

Image: Construction of the image: Con

Product Performance Report January 2019

Cardiac Rhythm Management Cumulative Survival Probability



Product Performance Report January 2019

Cardiac Rhythm Management Pacemakers ICDs Leads

Contents

	Quality Excellence	4
1.	Terms and Definitions	6
2.	Methodology for Pacemaker and ICD Survival Estimates	9
2.1	Cumulative Survival Probability	10
2.2	Data Acquisition	10
2.3	Returned Product Analysis	10
2.4	Product Performance Graphs and Data	11
3.	Performance of BIOTRONIK Pacemakers	12
3.1	Single-Chamber Pacemakers	13
3.2	Dual-Chamber Pacemakers	21
3.3	CRT Pacemakers	30
4.	Performance of BIOTRONIK ICDs	33
4.1	Single-Chamber ICDs	34
4.2	Dual-Chamber ICDs	41
4.3	CRT ICDs	57
5.	Methodology for Lead Survival Estimates	
	Based on Returned Product Analysis and Complaint Information	67
5.1	Cumulative Lead Survival Probability	68
5.2	Lead Data Acquisition	68
5.3	Returned Product Analysis	69
5.4	Lead Complications	69
5.5	Lead Product Performance Graphs and Data	70
6.	Performance of BIOTRONIK Leads	
	Based on Returned Products and Complaint Data	71
6.1	Pacing Leads	72
6.2	ICD Leads	83
6.2	CRT Leads	99
7.	Methodology for Lead Survival Estimates	
	Based on Clinical Studies	105
7.1	Introduction	106
7.2	BIOTRONIK's Clinical Studies	107
7.3	Lead Complications	109
7.4	Lead Product Performance Graphs and Data	110
8.	Performance of BIOTRONIK Leads Based on Clinical Study Data	111
8.1	Performance of Pacing Leads	112
8.2	Performance of ICD Leads	113
8.3	Performance of CRT Leads	117
9.	Advisories	122
	X-Ray Identifiers for Pacemakers and ICDs	124
	Contacting BIOTRONIK	125

Quality Excellence

BIOTRONIK has a long history of high quality in product design and performance. For more than 50 years, the name BIOTRONIK has been synonymous with excellent workmanship and reliable patient safety. Our quality concept follows an integrated approach and extends from preventative risk measures during a product's development phase through all the steps of the manufacturing and design process.

BIOTRONIK's quality assurance system guarantees strict adherence to internal quality standards as well as compliance with international standards and guidelines. Regular reviews of our product performance and manufacturing evaluations contribute significantly to the achievement of extraordinary quality. Our customers, patients, and physicians can rely on the highest degree of safety built into our products. We always welcome suggestions from users about how we can improve the quality of our products.

This Product Performance Report is an integral component of BIOTRONIK's commitment to provide detailed, accurate information regarding long term reliability. The Product Performance Report exemplifies BIOTRONIK's policy of transparent and timely communication with our customers.

As a means to obtain continuous improvement of the designs, **BIOTRONIK** carefully analyzes returned products and incorporates all findings into our quality assurance system. This Product Performance Report was prepared in accordance with International Standard ISO 5841-2: 2014 (E) and is in compliance with the recommendations from the U.S. Heart Rhythm Society Task Force on Device Performance Policies and Guidelines. As an active member of AdvaMed and their Pacemaker/ICD Working Group, BIOTRONIK has worked extensively with the CRM industry to ensure comparable product performance data is reported by all manufacturers. The data provided in BIOTRONIK's Product Performance Report incorporates the requirements and definitions as defined in AdvaMed's Requirements for Uniform Reporting of Clinical Performance of Pulse Generators, except as noted herein.

In BIOTRONIK's continuous efforts to provide accurate and transparent information and to ensure that a conservative estimate for device performance is reported, the Survival Probability calculations presented herein also consider reported pacemaker and ICD battery depletions as well as lead complications without the device having been returned for analysis.

> 1 The ISO 5841-2:2014(E) is replacing the previous version ISO 5841-2:2000. As part of the update, AdvaMed's Requirements for Uniform Reporting of Clinical Performance of Pulse Generators were incorporated in the new ISO 5841-2:2014(E).

Because a significant portion of this report is based on analyses of returned products, BIOTRONIK urges all physicians to return explanted devices and to notify us when a product is explanted or no longer in use for any reason.

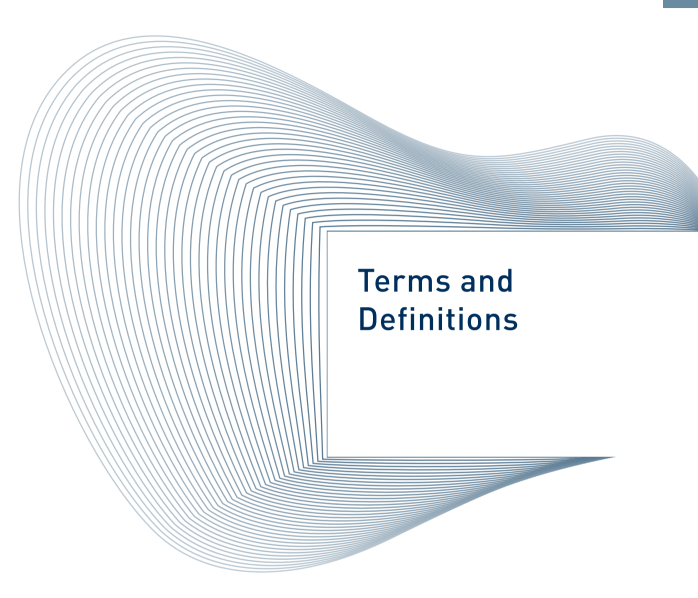
BIOTRONIK aims to continually improve and enhance the scope of this report while integrating the latest information and data concerning the performance of our products. Please contact Advanced Product Support (800) 547-0394 or the PPR Support Team at ppr@biotronik.com with any comments, suggestions or questions regarding this report. Your feedback is highly appreciated and will be used to further develop this report.

BIOTRONIK, January 2019



R. D.M.

Roman Borkowski Senior Vice President Quality Management & Regulatory Affairs CRM BIOTRONIK SE & Co. KG



1. Terms and Definitions

The following terms and definitions are used for Pacemakers and Implantable Cardioverter Defibrillators (ICDs) as well as pacing and ICD leads throughout this Product Performance Report. These definitions form the basis for this Product Performance Report by clearly articulating the status of each device return and product analysis classification.

Elective Replacement Indicator

All active implantable devices that are powered by an internal battery need to be replaced when their battery is depleted. BIOTRONIK pacemakers and ICDs have an Elective Replacement Indicator (ERI) feature aka Recommended Replacement Time (RRT) that notifies the health care provider when the device's battery is nearing the end of its useful life. Display of ERI is BIOTRONIK's recommendation to the user that the battery's present state will require device replacement in the near future. For further details please refer to the corresponding manual.

Battery Depletion

Battery depletions are classified as either normal (expected) or premature. Premature battery depletions are defined as device malfunctions, while normal battery depletions are not device malfunctions. Batteries of returned devices are considered to have depleted normally when (a) a device is returned with no associated complaint and the device has reached its elective replacement indicator(s) with implant time that meets or exceeds the nominal (50 percentile) predicted longevity at default (labeled) settings, or (b) a device is returned and the device has reached its elective replacement indicator(s) with implant time exceeding 75% of the expected longevity using the longevity calculation tool available at time of product introduction, calculated using the device's actual use conditions and settings.

For consistency with previous product performance reports, for ICDs released prior to Lumax and pacemakers released prior to Philos II, batteries of returned devices are considered to have depleted normally if they have reached their elective replacement indicator and testing indicates that the battery and associated circuitry are within specifications.

Out of Specification

Any component or software related event that causes the device's characteristics to not meet predefined performance specifications and requirements while implanted and in service. Returned product analysis that determines a device to be out of specification is considered a device malfunction. Normal battery depletions are not considered device malfunctions. BIOTRONIK defines the requirements and performance specifications for each product.

Device Malfunctions

Any component or software related event that causes the device's characteristics to be out of specification while implanted and in service are considered as device malfunctions. Because it is impossible to verify that a device has malfunctioned without analyzing it, only returned devices can be classified as malfunctions for this report. Each returned lead, ICD and pacemaker is analyzed to determine if it has malfunctioned. If the analysis determines that a pacemaker or ICD failed to meet its specifications while implanted and in service, it is further classified as either a malfunction with compromised therapy or as a malfunction without compromised therapy. Devices damaged during implant, revision or after explant, damaged due to external causes (i.e. electrocautery) or due to failure to follow instructions, warnings or contraindications in its associated

technical manual are not considered malfunctions. Devices damaged due to interaction with other implanted devices (i.e., leads) are also not considered as malfunctions for the purposes of this Product Performance Report.

