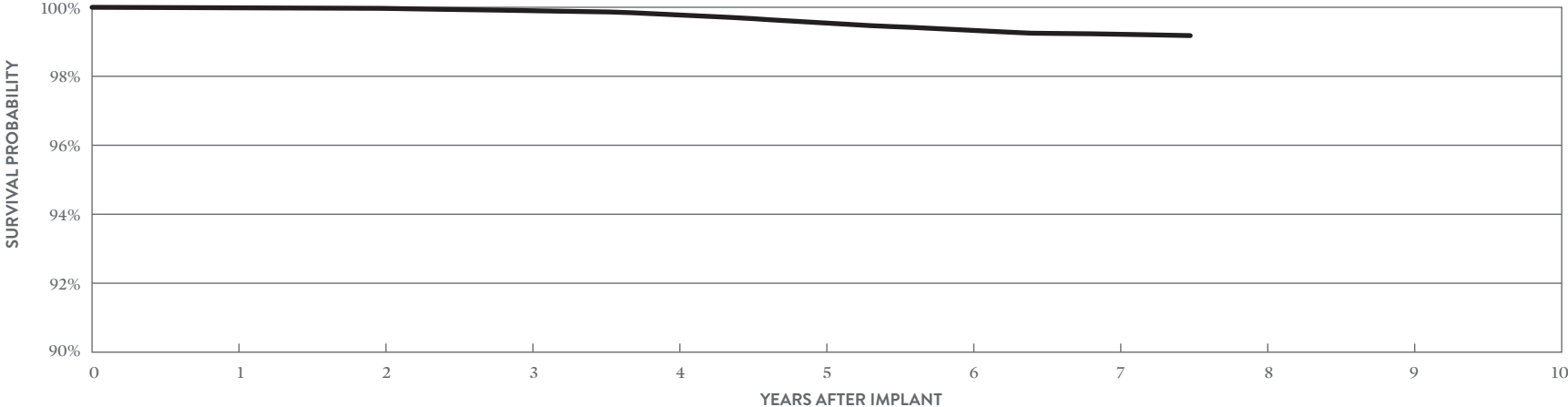
**Our intent is to provide these data to help explain the statement “battery depletion can occur in these devices within a day to a few weeks, which may result in the inability to deliver therapy” in the original field advisory notification.

These data are provided for information only and should not be used to make specific patient management decisions that would depart from recommendations previously communicated by the company. The patient management recommendations in the original advisory letter remain applicable.

The number of units in each of the groups above is not a predictor of future performance of a specific unit currently implanted.

Estimated Performance of Affected Fortify™, Fortify Assura™, Quadra Assura™, Unify™, Unify Assura™ and Unify Quadra™ Devices

SEVEN-YEAR COMBINED KAPLAN-MEIER (KM) SURVIVAL CURVE OF FREEDOM FROM PREMATURE BATTERY DEPLETION ASSOCIATED WITH LI DEPOSITS IN AFFECTED DEVICE POPULATION.



YEAR	1	2	3	4	5	6	7	AT 89 MONTHS
SURVIVAL PROBABILITY	99.999%	99.977%	99.872%	99.702%	99.509%	99.240%	99.016%	98.969%
SAMPLE SIZE	-224,000	-208,000	-173,000	-126,000	-85,000	-49,000	-15,000	-4,900

SURVIVAL CALCULATION GENERAL METHODS

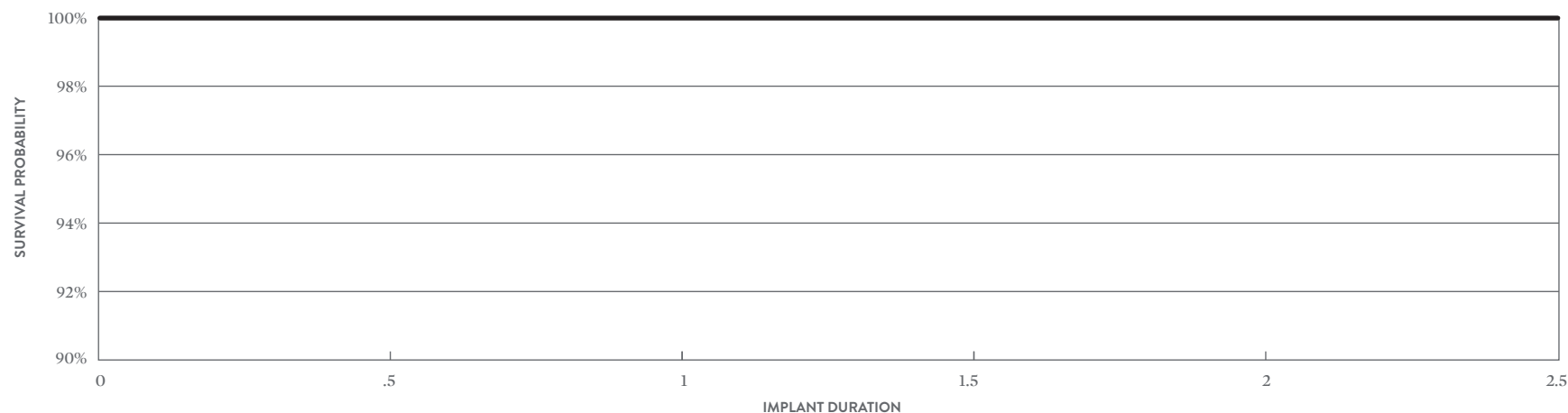
Internal modeling based on an analysis of field returns related to premature battery depletions associated with lithium clusters. Updated with data through February 2018.

Non-Advisory Population Update

Fortify™, Fortify Assura™, Quadra Assura™, Quadra Assura MP™, Unify™, Unify Assura™ and Unify Quadra™ Devices manufactured after May 23, 2015 were built with an improved battery design with additional insulation and thus not included in the advisory population. Through February 2018 there have been zero (N=0) occurrences, worldwide, of premature depletion due to Li clusters with the improved design.

In the US there have been ~ 100,000 implanted devices with the improved battery design with no occurrences of premature depletion associated with Li clusters. Of the implanted US population, ~60% (or ~62,000) have exceed 1 year of implant duration and ~17% (or ~17,000) have exceed 2.0 years of implant duration with no occurrences of premature depletion due to Li clusters.

**SURVIVAL PLOT FOR NON-ADVISORY POPULATION
KAPLAN-MEIER METHOD
CENSORING FOR NON-ADVISORY POPULATION WITH CLUSTER**



UNIFY/FORTIFY/ASSURA (NON-ADVISORY POPULATION)

YEAR	.5	1	1.5	2	2.5
SURVIVAL PROBABILITY	100%	100%	100%	100%	100%
SAMPLE SIZE	-82,000	-62,000	-35,000	-17,000	-2,500

Update on Riata™ Lead Performance

REGISTRY AND POST-MARKET STUDIES

Prospective, monitored registries continue to provide the best data to support clinical decision making. Abbott initiated the Riata Lead Evaluation Study (RLES), which began in December 2011 and has enrolled 782 patients with Riata leads at sites in U.S., Canada and Japan. Phase I of the study involved enrollment, collection of patient information, thorough (3 angle) cinefluoroscopic imaging, and adjudication of cinefluoroscopic data by experienced, independent electrophysiologists for the presence of externalized conductors. U.S., Canadian, and Japanese data from Phase I were first reported by Abbott in the November 2012 Product Performance Report 2nd Edition. RLES data was subsequently presented at the 2013 and 2014 Heart Rhythm Society Scientific Sessions and detailed in a peer-reviewed manuscript.^{1,2,3}

In 2013, Abbott expanded the RLES to include Durata™ and Quicksite™/Quickflex™ leads and to increase the quantity of Riata™ and Riata™ ST leads. The expanded study, known as the “St. Jude Medical Cardiac Lead Assessment Study” (CLAS), began enrollment in February 2013 to ensure at least 500 leads in each of the following lead families: Riata, Riata ST, Durata and Quicksite/Quickflex. Under the new CLAS protocol, patients will be followed every six months for three years and cinefluoroscopy will be done at the enrollment, 1-year, 2-year and 3-year follow-up visits. The main objective of the study is to determine the prevalence and incidence of lead compromise evidenced by imaging (includes externalized conductors and other visual lead anomalies) and electrical dysfunction in Riata, Riata ST, Quicksite/Quickflex and Durata leads. All images were adjudicated by an independent panel of experienced electrophysiologists for determining the presence of externalized conductors. Additionally, upon occurrence of a lead revision during follow-up, another physician panel determined whether electrical dysfunction had occurred based upon predefined criteria.⁴ Enrollment is now complete for CLAS. The following summaries for Riata/Riata ST and QuickSite/QuickFlex leads represent all data collected as of February 28, 2018. The Durata leads CLAS summary is available on page 317.

1 David Hayes, Roger Freedman, Mark Niebauer, Vance Plumb, Jay Dinerman, Scott Beau, Anne Curtis, *Incidence of New Externalized Conductors and Electrical Dysfunction in Riata and Riata ST Silicone ICD Leads: 1 year Results from a Prospective, Multicenter Study*, Heart Rhythm Society's Annual Scientific Sessions, San Francisco, CA, May 8, 2014.

2 David Hayes, Roger Freedman, Anne Curtis, Mark Niebauer, G.Neal Kay, Jay Dinerman, Scott Beau, Wilson Wong, *Prevalence of Externalized Conductors in Riata and Riata ST Silicone ICD Leads: Results from a Prospective, Multicenter Study*, Heart Rhythm Society's Annual Scientific Sessions, Denver, CO, May 9, 2013.

3 David Hayes, Roger Freedman, Anne B. Curtis, Mark Niebauer, G. Neal Kay, Jay Dinerman, Scott Beau, *Prevalence of Externalized Conductors in Riata and Riata ST Silicone Leads: Results from the Prospective, Multicenter, Riata Lead Evaluation Study*, *Heart Rhythm*, Vol. 10, Issue 12, Pages 1778-1782, December 2013.

4 Electrical dysfunction is defined as a lead that exhibits at least one of the following criteria: 1) Presence of non-physiologic noise not due to external interference. 2) Rise in pace/sense (p/s) conductor impedance to > 2000 Ω or increase of more than 200 Ω over previous 6 months or increase of 400 Ω over any period of time. 3) Decrease of more than 200 Ω over previous 6 months or to impedance < 200 Ω from baseline impedance > 300 Ω or decrease of 400 Ω over any period of time. 4) Change in any high voltage coil impedance of > 25 Ω or to > 125 Ω or < 20 Ω. 5) A capture threshold > 5 V or an increase of > 2 V from baseline (all measurements) of < 1 V.

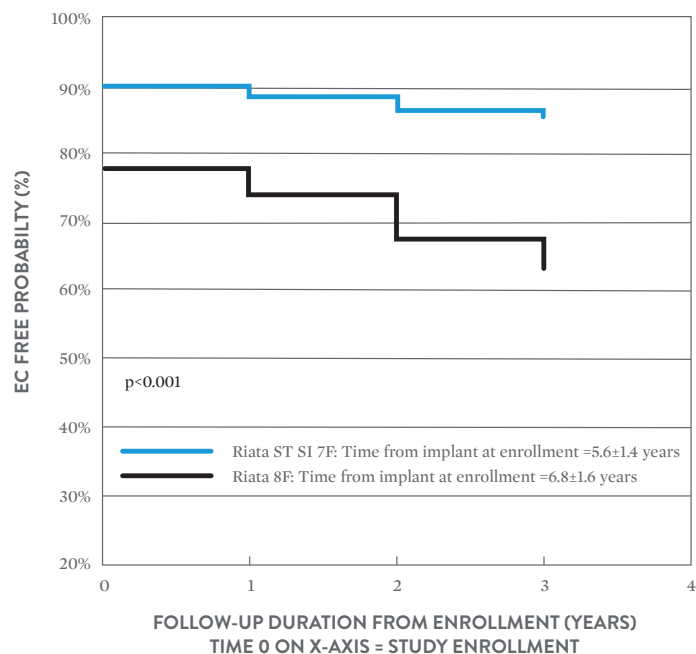
Focus on Clinical Performance

RIATA™/RIATA™ ST LEAD CLAS SUMMARY (AS OF FEBRUARY 28, 2018): This summary includes all Riata/Riata ST silicone leads that were enrolled in the initial RLES study as well as those enrolled in the subsequent CLAS study. A total of 1,128 patients with Riata/Riata ST silicone leads across 42 centers (8F/7F= 59.4%/40.6%) underwent fluoroscopic evaluation. The implant duration at enrollment was 6.8 ± 1.6 years for 8F Riata leads and 5.6 ± 1.4 years for 7F Riata ST leads. The prevalence of externalized conductors (EC) at enrollment was significantly lower in 7F Riata ST leads compared to 8F Riata leads (9.7% vs. 21.9%, $p < 0.0001$). A total of 754 patients (66.8%) completed at least 1 year of follow-up, 626 patients (55.5%) completed at least 2 years of follow up, and 466 (41.3%) completed at least three years of follow up with fluoroscopic evaluations. The event-free survival rate for externalized conductors in Riata and Riata ST leads through 3 years of follow-up is shown in Figure 1. At 3 years of follow up, the freedom from externalized conductors is 86.2% in Riata ST 7F leads and 63.5% in Riata 8F leads. In 1,128 Riata and Riata ST leads evaluated, 96.4% have been free from electrical dysfunction. Of the 39 leads (15 Riata ST 7F and 24 Riata 8F) exhibiting electrical dysfunction, 15 leads (5 Riata ST 7F and 10 Riata 8F) had externalized conductors. The electrical failure rate in leads with and without EC is statistically significant ($p = 0.0297$) as shown in Table 1.

All pending fluoroscopy data has been adjudicated and the minimum enrollment of Riata/Riata ST silicone leads has been met in the Cardiac Lead Assessment Study.

Focus on Clinical Performance

**FIGURE 1: KAPLAN-MEIER ESTIMATE OF FREEDOM FROM EXTERNALIZED CONDUCTORS (EC)
RIATA (8F) VS. RIATA ST SI (7F) LEADS**



RIATA ST (7F)

YEAR	0	1	2	3
AT RISK	452	314	247	181
CUMULATIVE EC EVENTS	44	49	55	56
EC FREE PROBABILITY	90.3%	88.8%	86.7%	86.2%

RIATA ST (8F)

YEAR	0	1	2	3
AT RISK	659	374	283	189
CUMULATIVE EC EVENTS	144	162	187	199
EC FREE PROBABILITY	78.1%	74.4%	67.8%	63.5%

Focus on Clinical Performance

TABLE 1: RIATA/ RIATA ST LEADS: CORRELATION BETWEEN EC AND ED

	PROPORTION OF LEADS WITH ELECTRICAL DYSFUNCTION (ED), %	P-VALUE*
With EC	5.9% (15/255) ¹	0.0297
Without EC	2.7% (24/873) ²	

*p-value was calculated using Fisher's exact test.

¹Denominator = Total # of leads with EC

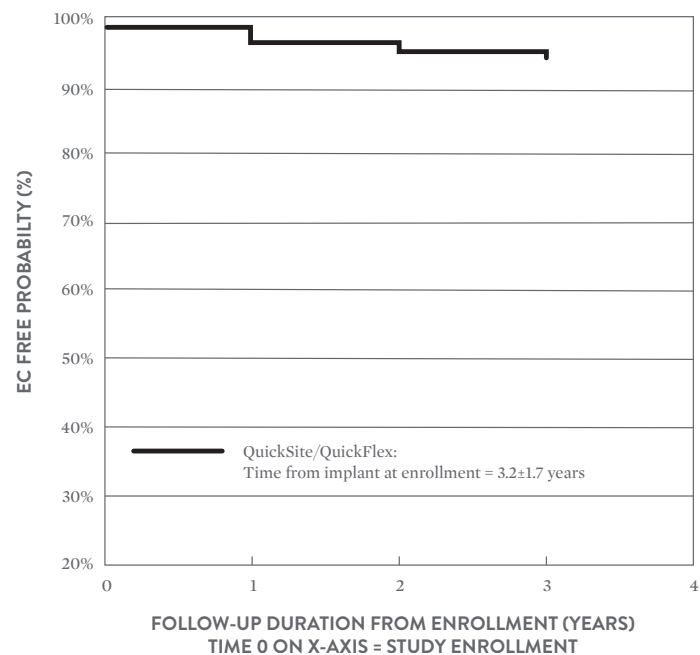
²Denominator = Total # of leads without EC

QUICKSITE™/QUICKFLEX™ LEAD CLAS SUMMARY (FEBRUARY 28, 2018): A total of 736 patients implanted with 800 QuickSite/QuickFlex Left Ventricular CRT leads at 43 centers underwent fluoroscopic evaluation. The implant duration at enrollment was 3.2 ± 1.7 years. A total of 585 patients (73.1%) completed at least 1 year of follow-up, 456 patients (57.0%) completed at least 2 years of follow up, and 302 (37.8%) completed at least three years of follow up with fluoroscopic evaluations. The event-free survival rate for externalized conductors through 3 years of follow-up is shown in Figure 2. The prevalence of externalized conductors (EC) was 1.7% (13/784) at enrollment. At 3 years of follow up, the freedom from externalized conductors is 94.0%. In 800 QuickSite/QuickFlex leads evaluated, 99.7% have been free from electrical dysfunction. Neither of the 2 leads with electrical dysfunction exhibited externalized conductors as shown in Table 2.