Malfunctions with Compromised Therapy

The condition when a pacemaker or ICD is found to have malfunctioned 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 critical patientprotective pacing or defibrillation therapy is not available. Examples include: sudden loss of battery voltage; accelerated current drain such that a depleted battery was not detected before loss of therapy; sudden malfunction during a tachycardia or fibrillation event resulting in aborted delivery of therapy; intermittent malfunction where therapy is sporadically unavailable

Malfunctions without Compromised Therapy

The condition when a pacemaker or ICD is found to have malfunctioned in a manner that did not compromise pacing or defibrillation therapy while implanted and in service. Therapy is not compromised as long as critical patient-protective pacing and defibrillation therapies are available as determined through device analysis.

Lead Complications

A lead performance issue where a complaint, associated with at least one of the clinical manifestations listed below, is reported and where the nonreturned lead is:

• Verified by medical records to have been implanted and in-service, and

- Reported to have been removed from service,
- Modified to remedy the malfunction, or
- Left in service based on medical judgment.

Complications for leads implanted greater than 30 days are reported as Qualifying lead complications, whereas complications occurring during the first 30 days are reported as Acute lead observations.

In accordance with the latest AdvaMed guidelines and ISO 5841-2:2014(E), the complications are classified in the following categories:

- Failure to Capture
- Failure to Sense
- Oversensing
- Abnormal Pacing Impedance
- Abnormal Defibrillation Impedance
- Insulation Breach
- Conductor Fracture
- Lead Dislodgement
- Extracardiac Stimulation
- Cardiac Perforation
- Other

Survival Probability Estimates

The probability that a device remains operational during a discrete time interval is defined as survival probability. Survival probability, as presented in this report, is related to device survival only and not survival of the patient. The survival probability estimates in this report are based on BIOTRONIK's analysis of returned products as well as events that are reported to BIOTRONIK (e.g., battery depletions or lead complications).

Cumulative Survival Probability Estimates

The survival probability over a device's service time is the cumulative survival probability. It is calculated from all discrete survival probabilities of previous time intervals. This characteristic is calculated separately for malfunction-free survival and allcause survival (including normal battery depletions). Specific populations that are subject to a safety advisory notification are excluded and shown separately.

Implanted Devices

Only devices remaining implanted for at least one calendar day after the implantation date are considered as implanted. Devices that are removed from the patient on the same calendar day as the implant procedure do not contribute to the survival statistics.

Active Implants

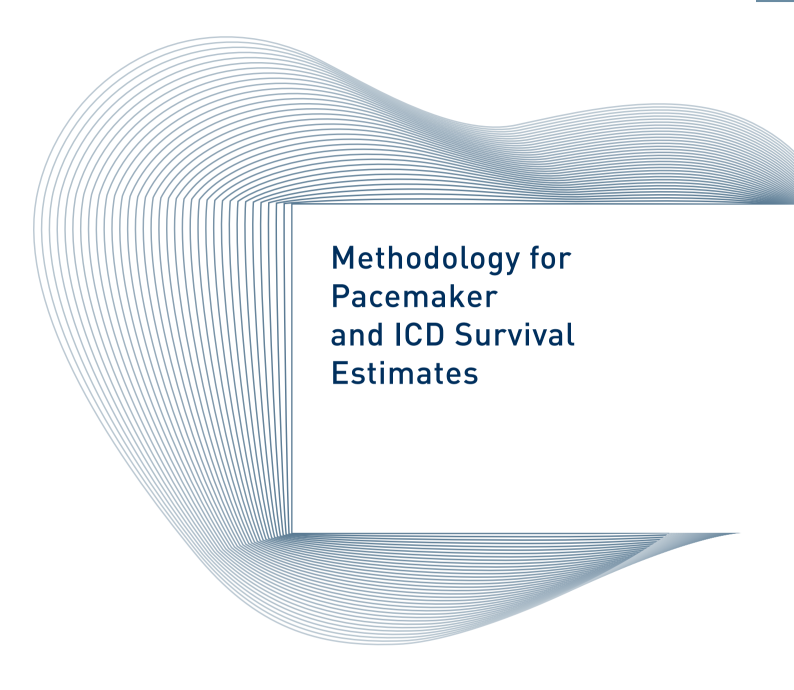
The number of devices that remain operational within a discrete observation interval are active implants. Units are removed from this cohort due to patient death or explant for any reason.

Underreporting

A device status may change without being accounted for in the Product Performance database due to a lack of information being provided to BIOTRONIK. Underreporting adjustments deemed to be necessary are detailed in this report.

Safety Advisory Notifications

Any action taken by the manufacturer to inform clinicians concerning a device performance issue that may cause the device to not meet its predefined specifications is referred to as a Safety Advisory Notification.



2. Methodology for Pacemaker and ICD Survival Estimates

2.1 Cumulative Survival Probability

This report has been prepared in accordance with ISO 5841-2:2014(E) applying actuarial analysis for the calculation of survival probabilities. Survival estimates given in this report are considered to be generally representative for worldwide performance of BIOTRONIK devices.

The Cumulative Survival Probability is an estimate based on the percentage of pacemakers and ICDs that remain implanted and operational at various points of the product's service time in absence of concurrent events such as morbidity and voluntary explants for various reasons (e.g., device upgrade). The Device Survival Estimate over time is displayed in cumulative curves (Kaplan-Meier). The product's performance is evaluated in discrete one-month intervals. The survival probability for each month is given by the number of devices that remain implanted and operational through this month divided by the number of devices that entered the interval. The cumulative survival probability for any period is given by multiplying all survival probabilities of previous months.

At the time of implantation, the cumulative survival probability is 100 %. Even though they are analyzed as part of our quality system monitoring, devices that are found to be out of specification prior to or during the implantation procedure are removed from the statistics as they do not contribute to a patient's risk of being subject to a device malfunction or replacement during the device's service time. Because this report is provided to describe product performance based on returned product analyses, the pacemaker and ICD data does not include information regarding medical complications such as erosion, infection or diaphragmatic stimulation.

In general, during the initial phase of the service time, devices which are out of specification are the primary contribution to reduction of survival probability. As the product lifecycle lengthens, normal battery depletion assumes a greater impact on the survival curve and becomes the dominating factor.

In order to make these two effects distinguishable, the cumulative survival probability curves are shown separately for devices that are confirmed to have malfunctioned only, and for total (allcause) cumulative survival. In case of a device being subject to a safety advisory notification that significantly impacts the survival probability, this factor is displayed separately.

2.2 Data Acquisition

This report is based on the observation of BIOTRONIK's U.S. products through review of our device registration and tracking systems and analyses of returned products from all sources. Because the ability to perform decedent searches of patients with BIOTRONIK devices via the U.S. Social Security Administration, the use of U.S. data more accurately represents the active patient population for reporting purposes. In addition, device tracking regulations and vigilance reporting regulations vary throughout the world; therefore use of the U.S. data is most appropriate for accurate and consistent reporting of product performance.

In order to be included in the population under observation, a device must be registered and implanted for at least one calendar day. The cutoff date for the data included in this report is June 30, 2018. The number of U.S. devices that are implanted and remain active as well as the total number of products distributed worldwide are provided for each product family in this report. Information is provided for separate product families, in that devices with nearly identical hardware and therapy functions are combined. For example, Lexos VR and VR-T (with Home Monitoring) ICDs are combined into a single family, Lexos Single Chamber ICDs.

Survival estimates are calculated for product families having accumulated at least 10,000 cumulative implant months. Because 10,000 implant months may take some time to accumulate, there may be a gap between U.S. market release and the start of graphical representation of survival probability. Products no longer being distributed with less than 500 active implants may be excluded from this report.

ISO 5841-2 describes a method for adjusting the device survival probability to compensate for underreported malfunctions and unrelated patient deaths. The factor for underreporting of malfunctions is unknown as currently no systematic data is available that reveals this factor. Consequently, this factor remains unaccounted for this report. Patient mortality is artificially elevated if the reported rate from our registration and tracking systems is below the annual mortality in clinical studies. Normal battery depletion rate is assumed if the reported rate of depletion decreases over time.

2.3 Returned Product Analysis

Information on malfunctioning for the pacemaker and ICD portions of this report is taken exclusively from the analysis of returned products. The outcome of this analysis is the basis for the final classification of the device's cause of explantation. Only analyzed products with confirmed device malfunctions are utilized in the calculation of malfunction-free survival probability. Every pacemaker and ICD returned to BIOTRONIK is analyzed per internal procedures and classified as functioning normally, normal battery depletion, or malfunctioning (including premature battery depletion) while implanted and in service. These device classifications are the basis for BIOTRONIK's cumulative survival estimates on pacemakers and ICDs.

As a significant portion of pacemakers and ICDs with normal battery depletion are not returned for analysis, BIOTRONIK also considers unconfirmed pacemaker and ICD battery depletions (reported, but device not returned) in the total survival estimates to ensure that a conservative estimate for device performance is reported.

2.4 Product Performance Graphs and Data

The product performance information is shown in each section in alphabetical order and by product type.

For each product, the report provides:

- Product versions that contribute to the evaluation
- U.S. and CE market release dates
- Worldwide quantity of products that have been distributed
- U.S. registered implants (number of products included in this report)
- Estimated active U.S. implants
- Number of U.S. normal battery depletions
- Number of U.S. confirmed malfunctions

The survival plots provide:

1. Total Survival

The combined cumulative survival probability for all causes that result in device removal or a system out of operation, excluding removals for clinical reasons unrelated to the device's performance (i.e., infections).

2. Malfunction-Free Survival

The cumulative survival probability free of component or software malfunctions excluding normal battery depletions, but including premature battery depletions. Normal battery depletions only have an impact on the total cumulative survival.

Products or subgroups of products may become subject to safety advisory notifications that can significantly impact the overall product performance. However, as these subgroups are clearly defined they are separated from the non-advisory devices. The impact of the advisory notification is then shown in a separate graph for total cumulative survival and for malfunction-free survival of the device population affected by the advisory notification. Current advisories are listed in chapter 11 of this report.

The cumulative survival data and the 95% confidence intervals according to the Greenwood's Formula¹ are shown in numerical form for the observed population.

Performance of BIOTRONIK Pacemakers

- 3.1 Single-Chamber Pacemakers
- 3.2 Dual-Chamber Pacemakers
- 3.3 CRT Pacemakers



Cylos and Cylos 990

Product Versions*	_ VR
NBG Codes	_VVIR
US Market Release	_ Jan 2006
CE Market Release	_ Nov 2005 / Mar 2008
Worldwide Distributed Devices	_ 25 900
Registered U.S. Implants	_ 6 150
Estimated Active U.S. Implants	_ 2800
U.S. Normal Battery Depletions	_ 664

	Quantity	Rate
U.S. Confirmed Malfunctions	_ 4	0.07%
Therapy Compromised	_ 1	0.02%
Therapy Available	_ 3	0.05%

- Malfunction-free Cumulative survival propability survival Total survival Years after implant 0 1 2 3 4 5 6 7 8 9 10 Total [%] 100.0 99.9 99.8 99.6 99.5 98.7 94.7 76.1 100.0 86.4 60.2 CI [±%] < 0.1 0.1 0.1 0.2 0.2 0.3 0.7 1.1 1.6 2.5 Malfunction-Free [%] 100.0 100.0 100.0 99.9 99.9 99.9 99.9 99.9 100.0 99.9 99.9 CI [±%] < 0.1 < 0.1 0.1 0.1 0.1 0.1 0.1 0.1 0.1 0.1

* While Cylos 990 VR is not distributed in the U.S., the performance is expected to be similar to the U.S. distributed products

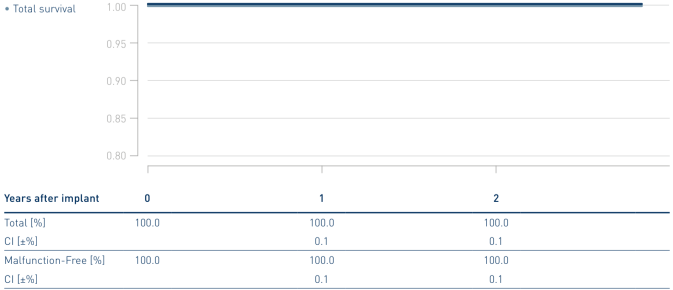
Eluna 8

Product Versions	_SR, SR-T
NBG Codes	_ AAIR, VVIR
US Market Release	_ Dec 2014
CE Market Release	_ Aug 2014
Worldwide Distributed Devices	_ 18800
Registered U.S. Implants	_ 5370
Estimated Active U.S. Implants	4760
U.S. Normal Battery Depletions	_ 0

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

survival



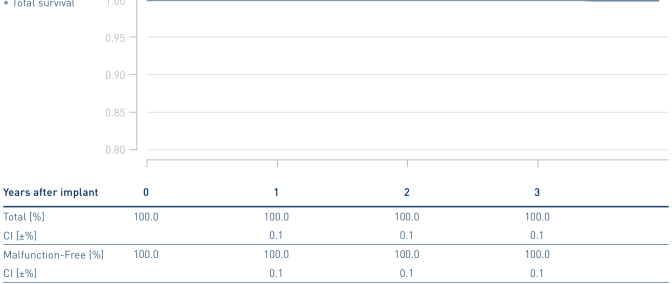
Entovis

Product Versions	SR, SR-T
NBG Codes	AAIR, VVIR
US Market Release	Jun 2010
CE Market Release	Nov 2009
Worldwide Distributed Devices	28 100
Registered U.S. Implants	2400
Estimated Active U.S. Implants	1920
U.S. Normal Battery Depletions	0

	Quantity	Rate
U.S. Confirmed Malfunctions	_0	0.00%
Therapy Compromised	_0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

survival • Total survival

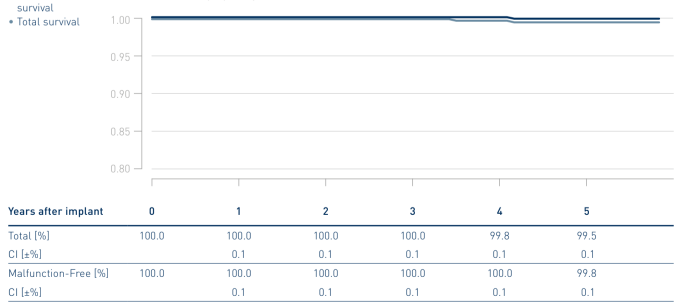


Estella

Product Versions	_SR, SR-T
NBG Codes	_ AAIR, VVIR
US Market Release	_ Feb 2011
CE Market Release	_ Feb 2011
Worldwide Distributed Devices	_ 31 000
Registered U.S. Implants	_ 609
Estimated Active U.S. Implants	421
U.S. Normal Battery Depletions	_ 1

	Quantity	Rate
U.S. Confirmed Malfunctions	. 1	0.16%
Therapy Compromised	0	0.00%
Therapy Available	. 1	0.16%

• Malfunction-free Cumulative survival propability



Etrinsa 8

Product Versions	SR-T
NBG Codes	AAIR, VVIR
US Market Release	Dec 2014
CE Market Release	Aug 2014
Worldwide Distributed Devices	18400
Registered U.S. Implants	1 580
Estimated Active U.S. Implants	1400
U.S. Normal Battery Depletions	_ 1
Estimated Active U.S. Implants	

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	_0	0.00%
Therapy Available	_0	0.00%

• Malfunction-free Cumulative survival propability



CI [±%]		0.1	0.1	
Malfunction-Free [%]	100.0	100.0	100.0	
CI [±%]		0.1	0.1	
Total [%]	100.0	99.9	99.9	
Years after implant	0	1	2	
0.80 —			1	
0.85 —				
0.90 —				
0.00				
0.95				

Evia

Product Versions	SR, SR-T
NBG Codes	AAIR, VVIR
US Market Release	May 2010
CE Market Release	Oct 2009
Worldwide Distributed Devices	59 200
Registered U.S. Implants	12000
Estimated Active U.S. Implants	8650
U.S. Normal Battery Depletions	16

	Quantity	Rate
U.S. Confirmed Malfunctions	. 1	0.01%
Therapy Compromised	. 1	0.01%
Therapy Available	0	0.00%