In QuickSite/QuickFlex leads the presence of externalized conductors was not significantly associated with an increased risk of electrical dysfunction. All pending fluoroscopy data has been adjudicated and the minimum enrollment of the QuickSite/QuickFlex leads has been met in the Cardiac Lead Assessment Study.

Focus on Clinical Performance

FIGURE 2: EVENT FREE SURVIVAL RATE FOR QUICKSITE/QUICKFLEX LEADS



YEAR	0	1	2	3
AT RISK	784	611	448	282
CUMULATIVE EC EVENTS	13	29	34	36
EC FREE PROBABILITY	98.3%	95.8%	94.7%	94.0%

TABLE 2: QUICKSITE/QUICKFLEX LEADS: CORRELATION BETWEEN EC AND ED

	PROPORTION OF LEADS WITH ELECTRICAL DYSFUNCTION (ED), %	P-VALUE*
With EC	0% (0/32) ¹	1.0000
Without EC	0.3% (2/768) ²	

*p-value was calculated using Fisher's exact test.

¹Denominator = Total # of leads with EC

²Denominator = Total # of leads without EC

Focus on Clinical Performance

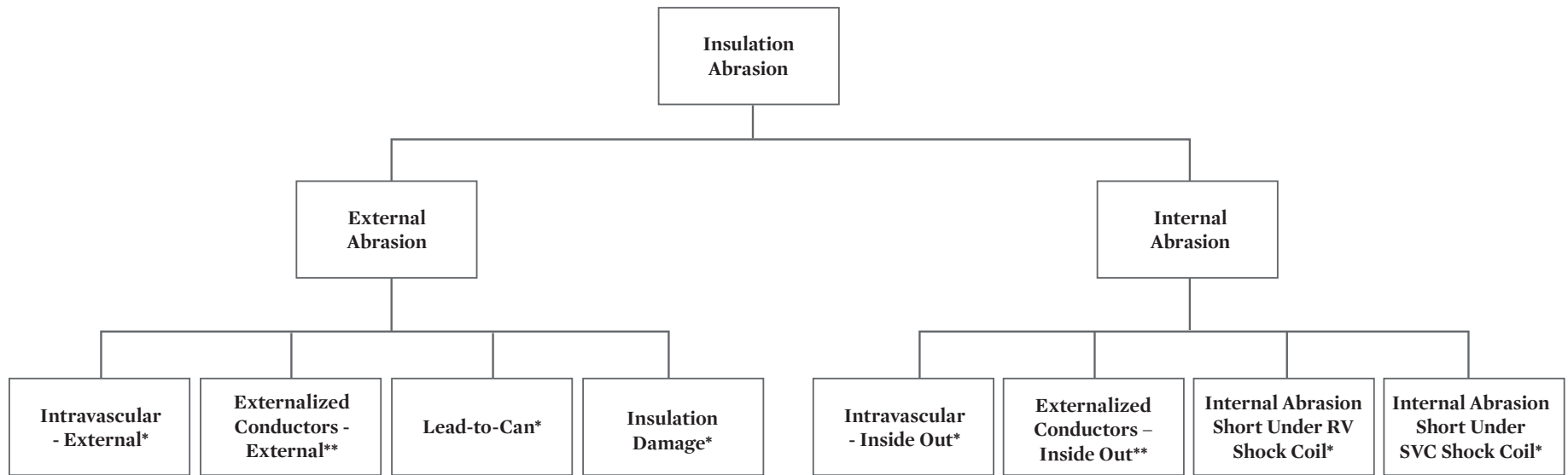
CUSTOMER REPORTED PERFORMANCE DATA

Abbott understands that the passive system of complaint reporting and returned product analysis results in under-reporting and hence underestimates the true failure rate associated with any given failure mechanism. This is especially true for externalized conductors since most manifest as visual anomalies only with normal electrical performance. While acknowledging these limitations, Abbott provides externalized conductor rates from the passive data system to maintain continuity with previously published data and to provide full disclosure of the data available to Abbott. As of February 28, 2018, there were 6,047 cases of externalized conductors reported to Abbott worldwide on Riata™ (8F) and Riata™ ST (7F) silicone defibrillation leads, equating to a 3.26% (5,090/156,000) incidence rate for Riata (8F) and 1.36% (957/70,600) for Riata ST (7F) leads. Of these 6,047 leads, 4,456 were not returned and 1,591 were returned for analysis.

As with any lead, there are failure mechanisms other than externalized conductors which result from insulation abrasion. Historically, the rate of all-cause insulation abrasion failures has been reported in the range of 3 to 10% (Kleemann et al., Annual Rate of Transvenous Defibrillation Lead Defects in Implantable Cardioverter Defibrillators over a Period > 10 years, *Circulation* 2007; 115:2474-2480). The most common form of insulation abrasion has been lead-to-can abrasion occurring in the pocket area. Externalization of conductors is another manifestation of insulation abrasion. It is most commonly caused by a mechanism referred to as inside-out abrasion, where the conductor cables become visible outside the insulation body. Approximately 87% of confirmed externalized conductors from product returns analysis are caused by inside-out abrasion, while 13% result from external sources of abrasion.

A flow diagram depicting specific insulation abrasion failure mechanisms for Riata™ and Riata™ ST silicone leads is shown in the following figure.

FLOW DIAGRAM OF INSULATION ABRASION TYPES AND FAILURE MECHANISMS



*Determined by returned product analysis.

**Includes cases determined by returned product analysis as well as cases identified only by fluoroscopy or visualization of explanted leads.

Focus on Clinical Performance

Definitions of the failure mechanisms are provided below:

- **External Abrasion:** Abrasion resulting from direct contact with an implanted device (e.g., pulse generator can, another lead), calcified anatomy, or anatomical structure that results in an outer insulation breach.
- **Internal Abrasion:** “Inside-out” abrasion between a lead conductor and the outer insulation that results in an insulation breach.
- **Intravascular Abrasion - External:** Direct contact with a foreign body (e.g., another lead), calcified anatomy, or cardiac structure within the vascular system or the heart that results in an outer insulation breach. The nature of the breach does not result in the conductors becoming visible outside the lead body. Determined by returned product analysis.
- **Externalized Conductors – External Source of Abrasion:** Direct contact with a foreign body (e.g., another lead), calcified anatomy, or cardiac structure within the vascular system or the heart that results in an outer insulation breach and the normally contained conductors becoming visible outside the lead body. Includes cases determined by returned product analysis as well as cases identified only by fluoroscopy or visualization of explanted leads. For those cases not returned, the assignment of an external source of abrasion is based on the fraction of externalized conductor cases identified by returned product analysis as external.
- **Lead-to-Can Abrasion:** Direct contact between the lead and the can (i.e. pacemaker, ICD, or CRT-D) that results in an outer insulation breach. Determined by returned product analysis.
- **Insulation Damage:** Insulation breaches that result from external mechanisms, including clavicular crush and outside-in abrasion by lead conductors. Determined by returned product analysis.
- **Intravascular Abrasion – Inside Out:** “Inside-out” abrasion between a lead conductor and the outer insulation within the vascular system or the heart that results in an outer insulation breach. The nature of the breach does not result in the conductors becoming visible outside the lead body. Determined by returned product analysis.
- **Externalized Conductors – Inside-Out:** Outward abrasion of conductors that results in an outer insulation breach within the vascular system or heart and the normally contained conductors becoming visible outside the lead body. Includes cases determined by returned product analysis as well as cases identified only by fluoroscopy or visualization of explanted leads. For those cases not returned, the assignment of an inside-out source of abrasion is based on the fraction of externalized conductor cases identified by returned product analysis as inside-out.
- **Internal Abrasion Short under RV Shock Coil:** Outward abrasion of the conductor cables under the RV shock coil that results in breaches of the outermost silicone insulation and the ETFE cable insulator, allowing the exposed metal surface of the conductor cables to make direct contact with, and potentially short against, the overlying RV shock coil. Determined by returned product analysis.
- **Internal Abrasion Short under SVC Shock Coil:** Outward abrasion of the conductor cables under the SVC shock coil that results in breaches of the outermost silicone insulation and the ETFE cable insulator, allowing the exposed metal surface of the conductor cables to make direct contact with, and potentially short against, the overlying SVC shock coil. Determined by returned product analysis.

Focus on Clinical Performance

The table below summarizes the incidence of insulation abrasion failure mechanisms confirmed by returns analysis of Riata™ and Riata™ ST leads. Approximately 13,860 Riata and Riata ST leads have been returned for analysis worldwide through February 28, 2018. Returned leads may exhibit more than one failure mechanism; hence the incidence rates presented in the table are not mutually exclusive. Note that the rates for externalized conductors also include visual-only observations that have been reported for leads remaining implanted.

RIATA™ (8F) AND RIATA™ ST (7F) LEAD INSULATION ABRASION FAILURE MECHANISMS FROM COMPLAINTS AND RETURNS

INSULATION FAILURE MECHANISM	ABRASION TYPE	RIATA (8F)	RIATA ST (7F)
		WORLDWIDE (WW) INCIDENCE RATE (WW SALES = 156,100)	WORLDWIDE (WW) INCIDENCE RATE (WW SALES = 70,600)
Intravascular – External*	External Abrasion	0.50%	0.52%
Externalized Conductors – External**	External Abrasion	0.42%	0.21%
Lead-to-Can*	External Abrasion	0.96%	0.89%
Insulation Damage*	External Abrasion	0.11%	0.06%
Intravascular - Inside Out*	Internal Abrasion	0.57%	0.39%
Externalized Conductors - Inside Out**	Internal Abrasion	2.86%	1.15%
Internal Abrasion Short Under RV Shock Coil*	Internal Abrasion	0.12%	0.05%
Internal Abrasion Short Under SVC Shock Coil*	Internal Abrasion	0.12%	0.02%

*Determined by returned product analysis.

**Includes cases determined by returned product analysis as well as cases identified only by fluoroscopy or visualization of explanted leads.

Update on Durata™ Lead Performance

REGISTRY AND POST-MARKET STUDIES

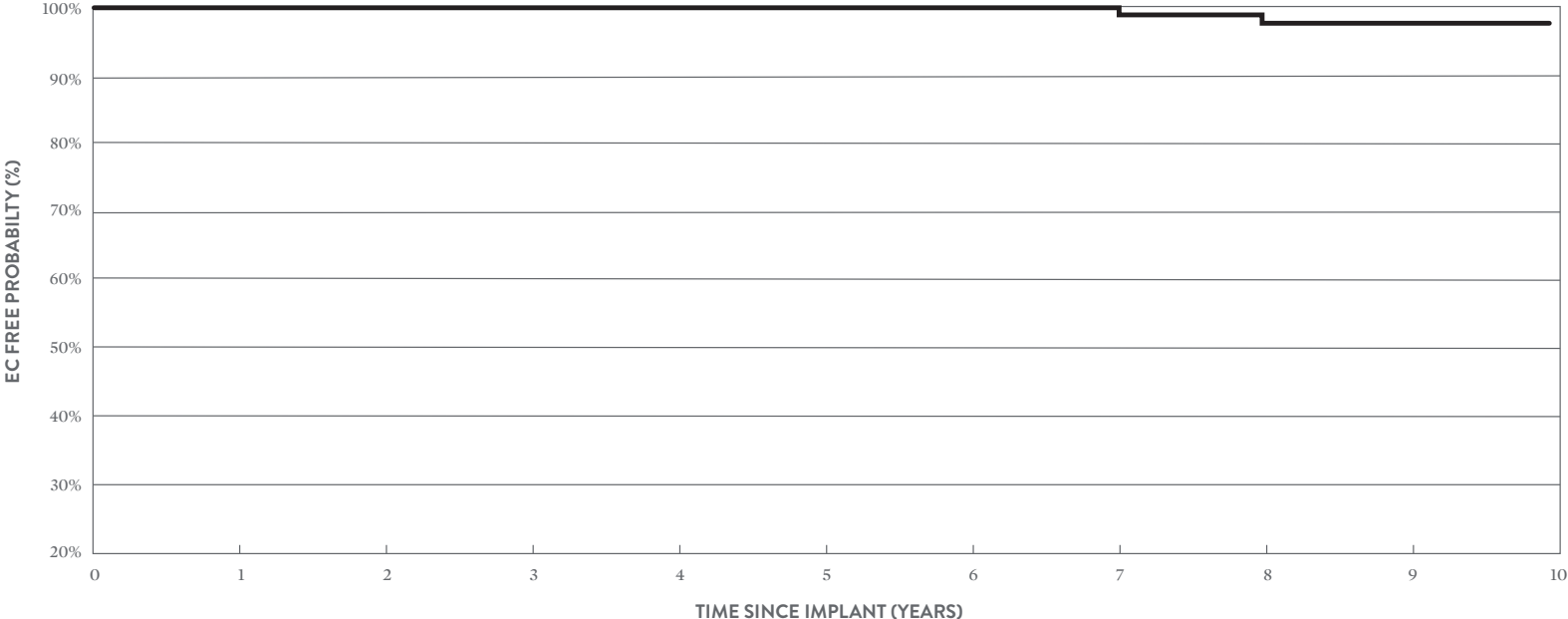
The safety and reliability of our Durata™ high voltage leads are supported by robust post-market surveillance monitoring. Data are collected on case report forms at each scheduled and unscheduled patient visit with additional information documented for any adverse event. We also employ a dedicated field monitoring organization to ensure that the data from the clinical site are accurately and completely submitted. Because of the size and scope of these actively monitored registries, they represent the true commercial experience with our current generation high voltage leads.

As described on page 308, the Durata lead family was added to the CLAS registry study in 2013 to determine the prevalence and incidence of lead compromise evidenced by imaging (includes externalized conductors and other visual lead anomalies) and electrical dysfunction. As of February 28, 2018, a total of 982 patients implanted with Durata leads at 43 centers underwent fluoroscopic evaluation. These include 286 leads implanted in 2008, 418 leads in 2009, and 278 leads in 2010. The implant duration at enrollment was 4.5 ± 1.1 years for the Durata leads. At enrollment, 100% of the 982 leads were free of externalized conductors (EC). A total of 757 patients (77%) completed 1 year follow-up, 624 patients (64%) completed 2 years of follow up, and 479 (49%) completed 3 years of follow up. Through February 28, 2018, the implant duration of these Durata leads is 7.8 ± 1.5 years with a follow up duration after enrollment of 3.3 ± 1.5 years. The event-free survival rate for Externalized Conductors is shown in Figure 1. Through 9 years since implant, freedom from EC is 98.2%. There were 5 cases of externalized conductors for which two leads were due to external abrasion (one due to clavicular crush and one due to a tricuspid annuloplasty ring); one lead, implanted for 7.5 years, showed EC in a short region just proximal to the RV coil not protected by Optim insulation; and two leads, implanted for 7.5 and 8.5 years, exhibited EC with no external mechanism identified on fluoroscopy. The electrical function of all 5 of these leads with externalized conductors was normal. In 982 Durata leads evaluated, 97.8% have been free of electrical dysfunction (ED). Of the leads with ED, none had externalized conductors.