- Malfunction-free Cumulative survival propability survival • Total survival 0.95 0.85 -0.80 -Т Τ Years after implant 0 1 2 3 4 5 6 7 Total [%] 100.0 100.0 99.9 99.9 99.9 99.9 99.8 99.6 CI [±%] 0.1 0.1 0.1 0.1 0.1 0.1 0.1 Malfunction-Free [%] 100.0 100 100 100 100 100 100 100 CI [±%] 0.1 0.1 0.1 0.1 0.1 0.1 0.1

Philos II and Talos

Product Versions*	_ S, SR
NBG Codes	_SSI, SSIR
US Market Release	_ Sep 2004
CE Market Release	_ Feb 2004 / May 2006
Worldwide Distributed Devices	_ 215 000
Registered U.S. Implants	5240
Estimated Active U.S. Implants	2660
U.S. Normal Battery Depletions	268

	Quantity	Rate
U.S. Confirmed Malfunctions	. 1	0.02%
Therapy Compromised	1	0.02%
Therapy Available	0	0.00%



survival

0	.95 — —												
0	.85 — —												
0	.80 — —												
Years after implant	0	1	2	3	4	5	6	7	8	9	10	11	12
Total [%]	100.0) 100.0	100.0	99.8	99.8	99.4	99.0	98.4	97.1	94.6	90.5	86.7	81.7
CI [±%]		< 0.1	< 0.1	0.1	0.1	0.2	0.3	0.4	0.6	0.9	1.2	1.6	2.2
Malfunction-Free [%]	100.0) 100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100
CI [±%]		< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1

* While Talos SR and Talos S are not distributed in the U.S., their performance is expected to be similar to the U.S. distributed products

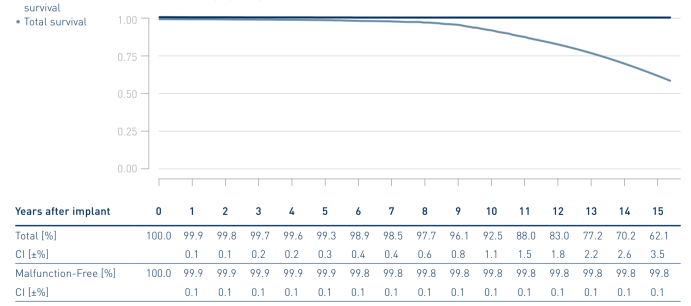
Philos

Product Versions	S, SR
NBG Codes	SSI, SSIR
US Market Release	Sep 2000
CE Market Release	Aug 2000
Worldwide Distributed Devices	109 000
Registered U.S. Implants	5 780
Estimated Active U.S. Implants	1 550
U.S. Normal Battery Depletions	268

	Quantity	Rate
U.S. Confirmed Malfunctions	7	0.12%
Therapy Compromised	0	0.00%
Therapy Available	7	0.12%

Malfunction-free

Cumulative survival propability



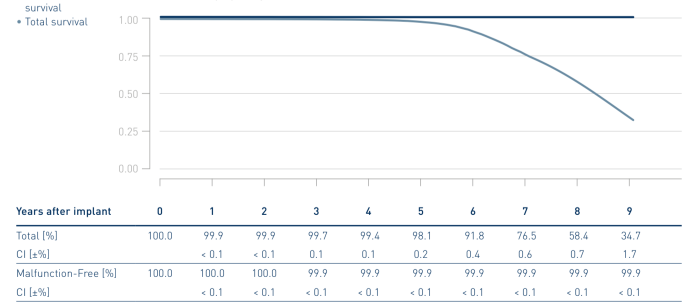
Cylos and Cylos 990

Product Versions*	_ DR, DR-T
NBG Codes	DDDR
US Market Release	Jan 2006
CE Market Release	_ Nov 2005 / Mar 2008
Worldwide Distributed Devices	81 300
Registered U.S. Implants	30400
Estimated Active U.S. Implants	9 0 3 0
U.S. Normal Battery Depletions	7169

	Quantity	Rate
U.S. Confirmed Malfunctions	27	0.09%
Therapy Compromised	7	0.02%
Therapy Available	20	0.07%

Malfunction-free

Cumulative survival propability



*While Cylos 990 DR and Cylos 990 DR-T are not distributed in the U.S., the performance is expected to be similar to the U.S. distributed products

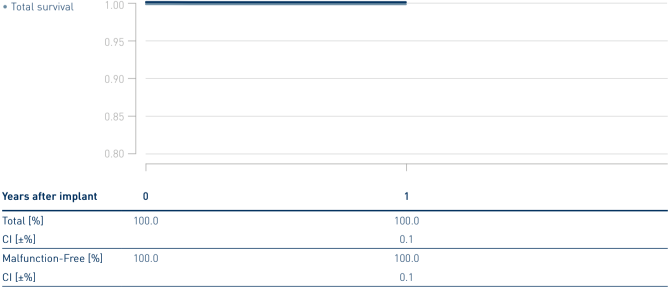
Edora 8

Product Versions	DR, DR-T
NBG Codes	DDDR
US Market Release	May 2017
CE Market Release	Jul 2016
Worldwide Distributed Devices	42000
Registered U.S. Implants	12400
Estimated Active U.S. Implants	12100
U.S. Normal Battery Depletions	_ 0

	Quantity	Rate
U.S. Confirmed Malfunctions	_ 1	0.01%
Therapy Compromised	_ 1	0.01%
Therapy Available	_ 0	0.00%

• Malfunction-free Cumulative survival propability

survival	
Total survival	



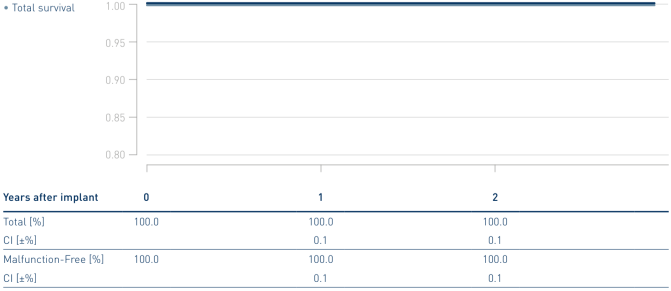
Eluna 8

Product Versions	DR, DR-T
NBG Codes	DDDR
US Market Release	_ Dec 2014
CE Market Release	_ Aug 2014
Worldwide Distributed Devices	_ 88 300
Registered U.S. Implants	_ 37 100
Estimated Active U.S. Implants	_ 33 200
U.S. Normal Battery Depletions	_ 9

	Quantity	Rate
U.S. Confirmed Malfunctions	. 1	0.00%
Therapy Compromised	0	0.00%
Therapy Available	.1	0.00%

• Malfunction-free Cumulative survival propability

survival

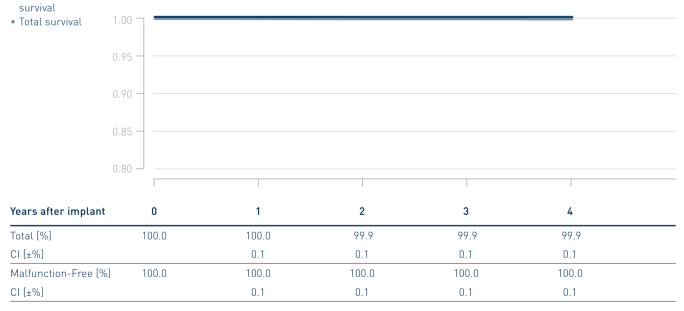


Entovis

Product Versions	DR, DR-T
NBG Codes	DDDR
US Market Release	Feb 2010
CE Market Release	Nov 2009
Worldwide Distributed Devices	106 000
Registered U.S. Implants	12200
Estimated Active U.S. Implants	9900
U.S. Normal Battery Depletions	8

	Quantity	Rate
U.S. Confirmed Malfunctions	2	0.02%
Therapy Compromised	1	0.01%
Therapy Available	1	0.01%

• Malfunction-free Cumulative survival propability



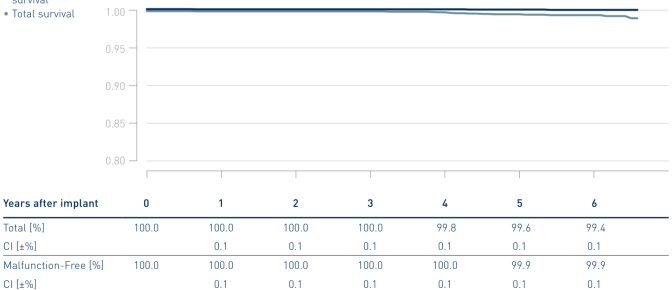
Estella

Product Versions	DR, DR-T
NBG Codes	DDDR
US Market Release	Feb 2011
CE Market Release	Feb 2011
Worldwide Distributed Devices	32 600
Registered U.S. Implants	2950
Estimated Active U.S. Implants	2150
U.S. Normal Battery Depletions	11
	Quantity

	Quantity	Rate
U.S. Confirmed Malfunctions	3	0.10%
Therapy Compromised	0	0.00%
Therapy Available	_ 3	0.10%

• Malfunction-free Cumulative survival propability

survival



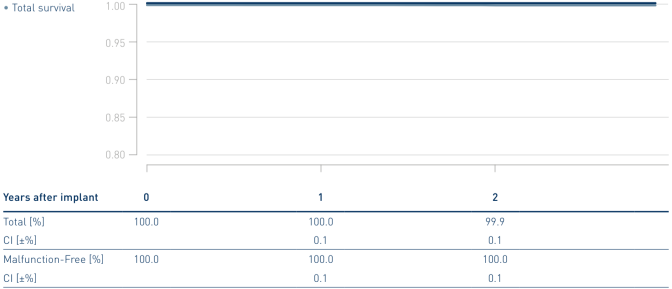
Etrinsa 8

Product Versions	DR-T
NBG Codes	DDDR
US Market Release	Dec 2014
CE Market Release	Aug 2014
Worldwide Distributed Devices	75 200
Registered U.S. Implants	10900
Estimated Active U.S. Implants	9 7 5 0
U.S. Normal Battery Depletions	5

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

survival

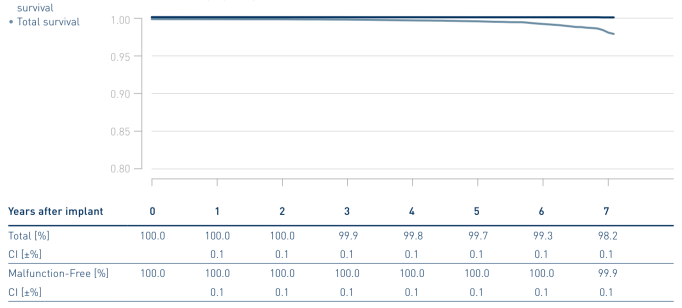


Evia

Product Versions	DR, DR-T
NBG Codes	DDDR
US Market Release	May 2010
CE Market Release	Oct 2009
Worldwide Distributed Devices	203 000
Registered U.S. Implants	61 900
Estimated Active U.S. Implants	45000
U.S. Normal Battery Depletions	239

	Quantity	Rate
U.S. Confirmed Malfunctions	23	0.04%
Therapy Compromised	10	0.02%
Therapy Available	13	0.02%

• Malfunction-free Cumulative survival propability



Philos

Product Versions NBG Codes	_ D, DR, DR-T, SLR _ DDD, DDDR, VDDR
US Market Release	_ Sep 2000
CE Market Release	_ Aug 2000
Worldwide Distributed Devices	_ 172 000
Registered U.S. Implants	_ 20 700
Estimated Active U.S. Implants	_ 5 200
U.S. Normal Battery Depletions	_ 2 583

	Quantity	Rate
U.S. Confirmed Malfunctions	28	0.14%
Therapy Compromised	5	0.02%
Therapy Available	23	0.11%

• Malfunction-free Cumulative survival propability



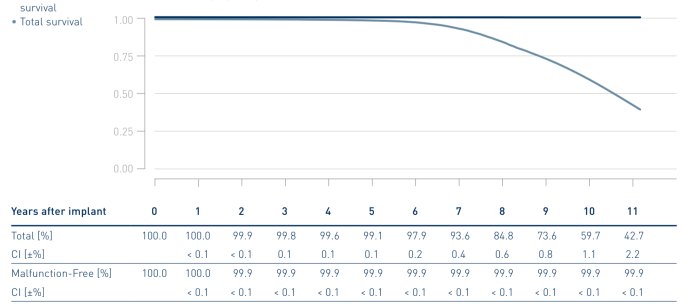
Philos II and Talos

Product Versions* NBG Codes	D, DR, DR-T (Philos II only), SLR DDD, DDDR, VDDR
US Market Release	Sep 2004
CE Market Release	Feb 2004 / May 2006
Worldwide Distributed Devices	372 000
Registered U.S. Implants	23 200
Estimated Active U.S. Implants	8490
U.S. Normal Battery Depletions	3 826

	Quantity	Rate
U.S. Confirmed Malfunctions	21	0.09%
Therapy Compromised	0	0.00%
Therapy Available	21	0.09%

Malfunction-free

Cumulative survival propability



* While Philos II SLR, Talos D, Talos DR and Talos SLR are not distributed in the U.S., their performance is expected to be similar to the U.S. distributed products

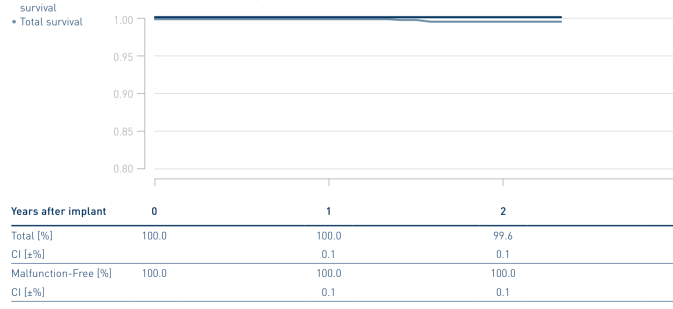
3.3 CRT Pacemakers

Etrinsa 8

Product Versions	HF-T
NBG Codes	DDDRV
US Market Release	_ Dec 2014
CE Market Release	_ Aug 2014
Worldwide Distributed Devices	_ 8410
Registered U.S. Implants	_ 1670
Estimated Active U.S. Implants	_1370
U.S. Normal Battery Depletions	_ 3
0.5. Normal Ballery Depletions	_ 3

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability



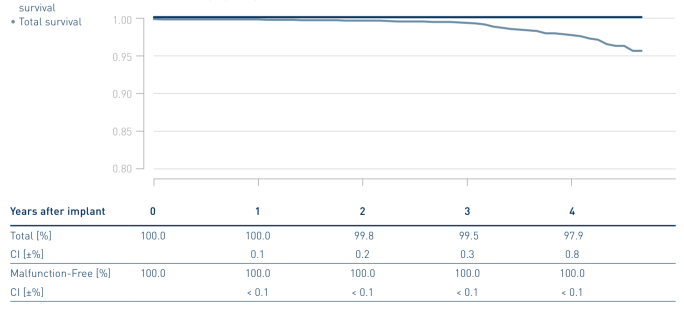
3.3 CRT Pacemakers

Evia

Product Versions	HF, HF-T
NBG Codes	DDDRV
US Market Release	_ May 2010
CE Market Release	Oct 2009
Worldwide Distributed Devices	8830
Registered U.S. Implants	2250
Estimated Active U.S. Implants	1360
U.S. Normal Battery Depletions	_ 46

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability



3.3 CRT Pacemakers

Stratos

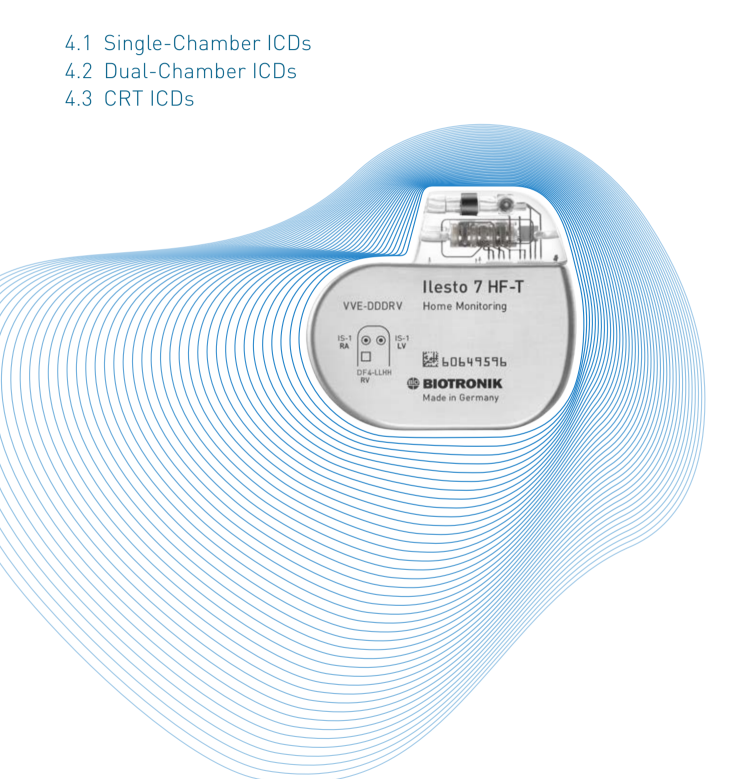
Product Versions	LV, LV-T
NBG Codes	DDDRV
US Market Release	May 2008
CE Market Release	Nov 2002
Worldwide Distributed Devices	21400
Registered U.S. Implants	1310
Estimated Active U.S. Implants	315
U.S. Normal Battery Depletions	254