During an average implant duration of 7.8 years, with complete ascertainment via annual fluoroscopy, performance of Durata leads remains strong, with 97.8% leads free of electrical dysfunction, and 98.2% without externalized conductors through 9 years. None of the leads with externalized conductors exhibited electrical dysfunction and none of the leads with electrical dysfunction were associated with an externalized conductor. All pending fluoroscopy data has been adjudicated and the minimum enrollment of Durata leads has been met in the Cardiac Lead Assessment Study.

Focus on Clinical Performance

FIGURE 1: KAPLAN-MEIER ESTIMATE OF FREEDOM FROM EXTERNALIZED CONDUCTORS (EC) FOR DURATA LEADS



YEAR	0	1	2	3	4	5	6	7	8	9
AT RISK	973	973	973	964	927	869	727	490	219	43
CUMULATIVE EC EVENTS	0	0	0	0	0	1	1	2	4	5
EC FREE PROBABILITY	100.0%	100.0%	100.0%	100.0%	100.0%	99.9%	99.9%	99.7%	99.1%	98.2%

Focus on Clinical Performance

Beginning in 2006, three prospective, outcome-oriented, actively monitored registry studies have enrolled Durata™ and Riata™ ST Optim™ leads: the OPTIMUM registry, SCORE registry, and the SJ4 Post Approval Study (PAS). A total of 11,130 Optim insulated leads (8,264 Durata and 2,866 Riata ST Optim leads) were enrolled in these studies at 293 sites. The raw data from these registries, current as of February 28, 2018, were independently analyzed by the Population Health Research Institute (PHRI) of McMaster University/Hamilton Health Science. Their results quantify the performance of the Durata and Riata ST Optim™ leads in the categories of externalized conductors, all-cause insulation breach, and all-cause mechanical failures. An externalized conductor represents an outer insulation breach within the vascular or cardiac systems resulting in the normally contained conductors becoming visible outside the lead body. All-cause insulation breach includes all types of abrasion and other mechanical types of insulation damage, including externalized conductors. The all-cause mechanical failure category includes any insulation breach (including abrasion), conductor fracture, failure of a crimp, weld, or bond, or other mechanical failure. Additionally, if the lead is reported to have been “taken out of service” (extracted, capped, or electrically abandoned) and the site reports (i) noise artifact, abnormal pacing impedance, or abnormal high-voltage lead impedance or (ii) a large impedance change coupled with any of elevated pacing threshold, loss of sensing, loss of capture, oversensing, or undersensing then that is also included in the all-cause mechanical failure category.¹ Overall incidence rates for these three failure categories are provided in the table below.

AN INDEPENDENT ANALYSIS OF DURATA™ AND RIATA™ ST OPTIM™ LEAD FAILURE RATES IN ACTIVE REGISTRIES BY PHRI (DATA THROUGH FEBRUARY 28, 2018)

FAILURE CATEGORY	DURATA AND RIATA ST OPTIM %	DURATA AND RIATA ST OPTIM 95% CI	FREEDOM FROM FAILURES AT 10 YEARS (%)
Externalized Conductors	0.00%	0.00% - 0.03%	100%
All-Cause Insulation Abrasion	0.28%	0.19% - 0.38%	99.3%
All-Cause Mechanical Failures	1.36%	1.15% - 1.58%	96.5%

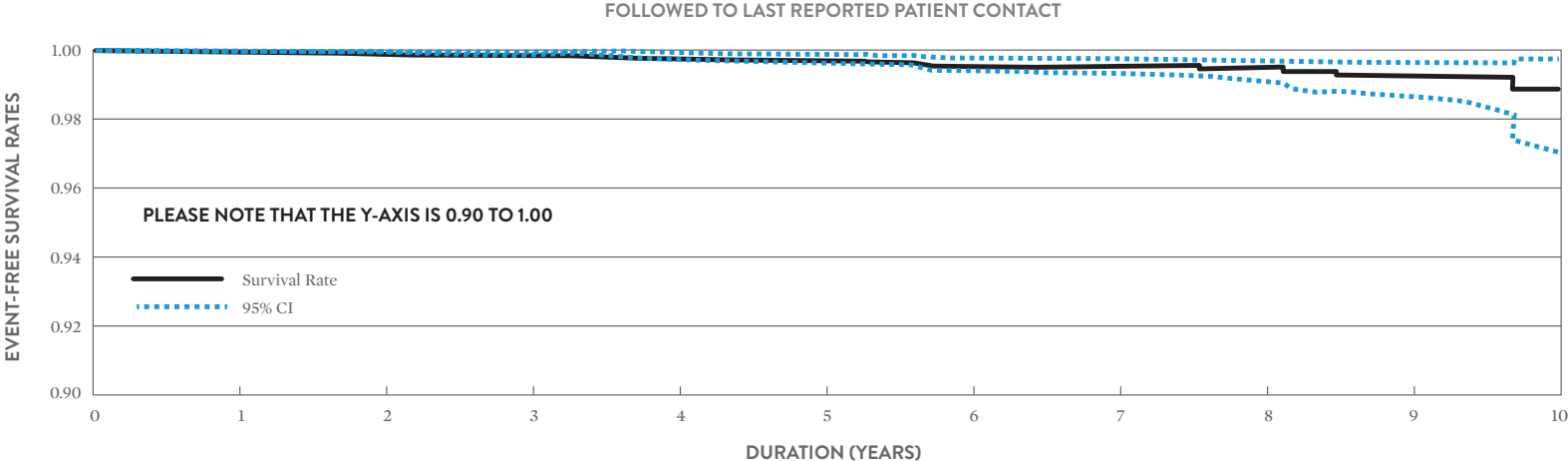
Event-Free Survival Rates for All-Cause Insulation Abrasion (Figure 2), and All-Cause Mechanical Failures (Figure 3) in Optim™ ICD leads were calculated by PHRI. The follow-up duration for active leads was based on last reported patient contact either in office or through remote monitoring. Lead implant date is used as time zero for these survival curves.

NOTE: The rates of all-cause mechanical failure, all-cause insulation abrasion and externalized conductors have been calculated based upon independent analyses of Abbott databases by PHRI. The total number of each event is derived from the sum of those events which have been independently adjudicated by PHRI and the recorded events awaiting adjudication. The final calculated rates may change slightly once adjudication is completed.

¹ John A. Cairns, Andrew E. Epstein, John Rickard, Stuart J. Connolly, Christopher Buller, Bruce L. Wilkoff, Janice Pogue, Ellison Themeles, Jeff S. Healey, *Prospective long-term evaluation of Optim-insulated (Riata ST Optim and Durata) implantable cardioverter-defibrillator leads*, Heart Rhythm, Vol 11, Issue 12, Pages 2156–2162, December 2014.

Focus on Clinical Performance

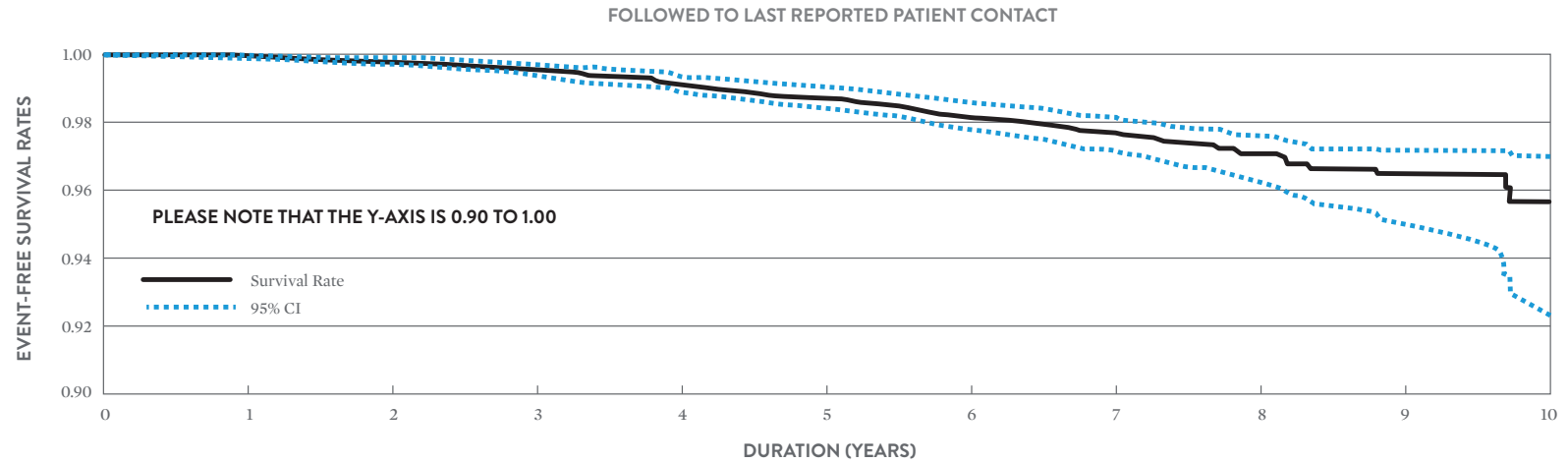
FIGURE 2: EVENT FREE SURVIVAL RATES FOR ALL-CAUSE INSULATION ABRASION IN OPTIM™ ICD LEADS AS CALCULATED BY PHRI



YEAR	0	1	2	3	4	5	6	7	8	9	10
LEADS AT RISK	11,130	9,885	8,578	7,357	6,225	4,916	3,742	2,787	1,783	838	239

Focus on Clinical Performance

FIGURE 3: EVENT FREE SURVIVAL RATES FOR ALL-CAUSE MECHANICAL FAILURE IN OPTIM™ ICD LEADS AS CALCULATED BY PHRI



YEAR	0	1	2	3	4	5	6	7	8	9	10
LEADS AT RISK	11,130	9,883	8,575	7,353	6,220	4,913	3,740	2,782	1,780	834	239

Focus on Clinical Performance

CUSTOMER REPORTED PERFORMANCE DATA

While large active registry data are robust for determining the true incidence rate of failures, passively collected data from worldwide complaints and returns analysis provides an important data source for better understanding the root cause of lead failures, as well as an appropriate method for comparing relative incidence rates of failure between lead models. The table below summarizes the incidence of insulation abrasion failure mechanisms confirmed by returns analysis of Riata™ ST Optim™ and Durata™ leads. Approximately 21,400 Riata ST Optim and Durata leads have been returned for analysis worldwide through February 28, 2018. Returned leads may exhibit more than one failure mechanism; hence the incidence rates presented in the table are not mutually exclusive.

DURATA™ (WW SALES 705,140) AND RIATA™ ST OPTIM™ (WW SALES = 33,000) LEADS INSULATION FAILURE MECHANISMS FROM COMPLAINTS AND RETURNS ANALYSIS

INSULATION FAILURE MECHANISM	ABRASION TYPE	OPTIM DEFIB LEAD WORLDWIDE (WW) INCIDENCE RATE (WW SALES = 738,250)
Intravascular – External*	External Abrasion	0.025%
Externalized Conductors – External**	External Abrasion	0.006%
Lead-to-Can*	External Abrasion	0.079%
Insulation Damage*	External Abrasion	0.025%
Intravascular - Inside Out*	Internal Abrasion	0.00176%***
Externalized Conductors - Inside Out**	Internal Abrasion	0.00014%***
Internal Abrasion Short Under RV Shock Coil*	Internal Abrasion	0.011%
Internal Abrasion Short Under SVC Shock Coil*	Internal Abrasion	0.008%

*Determined by returned product analysis.

**Includes cases determined by returned product analysis as well as cases identified only by fluoroscopy or visualization of explanted leads.

***These values reflect returns with a silicone insulation breach due to inside-out abrasion in the short region not protected by Optim insulation.

These incidence rates from complaints and returns analysis demonstrate the effectiveness of the Riata ST Optim and Durata lead design changes in reducing insulation-related failures when compared to the same type of data for Riata and Riata ST silicone leads (see page 316).

Update on Optim™ Lead Insulation

In 2006 Abbott brought to the cardiac rhythm management (CRM) market the first novel insulation technology in over 20 years: a silicone-polyurethane co-polymer known as Optim™ lead insulation, now featured in IsoFlex™ Optim™, Tendril™ STS, OptiSense™, QuickFlex™ μ, Quartet™, Durata™, and Optisure™ lead families. Optim lead insulation consolidates the best characteristics of two established CRM lead insulation materials, polyurethane and silicone.

The polyurethane content of Optim lead insulation imparts lubricity, strength, and abrasion resistance while the nearly 50% silicone content imparts flexibility and biostability.^{1,2} The clinical performance of >5.2 million Optim insulated pacing and tachycardia leads implanted worldwide continues to be excellent. All aspects of Optim lead insulation performance can be appreciated by referring to the Acute Observation, Chronic Complication, Lead Malfunction, and Survival Probability data found in this Product Performance Report. One noteworthy reliability benefit of Optim lead insulation is a significant reduction in the most common mode of lead malfunction: insulation abrasion.³ Insulation abrasion can occur as a result of lead motion and contact with pacemakers, ICDs, adjacent leads, or anatomical structures and can occur in the subcutaneous, intravascular, or intracardiac areas of the lead. Abrasion within a lead can occur as a result of contact between internal components, as noted in our November 2011 Riata™ lead advisory. The clinical effects associated with all types of insulation abrasion malfunctions can include sensing noise and changes in both pacing and defibrillation impedances and thresholds. As indicated in our December 2010 Riata communication, the presence of Optim™ lead insulation on the Riata™ ST Optim™ and Durata™ defibrillation lead family has greatly reduced the quantity of all abrasion types.

This Product Performance Report provides an up-to-date statistical assessment of the benefits of Optim lead insulation on Abbott tachycardia leads. A Kaplan-Meier analysis including all U.S. data through December 31, 2017 was performed on two groups of leads: (1) tachycardia leads with silicone insulation [Riata and Riata ST lead families], and (2) tachycardia leads with Optim lead insulation [Riata ST Optim and Durata lead families]. For each group, the U.S. registration and tracking data was combined with data from all U.S. confirmed abrasion malfunctions. This analysis does not include data from prospective registries or non-returned complaints. A Kaplan-Meier curve representing freedom from abrasion for both groups is provided below. The longest implant duration that is common to both model groups was 134 months. To provide a direct comparison of both model groups, the probability of an abrasion malfunction by 134 months of implant time is also presented in graphical format below.

1 C. Jenney, J. Tan, A. Karicherla, J. Burke, and J. Helland, "A New Insulation Material for Cardiac Leads with Potential for Improved Performance," HRS 2005, HeartRhythm, 2, S318-S319 (2005).

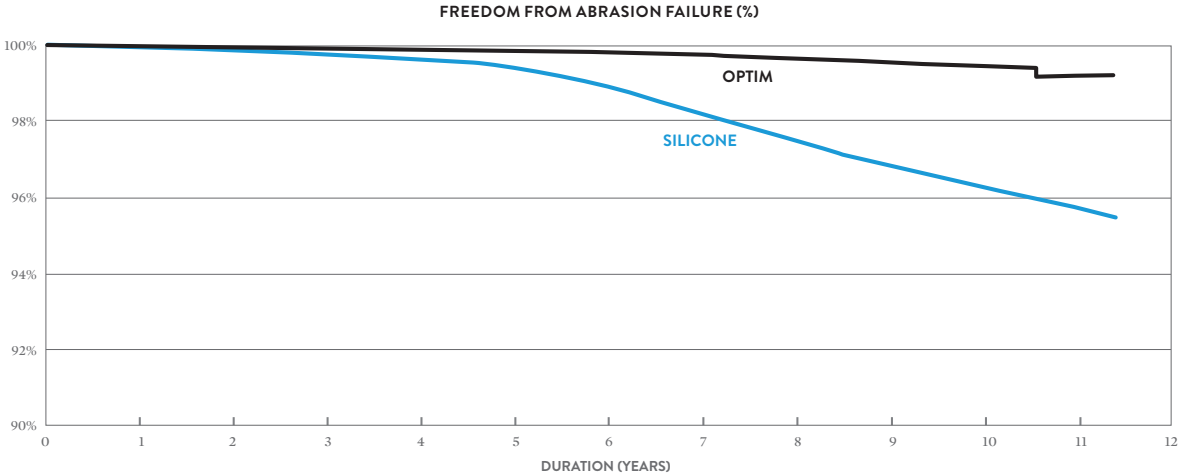
2 J. Tan and C. Jenney, "Comparative In Vivo Biostability Study of A New Lead Insulation Material Versus Polyurethanes," HRS2006, Heart Rhythm, 3, S146 (2006).