	Quantity	Rate
U.S. Confirmed Malfunctions	. 1	0.08%
Therapy Compromised	0	0.00%
Therapy Available	1	0.08%

• Malfunction-free Cumulative survival propability

survival • Total survival Т Years after implant 0 1 2 3 4 5 6 Total [%] 100.0 99.9 98.8 96.5 93.4 86.1 72.1 CI [±%] 0.2 0.7 1.2 1.6 2.4 3.4 Malfunction-Free [%] 100.0 99.9 99.9 99.9 99.9 99.9 99.9 CI [±%] 0.2 0.2 0.2 0.2 0.2 0.2

Performance of BIOTRONIK ICDs



4.1 Single-Chamber ICDs

Ilesto 7

Product Versions	VR-T
NBG Codes	VVE-VVIR
Maximum Energy J	40
US Market Release	Sep 2013
CE Market Release	Jun 2013
Worldwide Distributed Devices	2490
Registered U.S. Implants	1 2 7 0
Estimated Active U.S. Implants	1 0 2 0
U.S. Normal Battery Depletions	2

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.08%
Therapy Compromised	0	0.00%
Therapy Available	1	0.08%

• Malfunction-free Cumulative survival propability

_

survival

 Total survival 	1.00				
	0.95 —				
	0.90 —				
	0.85 —				
	0.80				
Years after implar	nt O	1	2	3	
Total [%]	100.0	100.0	99.8	99.7	
CI [±%]		0.1	0.1	0.1	
Malfunction-Free [[%] 100.0	100.0	100.0	99.9	
CI [±%]		0.1	0.1	0.1	

4.1 Single-Chamber ICDs

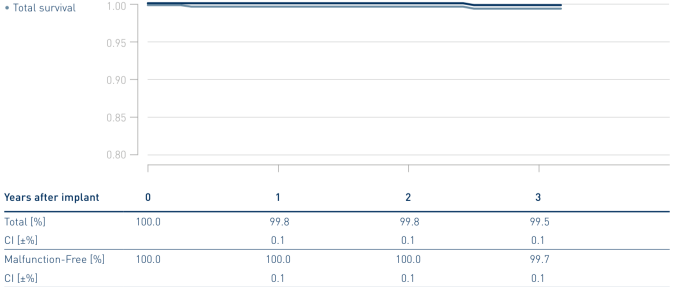
Ilesto 7 DF4

Product Versions	VR-T
NBG Codes	VVE-VVIR
Maximum Energy J	40
US Market Release	Sep 2013
CE Market Release	Jun 2013
Worldwide Distributed Devices	2420
Registered U.S. Implants	466
Estimated Active U.S. Implants	374
U.S. Normal Battery Depletions	1

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.21%
Therapy Compromised	_ 1	0.21%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

survival



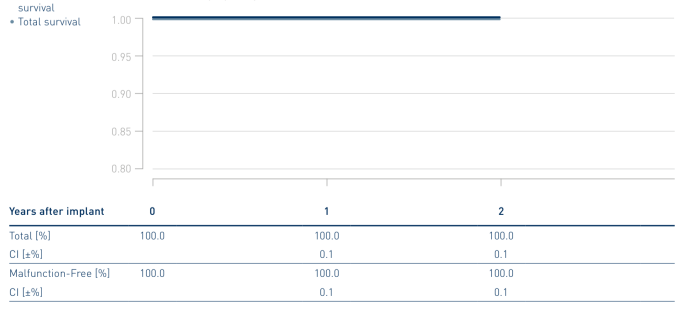
4.1 Single-Chamber ICDs

Itrevia 7

Product Versions	VR-T
NBG Codes	VVE-VVIR
Maximum Energy J	40
US Market Release	Mar 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	1 300
Registered U.S. Implants	607
Estimated Active U.S. Implants	536
U.S. Normal Battery Depletions	0

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

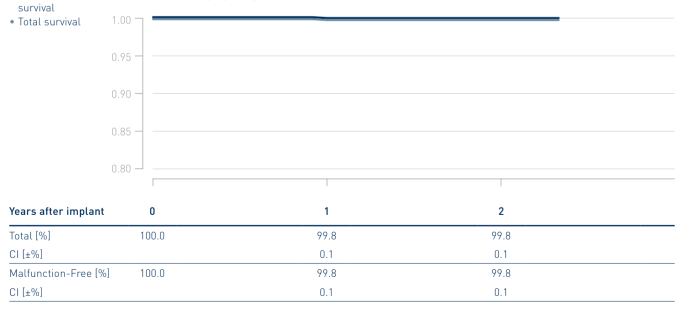


Itrevia 7 DF4

Product Versions	VR-T
NBG Codes	VVE-VVIR
Maximum Energy J	40
US Market Release	Mar 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	1460
Registered U.S. Implants	797
Estimated Active U.S. Implants	686
U.S. Normal Battery Depletions	0

	Quantity	Rate
U.S. Confirmed Malfunctions	. 1	0.13%
Therapy Compromised	0	0.00%
Therapy Available	.1	0.13%

• Malfunction-free Cumulative survival propability



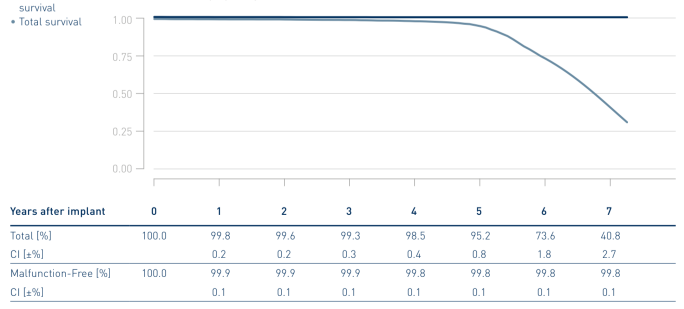
Lumax 340

Product Versions	VR, VR-T
NBG Codes	VVE-VVIR
Maximum Energy J	40
US Market Release	Feb 2007
CE Market Release	Feb 2007
Worldwide Distributed Devices	27 200
Registered U.S. Implants	3990
Estimated Active U.S. Implants	1080
U.S. Normal Battery Depletions	755

	Quantity	Rate
U.S. Confirmed Malfunctions	6	0.15%
Therapy Compromised	4	0.10%
Therapy Available	2	0.05%

• Malfunction-free

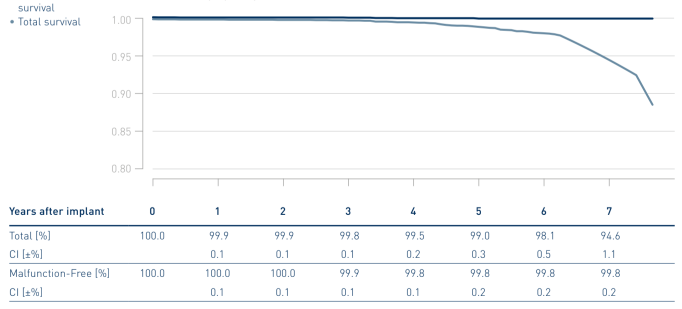
Cumulative survival propability



Product Versions	_VR-T	
NBG Codes	VVE-VVIR	
Maximum Energy J	_ 40	
US Market Release	_ May 2009	
CE Market Release	_ Jun 2008	
Worldwide Distributed Devices	_ 20100	
Registered U.S. Implants	4 5 5 0	
Estimated Active U.S. Implants	_ 2680	
U.S. Normal Battery Depletions	_ 106	
	Quantity	Rate