3 T. Kleemann, T. Becker, K. Doenges, M. Vater, J. Senges, S. Schneider, W. Saggau, U. Weisse, and K. Seidl, "Annual Rate of Transvenous Defibrillation Lead Defects in Implantable Cardioverter-Defibrillators Over a Period of >10 Years," Circulation, 115, 2474-2480 (2007).

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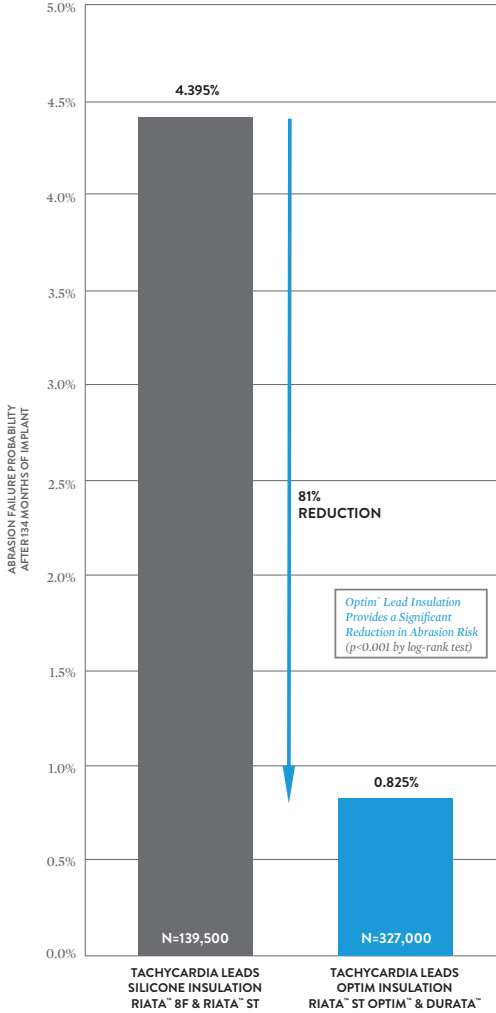
The data show that the presence of Optim™ lead insulation dramatically reduces the probability of abrasion malfunction in tachycardia leads at 134 months by 81%, which was confirmed to be statistically significant ($p < 0.001$) by a log-rank test.

OPTIM™ LEAD INSULATION EFFECTS ON ABBOTT TACHYCARDIA LEAD ABRASION KAPLAN-MEIER ANALYSIS OF U.S. RETURNS ANALYSIS DATA



YEAR	1	2	3	4	5	6	7	8	9	10	11	134 MONTHS
OPTIM	278,642	244,547	210,238	174,108	139,891	108,885	77,394	45,071	21,900	6,231	941	0
SILICONE	121,466	111,016	101,043	91,507	82,863	75,188	68,475	62,129	55,577	46,675	33,692	30,857

ABRASION MALFUNCTION PROBABILITY AFTER 134 MONTHS OF IMPLANT



Advisories & Safety Alerts

Advisories & Safety Alerts

The following table summarizes recalls, advisories and safety alerts regarding Abbott implantable devices since 1999. These advisories have been previously communicated to physicians. Advisories associated with non-implantable devices such as catheters are not included in the table. For more information please contact Abbott Technical Services at 1-800-722-3774.

ICD AND CRT-D DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
GLOBAL MODELS Excelsis Quadra™ (Models CD3281-40, CD3281-40Q) Excelsis™ + (Models CD3389-40C, CD3389-40QC) Excelsis™ CRT-D (Models CD3297-40, CD3297-40Q) Fortify Assura™ DR (Models CD2257-40, CD2257-40Q, CD2259-40, CD2259-40Q, CD2357-40C, CD2357-40Q, CD2359-40, CD2359-40C, CD2359-40Q, CD2359-40QC) Fortify Assura™ ST DR (Models CD2263-40, CD2263-40Q, CD2363-40C, CD2363-40Q) Fortify Assura™ ST VR (Models CD1263-40, CD1263-40Q, CD1363-40C, CD1363-40Q) Fortify Assura™ VR (Models CD1257-40, CD1257-40Q, CD1259-40, CD1259-40Q, CD1357-40C, CD1357-40Q, CD1359-40, CD1359-40C, CD1359-40Q, CD1359-40QC) Fortify™ DR (Models CD2231-40, CD2231-40Q, CD2233-40, CD2233-40Q) Fortify™ ST DR (Models CD2235-40, CD2235-40Q, CD2241-40, CD2241-40Q) Fortify™ ST VR (Models CD1235-40, CD1235-40Q, CD1241-40, CD1241-40Q) Fortify™ VR (Models CD1231-40, CD1231-40Q, CD1233-40, CD1233-40Q) HeartMinder™ + DR (Models CD2391-40C, CD2391-40QC) HeartMinder™ + VR (Models CD1391-40C, CD1391-40QC) HeartMinder™ ST DR (Models CD2299-40, CD2299-40Q) HeartMinder™ ST VR (Models CD1299-40, CD1299-40Q) Quadra + Excelsis™ (Models CD3385-40C, CD3385-40QC) Quadra Assura MP™ (Models CD3269-40, CD3269-40Q, CD3271-40, CD3271-40Q, CD3371-40, CD3371-40C, CD3371-40Q, CD3371-40QC) Quadra Assura™ (Models CD3265-40, CD3265-40Q, CD3267-40, CD3267-40Q, CD3365-40C, CD3365-40Q, CD3367-40, CD3367-40C, CD3367-40Q, CD3367-40QC) Unify Assura™ (Models CD3257-40, CD3257-40Q, CD3261-40, CD3261-40Q, CD3357-40C, CD3357-40Q, CD3361-40, CD3361-40C, CD3361-40Q, CD3361-40QC) Unify Quadra MP™ (Models CD3255-40, CD3255-40Q) Unify Quadra™ (Models CD3249-40, CD3249-40Q, CD3251-40, CD3251-40Q) Unify™ (Models CD3231-40, CD3231-40Q, CD3235-40, CD3235-40Q)	10/11/2016 Class I High voltage devices (ICDs and CRT-Ds) that utilize Lithium-based battery chemistries are subject to Lithium cluster formation during high voltage charging. Depending on their location, Lithium clusters may cause a short circuit that can lead to premature battery depletion. Our investigation indicates that if a short circuit occurs, battery depletion can occur in these devices within a day to a few weeks, which may result in the inability to deliver therapy. 8/28/2017 Class I Customers were made aware of the availability of a new battery performance management tool for detection of abnormal battery performance in devices subject to the October 2016 advisory.	In consultation with our Medical Advisory Board, we recommend the following: <ul style="list-style-type: none"> • Do not implant unused affected devices. • Conduct patient follow-up per standard practice. • Prophylactic device replacement is NOT recommended because complications following replacement have been reported to occur at a greater rate than the rate of harm associated with premature battery depletion due to lithium cluster induced shorts (see below for selected references). • In the event of an ERI indicator in these devices, immediate device change is recommended. At this time there is no factor, method or test to identify devices with this form of premature battery depletion approaching ERI or to accurately predict remaining battery life once ERI appears. • Physicians should reaffirm the availability of home monitoring to avoid or minimize time without device therapy for bradycardia and tachycardia events. • Enroll patients in Merlin.net™ Patient Care Network (PCN) utilizing the "DirectAlerts™" feature to provide you with an immediate alert notification in the event ERI is reached. For patients currently enrolled in Merlin.net PCN, remind them of the importance of using remote monitoring. • Review the most recent Programmed Parameters printout. <ul style="list-style-type: none"> • Ensure that under the "Trigger Alerts When" section, that the "Device at ERI" parameter is ON (it is normally ON) for both "Show on FastPath" and "Notify Patient" selections. • If the "Device at ERI" alert is OFF, we recommend that the patient be seen promptly to program this parameter ON. • Advise patients that an ERI indication triggers a vibratory alert. At the next scheduled office visit: <ul style="list-style-type: none"> • Interrogate the patient's device to determine if an ERI alert has been triggered. Premature battery depletion can be identified by physicians through home monitoring showing ERI or more advanced battery depletion. • Perform a patient notifier test to confirm that the patient feels and recognizes the vibratory alert. • Patients who cannot feel the vibratory alert may experience loss of battery and/or loss of device function without their awareness. • Advise the patient to contact your office promptly should they feel a vibratory alert. <ul style="list-style-type: none"> • In-office evaluation should be performed to determine the reason for the alert as other non-critical events can also trigger a vibratory alert. Device Replacement Complication Publications <ol style="list-style-type: none"> 1. John W. Moore III, William Barrington, et. al.; "Complications of replacing implantable devices in response to advisories: A single center experience"; International Journal of Cardiology 134 (2009) 42–46 (5.5% overall, 2.1% major complications) 2. Paul A. Gould, MBBS, PhD, Lorne J. Gula, MD, et. al.; "Outcome of advisory implantable cardioverter- defibrillator replacement: One-year follow-up"; Heart Rhythm, Vol 5, No 12, December 2008 (9.1% overall, 5.9% major complications, including two deaths) 3. Krystina B. Lewis, Dawn Stacey, R.N., Ph.D, et. al.; "Estimating the Risks and Benefits of Implantable Cardioverter Defibrillator Generator Replacement: A Systematic Review; PACE, Vol. 39, July 2016 (7.5% overall, 4.0% major complications) Current Status (February 28, 2018): At the time of the advisory, 841 returned devices (0.21%) of 398,740 devices worldwide have premature depletion in association with lithium clusters, including 549 in the US. As of February 28, 2018, there were additional occurrences for a cumulative worldwide total of 1,529, including 971 in the US, and the rate is now 0.38%. For additional information and to determine if a device serial number is subject to this advisory, please go to the following website: www.sjm.com/batteryadvisory

ICD AND CRT-D DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Ellipse[™] and Ellipse ST VR/DR US: CD1309, CD1311*, CD1409, CD1411*, CD2309, CD2311*, CD2409, CD2411* (all -36, -36Q, -36C and -36QC suffixes). *Denotes models also sold OUS. OUS: CD1277, CD1279, CD1293, CD1295, CD1377, CD1393, CD2277, CD2279, CD2293, CD2295, CD2377, CD2393 (all -36, -36Q, -36C and -36QC suffixes).</p>	<p>8/19/2014 Class II</p> <p>Extended Charge Time may result in delayed delivery of high voltage therapy or prevent delivery of part or all of a programmed high voltage therapy shock. The anomaly most commonly presents as a vibratory patient notification and upon interrogation, a programmer or Merlin.net[™] Patient Care Network (PCN) alert indicating a "Capacitor Charge Time Limit reached" message. This may occur during a capacitor maintenance or charging for high voltage therapy. The anomaly occurs as a result of internal damage to the capacitors used in the high voltage charging circuitry of the subject devices, which may result in an extended charge time. As designed, the device will deliver the available energy on the capacitors once the charge time limit of 32 seconds is reached, even if the energy is less than the programmed value. This condition is detectable as the device will initiate a vibratory patient alert and, for patients enrolled and actively being followed, a Merlin.net PCN notification. Additionally, upon device interrogation, an alert message will indicate "Capacitor Charge Time Limit reached" on a specific date. Approximately 97% of Ellipse ICD extended charge time events reported to St. Jude Medical have been detected during capacitor maintenance with the remainder detected during defibrillation threshold (DFT) testing. There have been no reported cases of an Ellipse device failing to deliver high voltage therapy to a patient when needed.</p>	<p>St. Jude Medical recommends that patients with affected devices be enrolled in Merlin.net Patient Care Network (PCN) so that any extended charge time alert ("Capacitor Charge Time Limit reached" message) will be transmitted to Merlin.net PCN for patients being actively monitored and can be viewed by your clinic staff. If your patient has received a vibratory notification and/or if a programmer or Merlin.net alert for an extended charge time has been observed:</p> <ul style="list-style-type: none"> • Schedule your Ellipse ICD patient for an in-office follow-up evaluation as soon as possible. • Interrogate the Ellipse ICD and perform a manual capacitor maintenance charge. Note the charge time to full charge; it should be approximately 15 seconds or less. • Contact St. Jude Medical's Technical Services Department at 800-722-3774 to review the results of the capacitor maintenance test and discuss if additional evaluation is required. • A device that has experienced repeated extended charge time out warnings should be considered for replacement. <p>As the large majority of the extended charge time events have presented at the routine 6 month automatic capacitor maintenance interval, programming the interval to every 4 months at your patient's next scheduled follow up visit may provide an earlier indication of this potential anomaly. It should be noted that changing the device programming to a 4 month capacitor maintenance interval will reduce device longevity by approximately 9%. Device replacement is not recommended for an Ellipse device exhibiting normal charge times, and patients should continue to be followed at routine follow up intervals, per HRS/EHRA Expert Consensus on Monitoring Cardiovascular Implantable Electronic Devices (CIED), April 2008.</p> <p>Current Status (December 31, 2017): At the time of the advisory, the worldwide event rate of extended charge time on the affected population was 0.42%, based on 179 extended charge time events out of 43,000 worldwide sales. As of December 31, 2017, there were additional reports and the rate is now 1.11%. There have been no reports of serious injury or death within this population.</p>

ICD AND CRT-D DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>AnalyST Accel[™] DR RF (Models CD2219-36, CD2219-36Q) AnalyST Accel[™] VR RF (Models CD1219-36, CD1219-36Q) Current Accel[™] DR RF (Models CD2215-36, CD2215-36Q) Current Accel[™] VR RF (Models CD1215-36, CD1215-36Q) Current[™] DR (Model 2207-36) Current[™] VR (Model 1207-36) Ellipse[™] DR (Models CD2277-36, CD2277-36Q, CD2377-36, CD2377-36Q, CD2377-36C, CD2377-36QC) Ellipse[™] VR (Models CD1277-36, CD1277-36Q, CD1377-36, CD1377-36Q, CD1377-36C, CD1377-36QC) Fortify Assura[™] DR (Models CD2259-40, CD2259-40Q, CD2359-40, CD2359-40Q, CD2359-40C, CD2359-40QC) Fortify Assura[™] VR (Models CD1259-40, CD1259-40Q, CD1359-40, CD1359-40Q, CD1359-40C, CD1359-40QC) Fortify[™] ST DR (Models CD2235-40, CD2235-40Q) Fortify[™] ST VR (Models CD1235-40, CD1235-40Q) Promote Accel[™] RF (Models CD3215-36, CD3215-36Q) Promote Quadra[™] (Models CD3239-40, CD3239-40Q) Promote[™] (Model 3213-36) Quadra Assura[™] (Models CD3267-40, CD3267-40Q, CD3367-40, CD3367-40Q, CD3367-40C, CD3367-40QC) Quadra Assura MP[™] (Models CD3371-40, CD3371-40Q, CD3371-40C, CD3371-40QC) Unify Assura[™] (Models CD3261-40, CD3261-40Q, CD3361-40, CD3361-40Q, CD3361-40C, CD3361-40QC) Unify Quadra[™] (Models CD3251-40, CD3251-40Q) Unify[™] (Models CD3235-40, CD3235-40Q)</p>	<p>1/23/2014 Outside US only</p> <p>In November 2013, St. Jude Medical released the Merlin[™] Programmer Software version 17.2.2 rev. 0 (herein after referred to as 17.2.2) as an upgrade to existing programmers. Testing has shown that, when using a programmer with the 17.2.2 software, an incorrect value for sinus redetection, potentially affecting the high voltage therapy delivery sequence can occur when a device is programmed to a single VF detection zone. The issue can be introduced during programming of certain families of St. Jude Medical[™] ICD/CRTD devices. The issue is not present when a device is programmed to a two or three zone configuration. When using the 17.2.2 software and any parameter is programmed as part of a single VF detection zone configuration, the sinus redetection value will be inappropriately set to zero milliseconds. As a result, any intrinsic activity following the first shock will be considered a "sinus rate" and the device will diagnose "return to sinus". Therefore, if the arrhythmia was not terminated by the initial high voltage therapy, the ongoing arrhythmia would be considered a new episode causing the next high voltage therapy to also be delivered at the first programmed energy level. For example, if the first shock is programmed to 20 joules and subsequent shocks are programmed to higher energy values, the only HV therapy the patient would receive if the arrhythmia continues and is redetected, would be 20 joules, rather than the increasing HV energy levels as programmed.</p>	<p>Immediate Resolution Steps:</p> <ul style="list-style-type: none"> Review your SJM[™] ICD/CRT-D patient records for patients with affected devices implanted or seen in clinic starting in September 2013 and programmed to a single VF detection zone with the 17.2.2 software. For patients identified during this review we recommend that you schedule an immediate follow-up visit. The programmer software version is printed on the bottom of each report page. For patient devices programmed as described above with 17.2.2 software, a new software version 17.2.3 will correct this issue and is expected to be available by February 2014. Your St. Jude Medical representative will assist you with obtaining and installing the 17.2.3 software on your programmer. Using this software, programming any parameter will reset the return to sinus criteria to normal function. If a patient is seen before the 17.2.3 software is installed, then program the device to a two or three zone configuration, even if one of the zones is strictly a monitor zone. This will resolve the issue when using a programmer with 17.2.2 software. <p>Current Status (December 31, 2017): No occurrences have been reported following the field communication and correction.</p>