		·
U.S. Confirmed Malfunctions	8	0.18%
Therapy Compromised	4	0.09%
Therapy Available	4	0.09%

• Malfunction-free Cumulative survival propability



Product Versions	VR-T
NBG Codes	VVE-VVIR
Maximum Energy J	40
US Market Release	Sep 2012
CE Market Release	Apr 2012
Worldwide Distributed Devices	4 8 1 0
Registered U.S. Implants	1 580
Estimated Active U.S. Implants	1120
U.S. Normal Battery Depletions	5

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.06%
Therapy Compromised	_ 1	0.06%
Therapy Available	0	0.00%

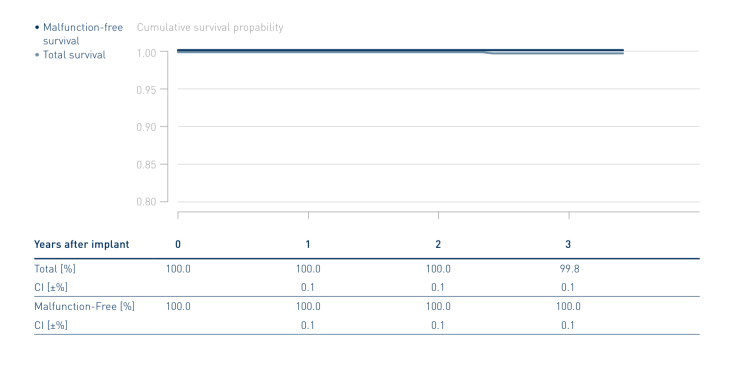
• Malfunction-free Cumulative survival propability

survival • Total survival	1.00						
	0.95 —						
	0.90 —						
	0.85 —						
	0.80						
Years after implant	0	1	2	3	4	5	
Total [%]	100.0	100.0	100.0	100.0	99.9	99.6	
CI [±%]		0.1	0.1	0.1	0.1	0.1	
Malfunction-Free [%	6] 100.0	100.0	100.0	100.0	100.0	100.0	
CI [±%]		0.1	0.1	0.1	0.1	0.1	

Iforia 7

Product Versions	_DR-T
NBG Codes	VVE-VDDR
Maximum Energy J	40
US Market Release	_Sep 2013
CE Market Release	_ Jun 2013
Worldwide Distributed Devices	1090
Registered U.S. Implants	614
Estimated Active U.S. Implants	493
U.S. Normal Battery Depletions	1

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%



Iforia 7 DX

Product Versions	VR-T
NBG Codes	VVE-VDDR
Maximum Energy J	40
US Market Release	Sep 2013
CE Market Release	Jun 2013
Worldwide Distributed Devices	2730
Registered U.S. Implants	1470
Estimated Active U.S. Implants	1210
U.S. Normal Battery Depletions	0

	Quantity	Rate
U.S. Confirmed Malfunctions	2	0.14%
Therapy Compromised	2	0.14%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

survival

survival • Total survival	1.00	_			
	0.95 —				
	0.90 —				
	0.85 —				
	0.80				
Years after impla	nt O	1	2	3	
Total [%]	99.9	99.9	99.9	99.9	
CI [±%]		0.1	0.1	0.1	
Malfunction-Free	[%] 99.9	99.9	99.9	99.9	
CI [±%]		0.1	0.1	0.1	

Ilesto 7

Product Versions NBG Codes	DR-T VVE-DDDR
Maximum Energy J	40
US Market Release	Sep 2013
CE Market Release	Jun 2013
Worldwide Distributed Devices	5140
Registered U.S. Implants	3490
Estimated Active U.S. Implants	2730
U.S. Normal Battery Depletions	3

	Quantity	Rate
U.S. Confirmed Malfunctions	_3	0.09%
Therapy Compromised	_3	0.09%
Therapy Available	_ 0	0.00%

• Malfunction-free Cumulative survival propability

Maturiction-nee survivalTotal survival						
	0.95 —					
	0.90 —					
	0.85 —					
	0.80					
	I	I			I	
Years after implant	0	1	2	3	4	
Total [%]	100.0	100.0	99.9	99.9	99.9	
CI [±%]		0.1	0.1	0.1	0.1	
Malfunction-Free [%	6] 100.0	100.0	100.0	100.0	100.0	
CI [±%]		0.1	0.1	0.1	0.1	

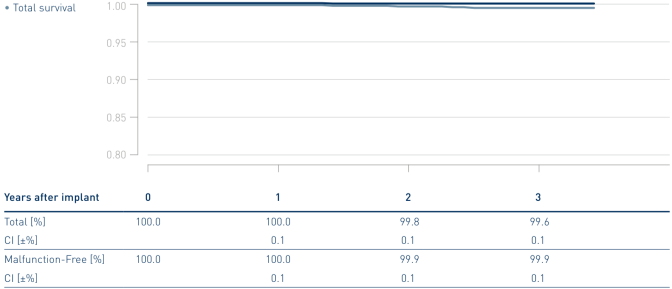
Ilesto 7 DF4

Product Versions	DR-T
NBG Codes	VVE-DDDR
Maximum Energy J	40
US Market Release	Jul 2014
CE Market Release	Jul 2013
Worldwide Distributed Devices	3780
Registered U.S. Implants	1150
Estimated Active U.S. Implants	931
U.S. Normal Battery Depletions	3

	Quantity	Rate
U.S. Confirmed Malfunctions	.1	0.09%
Therapy Compromised	0	0.00%
Therapy Available	.1	0.09%

• Malfunction-free Cumulative survival propability

survival

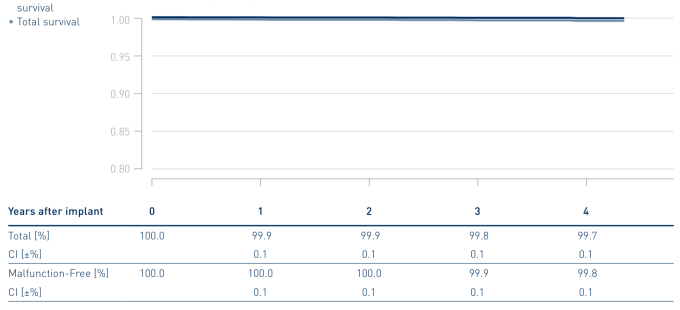


Ilesto 7 DX

Product Versions NBG Codes	VR-T VVE-VDDR 40
Maximum Energy J US Market Release	
CE Market Release	Jun 2013
Worldwide Distributed Devices	6620
Registered U.S. Implants	4720
Estimated Active U.S. Implants	3690
U.S. Normal Battery Depletions	6

	Quantity	Rate
U.S. Confirmed Malfunctions	5	0.11%
Therapy Compromised	3	0.06%
Therapy Available	2	0.04%

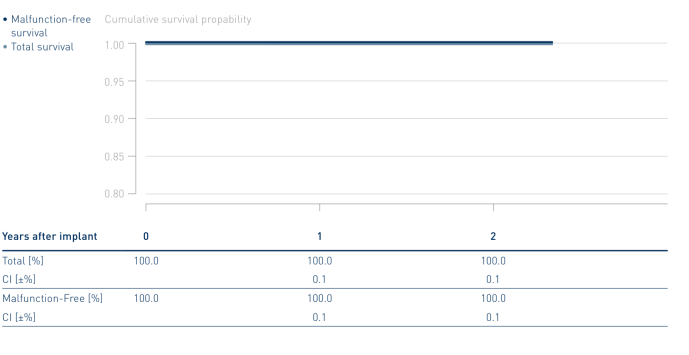
• Malfunction-free Cumulative survival propability



Inventra 7 DX

Product Versions	VR-T
NBG Codes	VVE-VDDR
Maximum Energy J	45
US Market Release	Mar 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	4 4 3 0
Registered U.S. Implants	3760
Estimated Active U.S. Implants	3410
U.S. Normal Battery Depletions	1

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%



Iperia 7

Product Versions	DR-T
NBG Codes	WVE-DDDR
Maximum Energy J	_ 40
US Market Release	_ Dec 2015
CE Market Release	_ Dec 2014
Worldwide Distributed Devices	_ 2 200
Registered U.S. Implants	_1130
Estimated Active U.S. Implants	_1060
U.S. Normal Battery Depletions	_ 0