ICD AND CRT-D DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Fortify™ ST (Models CD1235-40, CD1235-40Q, CD2235-40, CD2235-40Q)	<p>4/18/2013 Outside US only</p> <p>The Merlin™ PCS programmer software Model 3330 versions 14.2.2, 16.2.1 and 17.2.1.1 provide new features for St. Jude Medical™ ICDs, including an option to enhance the ST diagnostic features in Abbott Fortify™ ST ICD models CD1235-40, CD1235-40Q, CD2235-40 and CD2235-40Q via a device software upgrade. During a device software upgrade, implanted devices are temporarily placed into the back-up pacing (BVVI) and back-up defibrillation only (BDFO) mode. The back-up mode parameter settings will be in effect for the two minute upgrade process. Once the upgrade successfully completes, the device will revert to the previously programmed parameter settings. Depending on the individual patient, this temporary change in parameter values while in back-up defibrillation only mode could make the device susceptible to oversensing and potentially deliver high voltage therapy during the upgrade procedure.</p>	<p>In order to prevent the potential for inappropriate therapy during the software upgrade process, consider programming the “Tachy Therapy Enabled/Disabled” function to Disabled prior to proceeding with the software upgrade. It is imperative to re-interrogate the device and program the “Tachy Therapy Enabled/Disabled” function to Enabled after the upgrade has been successfully completed. As with any device evaluation and programming, ECG monitoring and availability of back up external defibrillation equipment is recommended during the entire software upgrade process.</p> <p>Current Status (December 31, 2017): At the time of the advisory there were 20 devices confirmed to be affected by this issue. As of December 31, 2017 there were an additional 52 devices confirmed with this issue. There have been no reports of erious injury or death.</p>

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Convert™+ (Model V-195)	<p>5/6/2010 Outside US only</p> <p>A condition where devices programmed to a two-zone tachy therapy configuration, using a Merlin™ Patient Care System (PCS) programmer running version 7.2.1, 8.2.1 or 10.2.0 software, can result in the VT Therapy Timeout parameter being programmed OFF and HV therapy not being available if ATP therapies are unsuccessful.</p>	<p>If a patient's device is already programmed to a two zone configuration with a Merlin™ PCS programmer running version 7.2.1, 8.2.1 or 10.2.0, a follow-up visit should be scheduled to perform the recommendations outlined below:</p> <p>A permanent correction is available in the new release of the Merlin™ PCS programmer software version 10.2.2 which has received regulatory approval. Subsequently using a Merlin™ programmer with 10.2.2 (or later) software and following the steps outlined below will ensure that the VT Therapy Timeout parameter is programmed ON.</p> <ol style="list-style-type: none"> 1. Interrogate the Convert+ ICD and verify that it is programmed to a two zone configuration. 2. Program the device to a single zone, fibrillation only tachycardia mode. This action will program the VT Therapy Timeout parameter ON. 3. Re-program the device to the desired two zone configuration. (VT Therapy Timeout will remain ON). <p>If your patient's device is programmed to a single zone (fib only) there is no need to perform any reprogramming action.</p> <p>As these actions fully correct the potential issue there is no need to consider any device explant.</p> <p>Current Status (December 31, 2017): At the time of the advisory there was one report of this issue out of approximately 330 Convert+ Model V-195 ICDs distributed in Europe and Asia. As of December 31, 2017, there have been no additional reports associated with this advisory.</p>

ICD AND CRT-D DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Epic[™] ICDs (Models V-197, V-235, V-337, V-338, V-339), Epic[™] + ICDs (Models V-196, V-233, V-236, V-239, V-350) Epic[™] II ICDs (Models V-158, V-255, V-258, V-355, V-356, V-357) Atlas[™] + ICDs (Model V-340, V-341, V-343, V-193, V-242, V-243) Atlas[™] II ICDs (Models V-168, V-265, V-268, V-365, V-366, V-367)</p>	<p>1/16/2008 Class II</p> <p>A very rare condition (incidence of eight in 143,000 devices worldwide; six in the US and two outside the US) that could lead to a ventricular sensing anomaly in Epic[™] and Atlas[™] family of implantable cardioverter defibrillators (ICDs) has been identified. A loss of ventricular sensing would prevent an ICD from being able to detect an arrhythmia. The loss of ventricular sensing anomaly can only occur when the device's software writes to a particular memory location and only if there is a precise alignment of two timing parameters that normally do not coincide during routine operation of the device. The precise alignment requires the software write to occur at the exact time that a comparison is made during a specific 61 microsecond (μsec) window.</p>	<p>A simple programmer software/device firmware upgrade will resolve the issue and prevent a future occurrence. Patients who are followed on a routine basis with scheduled follow-up visits every three to six months should continue with their scheduled visit. Upon interrogation of one of the subject devices, the Merlin[™] Patient Care System and Model 3510 programmers with the newly provided software will automatically identify a device that can benefit from a firmware upgrade and will instruct the clinician that an upgrade is available.</p> <p>St. Jude Medical, along with our independent Medical Advisory Board members, has determined that no other action is recommended.</p> <p>Current Status (December 31, 2017): At the time of the advisory, there were 8 worldwide (6 U.S.) devices confirmed to have been affected by this issue. As of December 31, 2017 there have been no additional devices confirmed to have this issue since the time of the advisory.</p>

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Photon[™] DR (V-230HV) (certain serial numbers), Photon[™] Micro VR/DR (Models V-194, V-232), Atlas[™] VR/DR (Models V-199, V-240)</p>	<p>10/7/2005 Class II</p> <p>A particular vendor-supplied memory chip can be affected at a low frequency rate by background levels of atmospheric ionizing cosmic radiation ("background cosmic radiation"). The anomaly can trigger a temporary loss of pacing function and permanent loss of defibrillation support.</p>	<p>In the unlikely event that a device chip is affected by background cosmic radiation, the high current drain condition will deplete the battery voltage rapidly. This can result in loss of output for a period up to approximately 48 hours. During this period, the patient would be without pacing or defibrillation therapy. After this initial period, the battery will reach a voltage level at which the device will enter its "Hardware Reset Mode." This safety mode is designed to preserve the device's ability to provide VVI pacing support. A device that has been reset to the Hardware Reset Mode will operate in the VVI mode at 60 ppm, but will not be capable of providing tachycardia detection or therapy. This will be noted by a warning message on the programmer screen upon device interrogation.</p> <p>To assist in your patient care and following discussions with our independent Medical Advisory Board, Abbott recommends:</p> <p>If it is not already your current practice, physicians should perform routine device monitoring every three months for patients with the affected models listed above.</p> <p>In determining whether additional patient management or follow up may be needed, consider the low failure rate for the anomaly and the unique medical needs and situation of each individual patient, including whether the patient is pacemaker dependent or at high risk for life-threatening arrhythmias.</p> <p>If a patient's device is found in the Hardware Reset Mode, you should arrange for device replacement as soon as possible. You should continue to provide patients with the usual admonitions to keep scheduled appointments and to report all changes in symptoms.</p> <p>Current Status (December 31, 2017): At the time of the advisory, there were 60 worldwide (38 U.S.) devices confirmed to have been affected by this issue. As of December 31, 2017 there were an additional 42 worldwide (28 U.S.) devices confirmed with this issue. This is within the 95% confidence interval prediction made at the time of the advisory. There have been no reports of serious injury or death.</p>

ICD AND CRT-D DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Epic[™] DR/HF (V-233, V-337, V-338), Epic[™] Plus DR/VR/HF (V-236, V-239, V-196, V-239T, V-196T, V-350), Atlas[™] DR (V-242), and Atlas[™] Plus DR/VR/HF (V-243, V-193, V-193C, V-340, V-341, V-343)</p>	<p>6/13/2005 Class II</p> <p>Two anomalies have been identified:</p> <ol style="list-style-type: none"> 1. Due to a device software anomaly, it is possible that when the device's battery is nearing its elective replacement indicator (ERI), a charging cycle may be skipped. 2. After a capacitor charge, if a rate responsive pacing mode (e.g., DDDR, VVIR, etc.) is programmed "On," this "noise" may be interpreted by the device's accelerometer (activity sensor) as physical activity, causing a temporary increase in the pacing rate that may persist after charging is completed. 	<p>Two anomalies were discovered during routine product monitoring. Neither of these anomalies presents a significant clinical risk to your patients, and no clinical complications have been reported to Abbott. Both are easily corrected by performing a simple, automated software download to the device. This potentially affects approximately 30,000 implanted ICDs in the United States and includes the following model numbers:</p> <p>Epic[™] DR/HF (V-233/V-337/V-338), Epic[™] Plus DR/VR/HF (V-236/V-239/V-196/V-239T/V-196T/V-350), Atlas[™] DR (V-242), and Atlas[™] Plus DR/VR/HF (V-243/V-193/V-193C/V-340/V-341/V-343). The first anomaly can occur when one of the affected devices attempts to deliver multiple shocks in rapid succession. Due to a device software anomaly, it is possible that when the device's battery is nearing its elective replacement indicator (ERI), a charging cycle may be skipped. If this were to occur, the first shock will always be delivered as programmed and, if needed, the next shock in the programmed sequence would be delivered after a delay of only two to four seconds. A skipped charge would result in less than the full number of programmed shocks being available for delivery during that episode, but all delivered shocks would be at their programmed energy. This behavior was discovered as an incidental finding during analysis of one returned device that had delivered a large number of high-voltage shocks over a short time period.</p> <p>A second anomaly is caused by electrical "noise" generated as a result of the charging of the device's high-voltage capacitors. After a capacitor charge, if a rate responsive pacing mode (e.g., DDDR, VVIR, etc.) is programmed "On," this "noise" may be interpreted by the device's accelerometer (activity sensor) as physical activity, causing a temporary increase in the pacing rate that may persist after charging is completed. The degree and duration of the rate increase will depend on a variety of factors, but the rate will never exceed the programmed Maximum Sensor Rate, and the device will gradually return to the appropriate rate. The anomalous behavior, which has been observed during the performance of manual capacitor maintenance, has been traced back to a component supplied to Abbott by one vendor; therefore, only the subset of the device models listed above that were manufactured with the affected component (device serial numbers below 141000 for any model) will exhibit this behavior.</p> <p>Abbott has developed programmer software that will automatically detect the affected ICDs and download device software that will correct the "skipped charge" anomaly and mitigate the response to electrical noise. Once the upgrade is performed, the potential for a skipped charge will be eliminated. Additionally, once the upgrade is performed if a rate responsive mode is programmed "On," devices with serial numbers below 141000 will have their rate response functions suspended for the time period during which the electrical noise could be present (i.e., while significant residual voltage remains on the high-voltage capacitors); non-rate responsive pacing at the programmed base rate will continue to be provided as appropriate. This period during which rate response is suspended may last anywhere from a few minutes up to approximately 90 minutes. If rate responsive pacing was ongoing prior to charging, the pacing rate will gradually decrease to the base pacing rate according to the normal rate response recovery algorithm and will remain there while rate responsive pacing is suspended. The rate response behavior for devices with serial numbers greater than 141000 will not be affected by the software download.</p> <p>The software download for potentially affected devices will automatically be initiated the next time the patient's device is interrogated with the v4.8.5 programmer software. Since a skipped charge is more likely to occur in devices that are closer to their elective replacement indicator (ERI), Abbott recommends that if the next patient follow-up is not scheduled to occur within the next six months that the patient be seen within this time period.</p> <p>In addition, if devices are programmed to pacing settings that result in high current consumption, such as high output bi-ventricular pacing, consideration should be given to scheduling the patient for a follow-up visit within three months if it is not scheduled to occur within that time period. As always, Abbott defers to your clinical judgment on any decisions regarding the management of your patients.</p> <p>Current Status (December 31, 2017): There have been no implanted devices confirmed to have been affected by this issue since the time of the advisory.</p>

ICD AND CRT-D DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Epic [™] (V-197, V-235), Epic [™] + (V-196, V-236), Epic [™] HF CRT-D (V-338), Epic [™] + HF CRT-D (V-350), Atlas [™] + (V-193, V-243), Atlas [™] + HF CRT-D (V-340), or Atlas [™] (model V-242) ICDs	3/10/2005 Class II A software parameter that affects the sensitivity of the reed switch in the listed devices was being set to an incorrect value which could prevent these devices from entering the magnet mode to inhibit tachy therapy when an external magnet is applied.	During routine product evaluation, Abbott Quality Assurance identified that a software parameter that affects the sensitivity of the reed switch in the listed devices was being set to an incorrect value during manufacturing beginning in late November of last year. This has the effect of preventing these devices from entering the magnet mode to inhibit tachy therapy when an external magnet is applied. This is a software controlled parameter that can be easily corrected via the programmer. All other bradycardia pacing and tachyarrhythmia detection and therapy features are not affected in devices subject to this notification. Until the magnet sensitivity parameter is corrected via the programmer, tachy therapy may not be properly inhibited as is customary with placement of an external magnet, but can be inhibited by using the programmer to program the device to Defib Off, and then back On as needed. The affected devices were manufactured during a three month period beginning November 22, 2004. To date, there have been no field reports of any magnet mode failures, nor have there been any clinical complications reported associated with this issue. Magnet mode application is usually used to inhibit tachycardia therapy such as when a patient is subjected to electrocautery during a surgical procedure. In order to remedy this situation, in addition to this notification, Abbott Sales Representatives and Field Clinical Engineers have been provided with a simple software tool that can be used to set, via the programmer, the reed switch's magnet sensitivity to the proper value. You may contact them to schedule this reprogramming at the patient's next scheduled follow-up visit or at your discretion. Current Status (December 31, 2017): There have been no implanted devices confirmed to have been affected by this issue since the time of the advisory.

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Profile [™] V-186	7/13/2001 Class II Failure of a ceramic capacitor could lead to sensing anomalies/early battery depletion	This Advisory applies to a well-defined group of Profile [™] MD (Model V-186HV3) implantable cardioverter-defibrillators (ICDs) which have been observed to exhibit decreased reliability. These devices had specific modules (subcircuits) manufactured during a five-month period during the first half of 1999. Those potentially affected devices that have not exhibited any anomalous behavior within 24 months of manufacture are not at increased risk of failure at a later date. These failures were caused by a component anomaly that is limited to a specific, traceable lot (group) of capacitors. If these capacitors fail, two different clinical manifestations may be observed, depending on the location of the failed capacitors: Low Voltage Module: The devices may exhibit sudden loss of sensing or oversensing of internally generated interference (noise) resulting in aborted or delivered shocks. Due to the potential for sudden loss of sensing, which would prevent the ICD from detecting ventricular tachyarrhythmias, device replacement should be considered for patients with devices incorporating the suspect capacitors in this location. In making this decision, you should weigh the likelihood that your patient will have one of the few additional devices expected to fail against the risk associated with the replacement procedure. For patients who do not have their devices replaced, Abbott recommends monthly monitoring for the remaining months during which additional failures are expected to occur and every three months thereafter. High-Voltage Module: The devices may exhibit premature battery depletion with no other clinical manifestations. While the battery voltage may drop rapidly to a value near or below the device's ERI, it stabilizes for several months at a value at which the device continues to detect arrhythmias and deliver therapy appropriately. In addition, some of these devices may exhibit oversensing of internally generated electrical interference (noise), which may result in false detection of tachyarrhythmias. However, during charging, the interference abates, and the therapy is usually aborted without the delivery of inappropriate shocks to the patient. Because these devices generally continue to respond appropriately to spontaneous tachyarrhythmias, and because the battery voltage stabilizes for several months at a level that does not compromise device function, Abbott recommends that patients with devices incorporating suspect capacitors in this location be monitored monthly for the remaining months during which additional failures are expected to occur and every three months thereafter.

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Nanostim™ Leadless Cardiac Pacemaker (Model SIDLCP)</p>	<p>11/17/2017 Outside US and US Investigational Device Exemption (IDE) only</p> <p>Abbott was made aware of docking button detachments that have occurred following implant or during attempted retrieval of Nanostim™ Leadless Cardiac Pacemaker (LCP) devices. The docking button is a small component (3.6 mm diameter) and is connected to the end of the LCP by two cables. This component is necessary for docking the LCP to the retrieval catheter during a retrieval procedure.</p>	<p>The following patient management recommendations have been developed in consultation with our Leadless Steering Committee members after discussions detailing the occurrences and the potential clinical impact associated with detached docking buttons:</p> <ul style="list-style-type: none"> • Continue following patients as per recommendations of the October 2016 Battery Malfunction for Nanostim™ LCP advisory. • Retrieval of an implanted Nanostim™ LCP with an intact docking button confirmed radiographically remains an option, but should only be considered if the procedure can be performed as per the specifications contained in the instructions for use. <ul style="list-style-type: none"> • If a detached docking button has been identified, Nanostim™ LCP retrieval is not recommended. In the rare situation where retrieval is the only management option, please contact the Abbott Clinical Study Team for further guidance. • Prophylactic imaging for the sole purpose of determining if the docking button is intact is not recommended due to the effects of radiation and lack of any clear clinical actions based on results of imaging alone. If the option of Nanostim™ LCP retrieval is being considered, final imaging decisions should take into account the individual patient circumstances and preferences. • If a detached docking button is identified, continue to follow the patient as per the Clinical Study Protocol and report the incident to Abbott and relevant Competent Authority, as appropriate. <p>Current Status: (December 31, 2017) At the time of the advisory, three (0.21%) of 1,423 devices implanted worldwide have been reported to have a detached docking button. As of December 31, 2017, there was one additional report and the rate is now 0.28%. There have been no reports of serious injury or death.</p>

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Global Models Accent™ MRI™ (Model PM1224) Accent™ DR RF (Models PM2210, PM2212) Accent MRI™ (Models PM2218, PM2224) Accent™ SR RF (Model PM1210) Accent™ ST DR RF (Models PM2216, PM2222) Accent™ ST MRI DR RF (Model PM2226) Accent™ ST MRI SR RF (Model PM1226) Accent™ ST SR RF (Model PM1222) Allure Quadra™ RF CRT-P (Model PM3242) Allure™ RF CRT-P (Model PM3222) Anthem™ RF CRT-P (Models PM3210, PM3212) Assurity™ + DR RF (Model PM2260) Assurity™ + SR RF (Model PM1260) Assurity™ DR RF (Model PM2240) Assurity MRI™ (Model PM2272) Assurity™ SR RF (Model PM1240) Assurity MRI™ (Model PM1272) Nuance™ DR RF (Model PM2214) Nuance™ MRI DR RF (Model PM2230) Nuance™ MRI SR RF (Model PM1230) Nuance™ SR RF (Model PM1214) Nuance™ ST DR RF (Model PM2228) Nuance™ ST SR RF (Model PM1228) Quadra Allure MP™ (Model PM3562) Quadra Allure MP™ RF CRT-P (Model PM3262) Quadra Allure™ (Model PM3542) Quadra Relieve MP™ (Model PM3564) Quadra Relieve MP™ RF CRT-P (Model PM3264) Quadra Relieve™ (Model PM3544) Quadra Relieve™ RF CRT-P (Model PM3244) Relieve™ RF CRT-P (Model PM3224) Zenex™ + DR RF (Model PM2270) Zenex™ + SR RF (Model PM1270) Zenex™ DR RF (Model PM2250) Zenex™ DR RF MRI (Model PM2282) Zenex™ SR RF (Model PM1250) Zenex™ SR RF MRI (Model PM1282)</p>	<p>8/28/2017 Class II</p> <p>New pacemaker firmware was developed to further mitigate the risk of unauthorized access to our pacemakers that utilize radio frequency (RF) communications. The firmware update provides an additional layer of security against unauthorized access to these devices that further reduces the potential for a successful cybersecurity attack.</p>	<p>Patient Management Recommendations</p> <p>Prophylactic replacement of affected devices is not recommended.</p> <p>While not intended to serve as a substitute for your professional judgment as to whether the firmware update is advisable for a particular patient, we, along with our Cyber Security Medical Advisory Board, recommend the following:</p> <ul style="list-style-type: none"> • Discuss the risks and benefits of the cybersecurity vulnerabilities and associated firmware update with your patients at the next regularly scheduled visit. As part of this discussion, it is important to consider patient specific issues such as pacemaker dependence, age of device, and patient preference and provide them with the "Patient Communication". • Determine if the update is appropriate given the risk of update for the patient. If deemed appropriate, install this firmware update following the instructions on the programmer (and listed below). • For pacing dependent patients, consider performing the cybersecurity firmware update in a facility where temporary pacing and pacemaker generator change are readily available, due to the very small estimated risk of firmware update malfunction. <p>Current Status (December 31, 2017): We have received no reports of device compromise related to the cybersecurity vulnerabilities in the implanted devices impacted by this communication. This release is part of planned system updates to ensure effective and secure products for our patients.</p> <p>If you have any questions about the cybersecurity firmware update you can contact your Abbott representative or our dedicated customer technical support hotline at 1-800-722-3774 (U.S.).</p> <p>Additional materials, including a Patient Communication, can be found on www.sjm.com/notices.</p>

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Nanostim™ Leadless Cardiac Pacemaker (Model SIDLCP)</p>	<p>10/28/2016 Outside US and US Investigational Device Exemption (IDE) only</p> <p>Abbott was made aware of seven (7) reports worldwide of lost telemetry and pacing output as a result of a battery malfunction associated with Nanostim Leadless Cardiac Pacemaker (LCP) devices. Due to these events, we have decided to pause Nanostim LCP implants in the Leadless II IDE/CAP study.</p> <p>Analysis of returned units has found decreased battery capacity due to reduced electrolyte, resulting in high internal battery resistance. This disrupts the required capacity for proper device function and reduces device longevity.</p> <p>Referring to a previously measured battery voltage may not provide an indication of continued normal operation as battery voltage remains normal under these circumstances. The Recommended Replacement Time (RRT) indicator will not be triggered as the battery voltage will remain above RRT in these devices. Battery malfunction may be indicated with a loss of telemetry/communication with the implanted device and/or loss of pacing and magnet mode operation.</p>	<p>In consultation with our Leadless II IDE and our Leadless Postmarket Study Steering Committees we recommend the following:</p> <ul style="list-style-type: none"> • Do not implant unused devices and return them to Abbott. • Do not rely on the RRT indicator to identify a battery that may potentially malfunction. However, if the RRT indicator does trigger, replace the device per standard practice. • Do not perform AV Node ablation in patients with an existing Nanostim LCP without another functional pacing system implanted. • For patients who have not previously been documented to be pacemaker dependent, re-assess patients in-office for pacemaker dependence. • For non-pacemaker dependent patients with devices of implant duration ≥ 24 months, more intensive follow-up and monitoring is recommended. <ul style="list-style-type: none"> • Implant Duration ≥ 24 months: Request follow-up as soon as possible to assess the status of the battery. Then, monthly follow-up is recommended through in-office visits or a reliable method of tele-monitoring of heart rate and electrocardiogram. • Implant Duration < 24 months: Continue follow up per protocol. • For pacemaker dependent patients, device replacement is recommended (priority should be for patients with implants of longer duration). <ul style="list-style-type: none"> • Identify and treat patients as quickly as possible. • Interrogate the device and identify the ability to communicate with the device and the patient's underlying rhythm. • Determine the strategy for management, including a decision whether to retrieve or abandon the Nanostim LCP, based on the individual patient's clinical history and overall medical condition. Use a temporary pacemaker for backup pacing while replacing the Nanostim device where clinically indicated. • If the device is to be retrieved, use the Nanostim retrieval system as per the standard procedure described in the instructions for use. • If the device will not be retrieved or if retrieval was attempted and not possible, implant a new pacemaker lead (bipolar) at a distance from the existing LCP to prevent long-term mechanical and electrical interactions. Confirm the location using multiple radiographic views. • After implantation of the new pacing system, if it is possible to communicate with the LCP, turn "OFF" the abandoned LCP system. If the LCP device cannot be turned "OFF", consider programming the newly implanted system a minimum of 5 bpm faster than the LCP in order to inhibit the Nanostim device. <p>Current Status (December 31, 2017): At the time of the advisory, seven (7) reported devices (0.50%) of 1,423 implanted devices worldwide have exhibited battery malfunction at 29 to 37 months after implant. As of December 31, 2017, there were additional reports and the rate is now 3.6%. There have been no reports of serious injury or death.</p>

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Accent [™] SR (Model PM1110) Accent [™] DR (Model PM2112)	<p>12/7/2012 Outside US Only</p> <p>Due to an incorrect software setting, a specific subset of the Accent[™] SR and Accent[™] DR devices shipped to certain countries outside the US will not provide a change in the sensor driven (rate responsive) pacing rates in response to patient physical activity. All other programmed parameters, features and functions operate as designed, e.g. an Accent DR device programmed to DDDR will appropriately track atrial activity and properly function in the DDD mode. A non-invasive programmer software solution that will correct the issue in all affected, implanted devices will be available once regulatory approval has been completed.</p>	<p>Abbott makes the following recommendations:</p> <p>Identify affected patient</p> <ul style="list-style-type: none"> Review your patient's clinical indications for pacing and determine the clinical need for rate responsive, sensor driven pacing. In the event that a patient requires rate responsive sensor driven activity pacing and exhibits clinical symptoms due to the lack of increased pacing rates with exercise, please contact your local Sales Representative or our Technical Support Continue to follow patients on their standard follow-up schedule. <p>Current Status (December 31, 2017): The programmer software update was released in April 2013. At the time of the advisory, approximately 6,000 affected devices were implanted. There have been no additional devices confirmed to have this issue since the time of the software release in April 2013.</p>

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Accent [™] DR (Models PM2110, PM2112, PM2210, PM2212), Anthem [™] CRT-P (Models PM3110, PM3112, PM3210, PM3212)	<p>9/22/2011 Class II</p> <p>A small amount of electrical charge may accumulate within an internal capacitor which results in a low or varying pacing lead impedance (PLI) value during the automatic daily measurements. An out of range lead impedance measurement may result in a patient notifier alert, a remote monitoring Merlin.net[™] Patient Care Network alert, or a prior alert message to be displayed on the programmer screen at the next in-clinic follow-up.</p>	<p>In order to prevent a false reading, a new Merlin[™] Patient Care System programmer software version is available. When used to interrogate an Accent DR or Anthem CRT-P pacemaker this software will eliminate the potential for this anomaly to occur. With the new software, the programmer will automatically activate circuitry in the pacemaker during the interrogation process to ensure that any residual charge on the capacitor is discharged prior to performing the daily PLI measurement. The onetime upgrade is performed automatically on affected devices and will not change the operation of the implanted device. Your St. Jude Medical Sales Representative will assist you in loading the new programmer software onto your Merlin programmer.</p> <p>If you are following any patients implanted with Accent DR pacemakers or Anthem CRT-P devices, St. Jude Medical makes the following recommendations, which are consistent with standard best practices:</p> <ul style="list-style-type: none"> Ensure that the new programmer software version is loaded on your programmers as soon as practical. Continue to follow patients on their standard follow-up schedule. Since the likelihood of the low lead impedance measurement is low, the device interrogation can be performed at the patient's next regular scheduled follow-up visit. In the event that a patient receives a low lead impedance alert before the new programmer software has been loaded, we suggest that you evaluate the device as you normally would for any such instance. If the daily pacing lead impedance value is out-of range, re-interrogate the device's measured data and look at the lead impedance values. This "in-clinic" measurement is not affected by the aforementioned capacitor charge build-up and will provide an accurate lead impedance measurement. <p>Current Status (December 31, 2017): Worldwide, 13 Accent DR (<0.01%) and 225 Anthem CRT-P (1.6%) devices have exhibited this diagnostic anomaly.</p>

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Identity [™] SR (Model 5172) Identity [™] DR (Model 5370) Identity [™] XL DR (Model 5376)	10/12/2006 Class II A programmer software anomaly can lead to incorrect reporting of battery voltage, expected battery longevity and elective replacement indicator (ERI) status in Abbott Identity [™] pacemakers. The anomaly does not affect the device's actual battery voltage, longevity or functionality, but could result in inaccurate reporting of the status of these measured data parameters. This software anomaly can appear in the Abbott Identity [™] family of pacemakers when programmed by the Abbott APS [™] III Model 3500/3510 or Merlin [™] Patient Care System Model 3650 programmers.	No follow-up is recommended at the time of advisory. Devices do not need to be replaced. A programmer software update, pending FDA and other regulatory agency approval, will mitigate this anomaly when the device is interrogated. Before the programmer software update is available, any subsequent measured data update that is performed during the session would be valid and the device operating magnet rate would be up-to-date. After the programmer software update is available, any device affected by this issue will be automatically corrected via the normal interrogation process. Current Status (December 31, 2017): At the time of the advisory, there were 53 worldwide (50 U.S.) devices confirmed to have been affected by this issue. As of December 31, 2017 there were an additional 78 worldwide (65 U.S.) devices confirmed with this issue, all prior to the distribution of the software fix.
Identity ADx [™] DR (Models 5286, 5380, 5386, 5480)	7/31/2003 Class II An anomaly in the pacemaker code, which under a certain set of programmed settings and specific event timing circumstances could deliver a short coupled pacing interval of 300 ms (200 ppm). Consecutive (up to a maximum of 12) shorter than anticipated pacing intervals could be experienced.	Abbott's software engineering department has identified an anomaly in the pacemaker code, which under a certain set of programmed settings and specific event timing circumstances has a potential to deliver a short coupled pacing interval of approximately 300 milliseconds (200 ppm). It is also possible, however unlikely, that a patient could experience consecutive (up to a maximum of 12) shorter than anticipated pacing intervals. To date, the only clinical observation has been the delivery of a single short interval. There have been no reports of patient injury as a result of the shortened pacing interval. In order for the above mentioned shortened pacing interval to potentially occur, the programmed settings of the dual-chamber device would have to satisfy both of the following conditions: Base rate, rate hysteresis or rest rate programmed to 55 ppm or lower AutoCapture [™] pacing system programmed ON Additionally, it should be noted that even with the following settings, the shortened pacing interval will not occur unless specific internal timing and other conditions are met. If the device is programmed to any other combination of settings (e.g. base rate, hysteresis rate and rest rate all at 60 ppm or above, or AutoCapture programmed OFF), the scenario described above cannot occur. Abbott recommends that physicians review their records to identify which, if any, of their patients implanted with a subject device have programmed settings as described above. For these patients, they should schedule a pacemaker programming session as soon as possible to temporarily program AutoCapture OFF or adjust the base rate, rate hysteresis and rest rate to 60 ppm or above. The root cause of potentially delivering the short pacing interval has been identified and a downloadable firmware correction has been developed to prevent any future occurrence. The revised firmware will be made available to the physicians via a new programmer software version immediately following FDA approval. The revised code will be downloaded automatically during pacer interrogation and will take approximately six seconds. At the time of the FDA approval, all newly manufactured and distributed products will also have the new pacemaker code. There is no need to consider replacement of the devices as the temporary programming outlined above and the forthcoming device firmware update will eliminate the potential for the described phenomenon from occurring in the future. Current Status (December 31, 2017): There have been no implanted devices confirmed to have been affected by this issue since the time of the advisory.

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Tempo [®] (Model 2102) Meta [®] (Model 1256D)	11/04/2002 Class II Dendritic growth at the battery/hybrid connection could cause a short circuit, which in turn could result in premature battery depletion.	Decreased reliability has been observed in a group of these devices. The devices in the advisory population were manufactured at the Teletronics' manufacturing facility between the second quarter of 1996 and the second quarter of 1997. Premature battery depletion resulting from bridging between adjacent connectors has resulted in no output or no telemetry in 1% of this population of devices to date. The following recommendations are made: Physicians are advised to review patients who have 1256D/2102 pacemakers at an early date to evaluate pacemaker function. For patients who are pacemaker dependent , the physician should give consideration to explantation and replacement as soon as practicable. For patients who are not considered pacemaker dependent , the physician should consider replacement only if any abnormal function is identified (such as no output, telemetry loss, telemetry abnormality or premature end of life indication). All remaining non-pacemaker dependent patients with 1256D/2102 pacemakers should subsequently be reviewed at least every six months.

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Tempo [®] (Models 1102, 1902, 2102, 2902) Meta [®] (Model 1256D)	11/04/2002 Class II Dendritic growth at the battery/hybrid connection could cause a short circuit, which in turn could result in premature battery depletion.	This Advisory applies to a well-defined group of Tempo [®] and Meta [®] 1256D pacemakers which have been observed to exhibit premature battery depletion. This is an extension of a previous Tempo [®] /Meta [®] advisory dated 6/6/00. The following recommendations are made: For patients who are significantly pacemaker dependent . The decision as to whether or not to replace a pacemaker prophylactically remains one of clinical judgment. We therefore recommend that you strongly consider replacing these pacemakers on a prophylactic basis for all patients who are significantly pacemaker dependent, unless pacemaker replacement is medically contraindicated. For patients who are not considered pacemaker dependent . Again, this is a matter of clinical judgment. In the event a device is not replaced, continued routine monitoring is advised.

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Meta [®] DDDR (Model 1256)	6/6/2000 Class II Integrated circuit failure due to electrostatic discharge during manufacturing resulting in no output or sensing anomalies.	This Advisory applies to a well-defined group of Meta [®] 1256 pacemakers, which have been observed to exhibit decreased reliability. The following recommendations are made: For patients who are significantly pacemaker dependent . The decision as to whether or not to replace a pacemaker prophylactically remains one of clinical judgment. We therefore recommend that you strongly consider replacing these pacemakers on a prophylactic basis for all patients who are significantly pacemaker dependent, unless pacemaker replacement is medically contraindicated. For patients who are not considered pacemaker dependent . Again, this is a matter of clinical judgment. In the event a device is not replaced, continued routine monitoring is advised.

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Tempo [™] (Models 1102, 1902, 2102, 2902) Meta [™] (Model 1256D)	6/6/2000 Class II Dendritic growth at the battery/hybrid connection could cause a short circuit, which in turn could result in premature battery depletion.	This Advisory applies to a well-defined group of Tempo [™] and Meta [™] 1256D pacemakers, which have been observed to exhibit premature battery depletion. The following recommendations are made: For patients who are significantly pacemaker dependent. The decision as to whether or not to replace a pacemaker prophylactically remains one of clinical judgment. We therefore recommend that you strongly consider replacing these pacemakers on a prophylactic basis for all patients who are significantly pacemaker dependent, unless pacemaker replacement is medically contraindicated. For patients who are not considered pacemaker dependent. Again, this is a matter of clinical judgment. In the event a device is not replaced, continued routine monitoring is advised.

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Trilogy [™] (Models 2250L, 2260L, 2264L, 2308L, 2318L, 2350L, 2360L, 2364L)	3/10/2000 Class II Risk of malfunction due to an anomaly of the pacemaker's microprocessor. Typical manifestations of this anomaly are dashes observed on the programmer screen, unexpected rate variation, abnormally high battery current drain, and mode change.	Continued monitoring of Trilogy [™] devices affected by an Advisory issued in July 1999 has revealed an additional low level of adverse events. A sub-population of the above models, principally those manufactured before January 1999, is potentially affected by a risk of malfunction due to an anomaly of the pacemaker's microprocessor. To date this has been reported in only 0.11% of the implanted population. Typical manifestations of this anomaly are: <ul style="list-style-type: none"> Interrogation/programming difficulties, including the presence of dashes (—) on the programmer screen for some parameter values after interrogation Unexpected rate variations Abnormally high battery current drain Mode change The presence of dashes on the programmer screen is typically caused by partial programming of one or more of the programmed parameters and may be corrected by reprogramming. In the event that observed dashes (—) cannot be resolved by reprogramming, please contact your Abbott representative, as it may be possible to resolve the situation non-invasively via special programming and may not be indicative of a microprocessor anomaly. Considering the low level of incidence of this anomaly, the following steps are recommended: <ol style="list-style-type: none"> 1. Assess the patient population, relative to the potential for an inappropriate mode change to single-chamber atrial pacing, to determine which patients are pacemaker-dependent and have an inadequate ventricular escape rhythm. 2. Replacement of a device always remains a matter of clinical judgement, including balancing the clinical risk associated with pacemaker replacement against the risk of device malfunction. 3. Almost all microprocessor malfunctions described in this notification were found during routine follow-up. This reinforces the importance that pacemaker patients should be followed regularly in accordance with best prevailing medical practices.

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Affinity™ (Models 5130L, 5130R, 5330L, 5330R, 5230L, 5230R)	<p>2/14/2000 Class II</p> <p>An unsecured resistor connection to the hybrid circuit might cause abnormal measured data, false RRT indication, backup VVI mode, or intermittent loss of output.</p>	<p>This advisory applies to a very specific, well-defined group of Affinity devices that have been implanted worldwide. The device may exhibit any or all of the following output anomalies:</p> <ul style="list-style-type: none"> Abnormal measured battery data, A false recommended replacement (RRT) indication, Reversion to back-up VVI mode, Intermittent loss of output. One marker of an affected device that may be apparent during a routine follow-up interrogation is an abnormally high reading for battery current (taking into consideration the device's programmed output parameters). <p>If this or an early indication of RRT or reversion to back-up VVI mode is observed, please contact your local Abbott representative or Technical Services. Although the time course is variable, these behavioral anomalies will occur before there is any affect on device output.</p>

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
Trilogy™ (Models 2250L, 2260L, 2264L, 2308L, 2318L, 2350L, 2360L, 2364L)	<p>7/19/1999 Class II</p> <p>Premature battery depletion caused by a current leakage path that could be created during the laser welding process to attach the battery to the device hybrid</p>	<p>Analysis of the clinical data associated with the failed devices has shown that an early increase in battery impedance could be observed at one or more previous routine follow-up visits prior to reaching RRT. The observed rise in battery impedance is an earlier and more reliable indicator than the reported battery voltage. The battery depletion, although accelerated, does not occur abruptly and patients can be monitored using the measured data telemetry in the pacemaker. The following follow-up schedule and device monitoring is advised.</p> <ol style="list-style-type: none"> 1. For patients who have had a follow-up visit at least 12 months after their device was implanted, check the measured data printout from the last follow-up evaluation. If a battery impedance of “< 1 kOhm” was recorded, continue to monitor the battery impedance with your routine follow-up schedule for that patient (6-month intervals recommended). If follow-up visits every 6 months is not your routine schedule, then an additional visit at 18 months should be performed. 2. For patients who have not yet had a follow-up visit at least 12 months after their device was implanted, the following is recommended: <ul style="list-style-type: none"> If the patient has had their device implanted for less than two months or has not had a device interrogation with measured data within the last three months, an evaluation of the battery impedance is recommended as soon as possible. This should be printed and a copy retained in the patient's pacemaker or office chart. Otherwise, if the battery impedance from the most recent evaluation is “< 1 kOhm,” begin an every 3-month follow-up schedule with respect to measured data telemetry until that value has been recorded at least 12 months post-implant. Thereafter, continue to monitor the battery impedance with your routine follow-up schedule for that patient (6-month intervals recommended). If follow-up visits every six months is not your routine schedule, then an additional visit at 18 months should be performed. <p>If the battery impedance reading is 1 kOhm or higher and the pacemaker has been implanted for less than two years, it is likely that the system is demonstrating accelerated battery depletion. Please contact your local Field Representative or Technical Services.</p>

LEFT-HEART LEADS

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>QuickSite™ (Models 1056T, 1058T) QuickFlex™ (Models 1156T, 1158T)</p>	<p>4/3/2012 Class II</p> <p>Abrasion of the silicone insulation in the distal portion of QuickSite and QuickFlex leads has led to visual observations of externalized conductors.</p> <p>There have been no reports of death or serious injury associated with the externalized conductors; likewise there have been no electrical dysfunctions attributable to the externalized conductors.</p> <p>The reported rate of externalized conductors in the QuickSite and QuickFlex leads is 0.023%, based on 39 confirmed cases of externalized conductors in a population of approximately 82,000 QuickSite and 89,000 QuickFlex leads sold worldwide.</p> <p>This issue is under-detected because these cases are visual observations without any signs of electrical dysfunction and fluoroscopic/xray imaging is not routine. Based on a review of returned leads and available fluoroscopic and x-ray images of patients with QuickSite and QuickFlex leads (1,219 leads), it is estimated that the incidence of conductor externalization on these leads may be 3% to 4%.</p>	<p>Abbott and its Medical Advisory Board recommend that physicians continue to monitor their patient's implanted system at regularly scheduled intervals with attention paid to diagnostic information related to LV pacing performance, in particular LV lead impedance and capture thresholds. Programming of alerts that monitor lead impedance changes outside of the nominal range and enabling the patient notifier should be considered. A special X-ray or fluoroscopic imaging is not recommended for LV CRT leads with normal electrical function. CRT pacing functionality should be evaluated during routine device checks and only leads exhibiting electrical anomalies that cannot be reprogrammed to deliver effective CRT pacing should be considered for replacement.</p> <p>Current Status (December 31, 2017): At the time of the advisory there was a worldwide reported externalized conductor rate of 0.023% in QuickSite and QuickFlex leads. As of December 31, 2017, the worldwide reported externalized conductor rate for QuickSite and QuickFlex leads was 0.25%.</p>

DEFIBRILLATION LEADS

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Optisure™ Defibrillation Lead (Models LDA220, LDA220Q, LDA230Q, LDP220Q)</p>	<p>11/3/2015 Class I</p> <p>A limited number of dual coil Optisure defibrillation leads may have been compromised during the manufacturing process. A trim technique to remove excess medical adhesive around the SVC shock coil may have introduced damage to the lead's insulation.</p> <p>A thorough investigation has determined the probability of a lead malfunction as a result of this trim technique is very low. A total of 447 leads subjected to the trim technique were distributed globally. Of those, 278 were implanted in the United States. St. Jude Medical is not aware of any adverse clinical events related to this matter. Furthermore, an analysis of patients implanted with the subject leads that are being actively monitored via Merlin.net™ Patient Care Network has shown that none of these patients have experienced any recorded electrical issues.</p>	<p>Abbott recommends the following actions depending on the device the affected patients have implanted. According to our records the vast majority of patients with the subject leads have devices with the DynamicTx™ feature that provides additional protection to help ensure therapy delivery in the case of a compromised lead.</p> <p>For patients implanted with a potentially-impacted Optisure lead connected to a device WITH DynamicTx™ technology, we recommend:</p> <p>Review the Patient Records:</p> <ol style="list-style-type: none"> 1. Ensure DynamicTx™ technology is programmed "On" 2. Enroll these patients in our Merlin.net™ Patient Care Network 3. Monitor patients as normal, with no additional testing or follow-up needed. <p>For patients implanted with a potentially-impacted Optisure lead connected to a device WITHOUT DynamicTx™ technology we recommend:</p> <ol style="list-style-type: none"> 1. Enroll these patients in our Merlin.net™ Patient Care Network 2. Where clinically appropriate, consider turning off the SVC coil (select RV-to-Can vector) 3. If dual coil shocking configuration is desired, consider performing a high voltage test using maximum energy. <ol style="list-style-type: none"> a. If shock delivery is normal - no additional testing is required b. If shock delivery identifies a short circuit – consider lead replacement <ul style="list-style-type: none"> • DynamicTx™ technology automatically adjusts shock configurations to ensure the delivery of high-voltage therapy even if an electrical short were to occur. <p>We recommend at your patient's next follow-up visit a St. Jude Medical representative be present to program an alert message into the implanted device. This will provide clinicians following patients with impacted subject lead an alert message on the Merlin™ Programmer upon interrogation, ensuring that future caregivers assessing the diagnostics of these devices receive the latest information and be made aware of this corrective action. We believe such actions will further the ability of our clinician partners to most optimally manage the care of their patients.</p>

DEFIBRILLATION LEADS

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Riata[™] Defibrillation Lead (Models 1570, 1571, 1572, 1580, 1581, 1582) Riata[™] i Defibrillation Lead (Models 1560, 1561, 1562, 1590, 1591, 1592) Riata[™] ST Defibrillation Lead (Models 7000, 7001, 7002, 7010, 7011, 7040, 7041, 7042)</p>	<p>11/28/2011 Class I</p> <p>Externalized conductors occur when an abrasion results in an outer insulation breach, allowing the normally contained conductors to become visible outside the lead body. Even though causality cannot be established, when externalized conductors are accompanied by reports of electrical malfunction, these reports typically include pacing or defibrillation impedance changes, inappropriate therapy, noise and oversensing, and pacing threshold rise. Externalized conductors have not been observed in Riata ST Optim[™] and Durata[™] models due to the presence of an abrasion resistant outer Optim[™] lead insulation sheath.</p> <p>A summary of the types and incidence rates of Riata lead abrasion malfunctions is presented on pages 308-316 of this Product Performance Report.</p>	<p>Abbott and its Medical Advisory Board (MAB) make the following recommendations, which are consistent with standard best practices and our December 2010 product communication.</p> <p>Whenever possible, monitor devices and leads remotely and advise your patients of the importance of contacting you should they experience any adverse events. Abbott[™] remote monitoring features can be used to detect electrical changes early that may be associated with externalized conductors.</p> <p>Continue to monitor your patient's implanted system at regularly scheduled intervals with particular attention to diagnostic information related to defibrillation lead performance. The recommendations for frequency of in-person are a follow-up period of every 3 - 6 months for ICD/CRT-D devices per the HRS/EHRA consensus.</p> <p>Review lead measurements including pacing and high voltage lead impedances per your standard follow-up procedures in particular looking for significant changes from the patient's previous follow-up visits.</p> <p>If there is evidence of a lead electrical failure, manage the patient per standard practice.¹ This may include x-ray or fluoroscopy. Additional testing, if necessary, could include provocative methods such as shoulder and arm movements and deep respiration while looking at the surface ECG and intracardiac electrograms with the programmer, which may reveal an intermittent problem associated with any source of lead electrical failure if one exists.</p> <p>The value of routine x-ray or fluoroscopy for patients with leads having no electrical abnormalities is unknown at this time and is therefore not recommended.</p> <p>In addition, prophylactic explant or replacement of a lead without electrical dysfunction is not recommended.</p> <p>Currently there is no expert consensus regarding whether patients undergoing pulse generator replacement should undergo fluoroscopy or lead replacement should an externalized conductor without electrical anomalies be present. This is, in part, because the risk versus benefit of replacing a lead in such a patient may vary from patient to patient and center to center. Clinical decisions in this setting should be individualized based on specific patient conditions and circumstances.</p> <p>Based on input from the MAB, Abbott is conducting a prospective study to evaluate further the incidence of externalized conductors and the long-term performance of leads with externalized conductors that do not exhibit electrical abnormalities.</p> <p>Current Status (February 28, 2018): At the time of the advisory there was a worldwide reported all-cause insulation abrasion rate of 0.63% for Riata silicone leads. The worldwide reported rate for the subcategory of externalized conductors in Riata silicone leads was 0.10%. As of February 28, 2018, there have been additional reports. The worldwide reported rates of all-cause abrasion and externalized conductors for Riata silicone leads was 4.44% and 2.69% respectively. The latest information related to the silicone Riata lead advisory, including references to independent studies of Riata lead performance, can be obtained at www.RiataCommunication.com.</p> <p>¹ Epstein, A.E. "Troubleshooting of Implantable Cardioverter-Defibrillators." <i>Clinical Cardiac Pacing, Defibrillation, and Resynchronization Therapy, 4th ed.</i> Eds. Ellenbogen, K.A., Kay G.N., Lau, C-P., Wilkoff, B.L. Philadelphia: Elsevier, 2011. 889-910.</p>

DEFIBRILLATION LEADS

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Riata[™] Defibrillation Lead (Models 1570, 1571, 1572, 1580, 1581, 1582) Riata[™] i Defibrillation Lead (Models 1560, 1561, 1562, 1590, 1591, 1592) Riata[™] ST Defibrillation Lead (Models 7000, 7001, 7002, 7010, 7011, 7040, 7041, 7042)</p>	<p>12/15/2010 Outside US Only</p> <p>Abrasion of silicone defibrillation leads is acknowledged within the clinical community as a well known clinical risk and is documented in the literature as the number one cause of lead failure across the industry with reported failure rates ranging from 3 to 10%. After more than 9 years of clinical use and approximately 227,000 implants, silicone insulated Riata[™], Riata[™] i, and Riata[™] ST defibrillation leads have exhibited an insulation abrasion rate of 0.47% (inclusive of confirmed returns and complaints/observations with no associated return). There are several factors that can contribute to lead abrasion in implanted pacing and defibrillation systems, including physiological stresses placed on the lead due to patient anatomy, implant orientation, and mechanical stresses applied from concomitant devices in the body.</p> <p>A summary of the types and incidence rates of Riata lead abrasion malfunctions is presented on pages 308-316 of this Product Performance Report.</p>	<p>Continue to monitor your patient's implanted system at regularly scheduled intervals with particular attention to diagnostic information related to defibrillation lead performance. The recommendations for frequency of in-person or remote monitoring are a follow-up period of every 3 - 6 months for ICD/CRT-D devices per the HRS/EHRA consensus.</p> <p>Review lead measurements including pacing and high voltage lead impedances per your standard follow-up procedure in particular looking for significant changes from the patient's previous follow-up visits.</p> <p>If there is evidence of a lead electrical failure, manage the patient per standard practice.¹ This may include x-ray or fluoroscopy. Additional testing if necessary could include provocative methods such as shoulder and arm movements and deep respiration while looking at the surface ECG and intracardiac electrograms with the programmer, which may reveal an intermittent problem if one exists.</p> <p>Consider remote monitoring and advise your patients of the importance of contacting you if they experience any adverse events.</p> <p>Current Status (February 28, 2018): At the time of the advisory there was a worldwide insulation abrasion rate of 0.47% for Riata silicone leads. As of February 28, 2018, there have been additional reports and the worldwide reported insulation abrasion rate is 4.44%.</p>

¹ Epstein, A.E. "Troubleshooting of Implantable Cardioverter-Defibrillators." *Clinical Cardiac Pacing, Defibrillation, and Resynchronization Therapy, 4th ed.* Eds. Ellenbogen, K.A., Kay G.N., Lau, C-P., Wilkoff, B.L. Philadelphia: Elsevier, 2011. 889-910.

ICM DEVICES

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>SJM Confirm™ ICM (Models DM2100, DM2102)</p>	<p>3/11/2011 Class II US and Germany</p> <p>A product firmware upgrade using the Merlin™ Patient Care System (PCS) programmer running software versions 10.1.1.3, 10.1.1.2, or 11.2.2 leaves the implantable cardiac monitor device in a state which results in increased current usage. If not corrected this state could result in premature battery depletion.</p>	<p>If you are following any patients implanted with the SJM Confirm ICM Models DM2100 or DM2102 and their device was upgraded using the Merlin programmer with the above mentioned software versions it is recommended to determine the patient's clinical reason for the implant and their continued need for the device:</p> <ul style="list-style-type: none"> • If the device has previously been used to record and assist in the diagnosis of an arrhythmia and is no longer needed, no further action is required. The device will exhaust its battery capacity prior to the 3 year expected longevity. • If the unit is still indicated for diagnosing a potential clinical arrhythmia, contact your Field Clinical Engineer and he/she will assist in calculating the projected remaining longevity. If appropriate, the microprocessor operation can be reset to the nominal current drain. • If the device is no longer indicated it can be left implanted until such time that a routine explant is desired. <p>If the device is determined to be necessary and is experiencing increased current usage as described above, it can be corrected with assistance from your Sales Representative or St. Jude Medical Technical Services.</p> <p>St. Jude Medical is in the process of developing new Merlin PCS programmer software that will properly upgrade SJM Confirm devices.</p> <p>Current Status (December 31, 2017): At the time of the advisory, 83 implanted devices world-wide were identified as having undergone the problematic firmware upgrade. All of these devices have been corrected using the Merlin PCS programmer or were determined by the clinician to not require further action because the device had already provided the necessary diagnostic information and was no longer required. There have been no additional implanted devices confirmed to have been affected by this issue. Updated Merlin PCS programmer software has been implemented which prevents this issue from occurring in the future.</p>

REMOTE MONITORING/TRANSMITTERS

MODEL IDENTIFICATION	ADVISORY	FOLLOW-UP RECOMMENDATIONS AT TIME OF ADVISORY
<p>Merlin@home™ RF Remote Monitoring Transmitter EX1150</p>	<p>12/18/2014 Class II</p> <p>A low incidence of Merlin@home transmitters initiating a software reset resulting in backup operation in some implanted Abbott Radio Frequency (RF) enabled Implantable Cardioverter Defibrillators (ICDs) and Pacemakers. Potentially affected RF devices include the Abbott Ellipse™, Fortify Assura™, Unify Assura™, and Quadra Assura™ ICDs and Assurity™ and Allure™ Pacemakers.</p> <p>In the event that an Ellipse, Fortify Assura, Unify Assura, or Quadra Assura ICD enters backup mode, the nominal operational settings will be VVI pacing mode, 67 ppm, 5.0v/0.6ms with bipolar pacing output and defibrillation settings of a VF detection rate of 146 bpm and 36J high voltage therapy. In the event an Assurity or Allure pacemaker enters backup mode, it will have output settings of VVI pacing mode, 67 ppm, 5.0v/0.6ms with unipolar pacing. This issue can only occur when the patient is being actively monitored by a Merlin@home™ RF bedside transmitter. If a device enters backup mode, the Merlin@home system will detect it and an alert will be provided to the clinic. Additionally, the ICD will deliver a patient vibratory alert and the pacemaker will deliver a patient audible alert.</p> <p>For Ellipse, Fortify Assura, Unify Assura and Quadra Assura ICDs, the rate of occurrence is 0.25% based on 55,000 devices followed via Merlin.net™ remote monitoring. For Assurity and Allure pacemakers, the rate of occurrence is 0.016% based on 12,000 devices followed remotely. All pacemakers and the vast majority (approximately 90%) of ICDs reported to exhibit backup operation as a result of this anomaly were non-invasively restored to normal operation. In approximately 10% of the ICD cases, software was unable to be successfully restored and a device replacement was performed. The software download procedure was revised to ensure a successful download if an incident of a software reset were to occur in the future.</p> <p>There have been no reported cases of serious injury or death associated with this anomaly. The issue will be resolved with a software update of the Merlin@home transmitter that will be performed automatically over its telephone, broadband, or cellular connection without requiring any action from you or your patients.</p> <p>9/19/2015 An additional software upgrade was implemented to address a second software anomaly coexisted in the Merlin@home transmitter system that also had the potential to cause software resets for potentially affected Abbott devices.</p>	<p>The Merlin@home transmitter software has been modified to prevent this issue from occurring and has received FDA approval. A Merlin@home software update will be performed automatically over its telephone, broadband, or cellular connection without requiring any action from you or your patients. No changes to your patient's remote or in-clinic follow up schedules are required. The process of automatically "uploading" this new version of software to patient transmitters has begun. Patients with implanted devices not mentioned above, patients who are being remotely followed with inductive telemetry (wand directly over the device) and patients not being followed remotely are not affected by this issue.</p> <p>Current Status (December 31, 2017): The worldwide event rate of Merlin@home transmitters initiating a software reset resulting in backup operation for Ellipse, Fortify Assura, Unify Assura and Quadra Assura ICDs, was 0.30% based on 83,000 devices followed via Merlin.net™ Patient Care Network (Merlin™ remote monitoring). For Assurity and Allure pacemakers, the rate of occurrence was 0.06% based on 12,000 devices followed remotely. As of December 31, 2017, there were additional reports and the rate for Ellipse, Fortify Assura, Unify Assura and Quadra Assura ICDs, was 0.60%. For Assurity and Allure pacemakers, the rate of occurrence was 0.14%.</p>

Healthcare Professional Communications

PACEMAKER AND CRT-P DEVICES

MODEL IDENTIFICATION	COMMUNICATION	DETAILS
<p>Affinity[™], Entity[™], Integrity[™], Identity[™], Sustain[™], Frontier[™], Victory[™] and Zephyr[™] models</p>	<p>1/29/2014 Worldwide</p> <p>As part of Abbott's commitment to communications on device performance, and in consultation with our Medical Advisory Board, we provided Health Care Professionals information regarding possible effects of electrocautery on older generation Abbott pacemakers.</p>	<p>Abbott has reviewed incident reports on specific older generation pacemaker models exposed to electrocautery. When devices from these pacemaker families are exposed to electrocautery (as well as the PEAK PlasmaBlade[™] blade), they may exhibit a temporary change in function that could persist for 30 seconds or longer after the electrocautery exposure has been terminated. The duration of the effect depends on several factors including the battery voltage of the device, the energy of the electrocautery output, and the distance from the electrocautery source to the implanted system. The most clinically significant observation has been loss of capture due to a transient reduction in the pacing output voltage. Placing a magnet over the device or programming to an asynchronous pacing mode will not prevent this temporary reduction in pacing output.</p> <p>The effects of electrocautery on cardiac implantable electronic device operation are well documented in the scientific literature and most, if not all, pacemaker and implantable cardioverter defibrillator (ICD) User's Manuals include labeling about the use of electrosurgery equipment and its possible effects on the operational characteristics and/or internal circuitry of these devices.</p> <p>As is the case with all perioperative assessments in patients with cardiac implantable electronic devices, evaluating the individual patient's dependence on the implanted device should be assessed prior to any procedure that would ordinarily require electrocautery, particularly a pacemaker procedure. If pacemaker dependency is identified, either do not use electrocautery or employ appropriate precautions to ensure that the heart rate will be supported in the presence of electrocautery. Consideration of placing a temporary transvenous pacemaker is appropriate.^{1,2}</p> <p>All Abbott pacemaker and ICD User's Manuals provide Warnings and Precautions regarding the use of electrosurgical devices in the vicinity of an implanted device.</p> <p>Importantly, the more recent families of Abbott pacemakers (Accent and Anthem) and all ICDs are not subject to this temporary change in function from the extended effects of electrocautery.</p> <p>References: ¹ Hayes and Friedman, Cardiac Pacing, Defibrillation and Resynchronization, 2nd Edition, p. 192 ² Ellenbogen and Wood, Cardiac Pacing and ICDs, 4th Edition, p. 227</p>

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IMPLANTABLE CARDIAC MONITORS

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Index of Phased-out Models

Phased-out Models

As stated in the introduction of this Product Performance Report, product performance is plotted over a maximum range of 20 years, with a minimum of 500 registered implants required for inclusion in the report. As such, models that no longer meet the criteria for inclusion have been phased-out of the Product Performance Report over time. In order to provide our customers with information on these phased-out models, an index including the final edition for each phased-out model has been included. Previous Product Performance Reports can be viewed on the web at www.SJM.com.

CRT DEVICES

Atlas™ + HF (V-340)
Atlas™ II HF (V-365)
Atlas™ II + HF (V-366)
Epic™ HF (V-337)
Epic™ HF (V-338)
Epic™ II HF (V-355)
Frontier™ (5508)
Promote™ (3107-36)
Promote™ RF (3207-30)

FINAL EDITION

Apr 2011
Dec 2015
Dec 2015
Apr 2011
May 2010
Apr 2011
May 2010
Nov 2010
May 2014

ICDS

Atlas™ DR (V-240)
Atlas™ DR (V-242)
Atlas™ II DR (V-265)
Atlas™ VR (V-199)
Contour™ II (V-185, V-185AC, V-185B, V-185C, V-185D)
Contour™ MD (V-175, V-175AC, V-175B, V-175C, V-175D)
Current™ DR (2107-36)
Current™ DR RF (2207-30)
Current™ VR (1107-36)
Current™ VR (1207-30)
Epic™ + DR (V-236)
Epic™ + DR (V-239)
Epic™ DR (V-233)
Epic™ DR (V-235)
Epic™ II DR (V-255)
Epic™ II DR (V-258)
Epic™ II VR (V-158)
Epic™ + VR (V-196)
Epic™ VR (V-197)
Photon™ DR (V-230HV)

FINAL EDITION

May 2010
Dec 2014
May 2014
Nov 2010
May 2008
May 2010
Nov 2010
Dec 2015
May 2010
Nov 2013
May 2010
May 2014
Apr 2011
Nov 2010
May 2010
Nov 2013
Nov 2013
Dec 2015
Nov 2010
Oct 2007

ICDS

Photon™ μ DR (V-232)
Photon™ μ VR (V-194)
Profile™ (V-186F, V-186HV3)

DEFIBRILLATION LEADS

Riata™ i (1560, 1561)
Riata™ ST Optim™ (7030, 7031)
TVL™ RV (RV01, RV02, RV03, RV06, RV07)
TVL™ SVC (SV01, SV02, SV03)

FINAL EDITION

Oct 2009
May 2010
Oct 2007

FINAL EDITION

Dec 2016
Nov 2013
May 2010
May 2010

PACEMAKERS

AddVent™ (2060)
Affinity™ VDR (5430)
Integrity™ μ SR (5136)
Integrity ADx™ DR (5360)
Integrity ADx™ SR (5160)
Integrity™ μ DR (5336)
Meta™ DDDR (1256)
Meta™ DDDR (1256D)
Paragon™ (2010, 2011, 2012)
Paragon™ II (2016)
Paragon™ III (2304, 2314, 2315)
Phoenix™ II (2005, 2008, 2009)
Phoenix™ III (2204, 2205)
Regency™ SC+ (2400L, 2402L)
Solus™ (2002, 2003)
Solus™ II (2006, 2007)
Synchrony™ II (2022, 2023)
Synchrony™ III (2028, 2029)
Tempo™ D (2902)
Tempo™ DR (2102)
Tempo™ V (1102)

FINAL EDITION

May 2010
May 2010
Nov 2013
Nov 2013
Nov 2013
Nov 2010
Oct 2008
Oct 2008
Nov 2010
Nov 2010
May 2010
Nov 2010
Apr 2009
May 2010
Nov 2010
Nov 2010
Oct 2009
May 2010
Oct 2008
Oct 2008
May 2010

Phased-out Models

PACEMAKERS

Tempo™ VR (1902)
Trilogy™ DC (2308)
Trilogy™ DC+ (2318)
Trilogy™ DR (2350)
Trilogy™ DR+ (2360, 2364)
Trilogy™ SR (2250)
Trilogy™ SR+ (2260, 2264)

PACING LEADS

ACE™ (1015M, 1025M)
Fast-Pass™ (1018T, 1028T)
IsoFlex™ P (1644T)
Passive Plus™ (1135K, 1143K, 1145K, 1235K, 1243K, 1245K)
Passive Plus™ (1136T, 1142T, 1146T, 1222T, 1226T, 1236T, 1242T, 1246T)
Passive Plus™ DX (1343K, 1345K)
Permathane™ ACE (1035M)
Permathane™ ACE (1036T, 1038T)
Tendril™ (1148T, 1188T)
Tendril™ (1188K)
Tendril™ DX (1388K)
Tendril™ DX (1388T, 1388TC)
Unipolar Lead (1007)

FINAL EDITION

May 2010
Oct 2006
Oct 2009
Apr 2007
May 2010
Oct 2009
Nov 2010

FINAL EDITION

Oct 2009
Oct 2009
Apr 2011
May 2010
Dec 2014

May 2010
May 2010
May 2010
Dec 2015
May 2010
May 2010
May 2017
May 2010

Abbott

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SJM.com

St. Jude Medical is now Abbott.

Rx Only

Brief Summary: Prior to using these devices, please review the Instructions for Use for a complete listing indications, contraindications, warnings, precautions, potential adverse events and directions for use.

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