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

	survival	
•	Total survival	

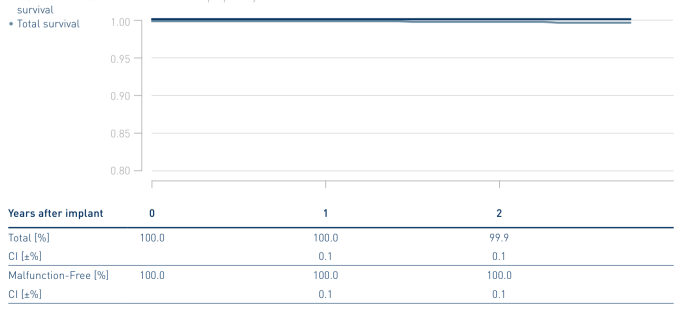


Iperia 7 DF4

Product Versions NBG Codes	_DR-T _VVE-DDDR
Maximum Energy J	40
US Market Release	Dec 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	_ 7 220
Registered U.S. Implants	_3620
Estimated Active U.S. Implants	_3280
U.S. Normal Battery Depletions	_1

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

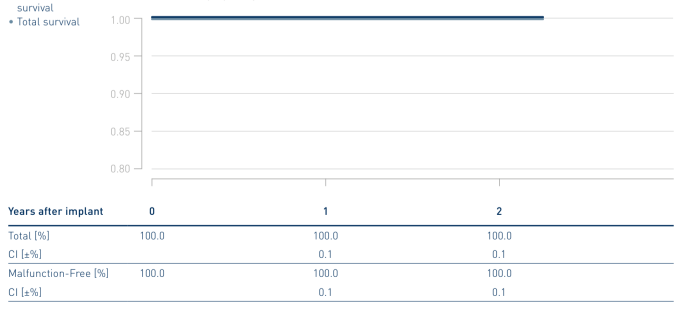


Iperia 7 DX

Product Versions	VR-T
NBG Codes	VVE-VDDR
Maximum Energy J	40
US Market Release	Dec 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	6370
Registered U.S. Implants	4340
Estimated Active U.S. Implants	3990
U.S. Normal Battery Depletions	0

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	_ 0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

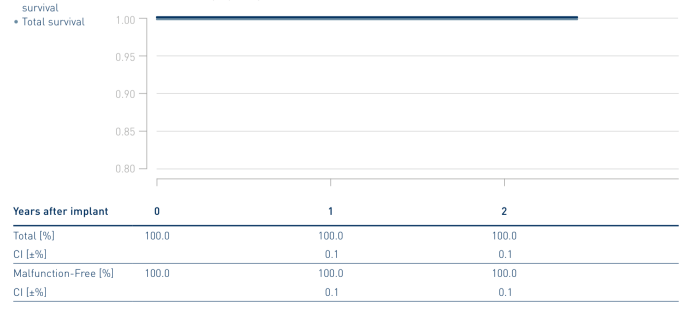


Itrevia 7

Product Versions	_DR-T
NBG Codes	_VVE-DDDR
Maximum Energy J	_ 40
US Market Release	_ Mar 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	2220
Registered U.S. Implants	_1330
Estimated Active U.S. Implants	_ 1 200
U.S. Normal Battery Depletions	_ 0

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

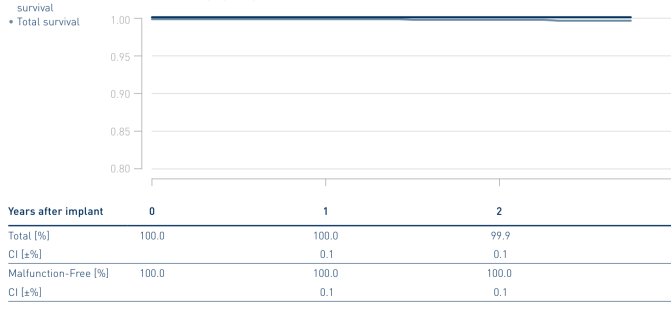


Itrevia 7 DF4

Product Versions NBG Codes	DR-T VVE-DDDR
Maximum Energy J	
US Market Release	40 Mar 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	2360
Registered U.S. Implants	1270
Estimated Active U.S. Implants	1 080
U.S. Normal Battery Depletions	2

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability



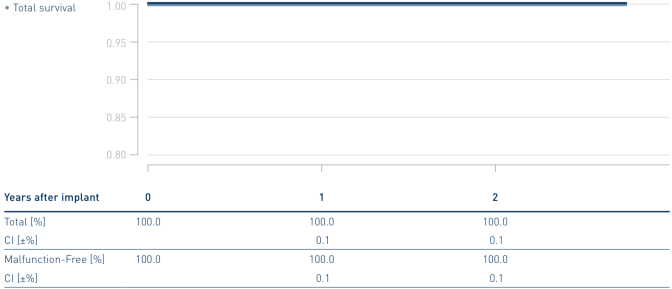
Itrevia 7 DX

Product Versions	_VR-T
NBG Codes	VVE-VDDR
Maximum Energy J	40
US Market Release	_ Mar 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	_ 2790
Registered U.S. Implants	1250
Estimated Active U.S. Implants	_ 1090
U.S. Normal Battery Depletions	_ 0

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

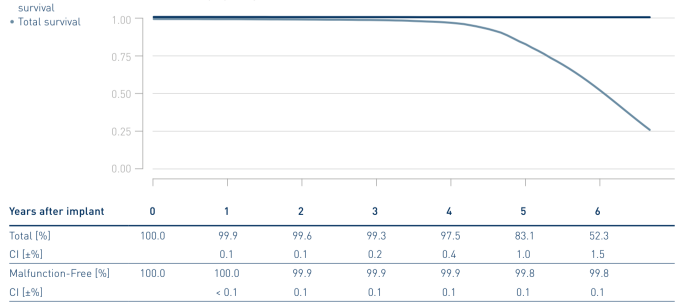
survival



Product Versions	DR, DR-T
NBG Codes	VVE-DDDR
Maximum Energy J	40
US Market Release	Feb 2007
CE Market Release	Feb 2007
Worldwide Distributed Devices	26 500
Registered U.S. Implants	8 2 2 0
Estimated Active U.S. Implants	1950
U.S. Normal Battery Depletions	1862

	Quantity	Rate
U.S. Confirmed Malfunctions	10	0.12%
Therapy Compromised	8	0.10%
Therapy Available	2	0.02%

• Malfunction-free Cumulative survival propability



Lumax 540

Product Versions	_DR-T
NBG Codes	_VVE-DDDR
Maximum Energy J	_ 40
US Market Release	_ May 2009
CE Market Release	_Jun 2008
Worldwide Distributed Devices	_26100
Registered U.S. Implants	_ 11 600
Estimated Active U.S. Implants	_6020
U.S. Normal Battery Depletions	_ 906

	Quantity	Rate
U.S. Confirmed Malfunctions	16	0.14%
Therapy Compromised	8	0.07%
Therapy Available	8	0.07%

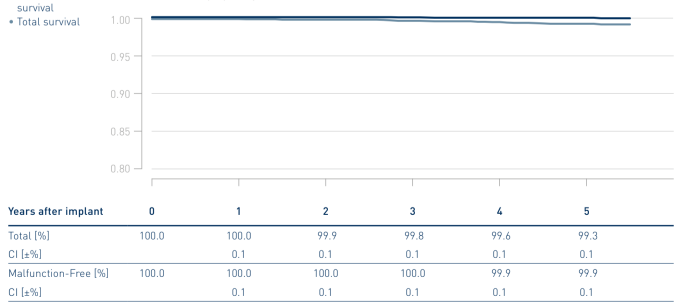
• Malfunction-free Cumulative survival propability

survival Total survival 7 Years after implant 0 1 2 3 4 5 6 Total [%] 100.0 99.9 99.8 99.6 99.4 98.7 97.2 91.8 CI [±%] < 0.1 0.1 0.1 0.2 0.2 0.4 0.8 Malfunction-Free [%] 100.0 100.0 100.0 99.9 99.9 99.8 99.8 99.8 CI [±%] 0.1 0.1 0.1 0.1 0.1 < 0.1 < 0.1

Product Versions	DR-T
NBG Codes	VVE-DDDR
Maximum Energy J	40
US Market Release	Sep 2012
CE Market Release	Apr 2012
Worldwide Distributed Devices	7980
Registered U.S. Implants	3820
Estimated Active U.S. Implants	2750
U.S. Normal Battery Depletions	18

	Quantity	Rate
U.S. Confirmed Malfunctions	4	0.10%
Therapy Compromised	2	0.05%
Therapy Available	2	0.05%

• Malfunction-free Cumulative survival propability



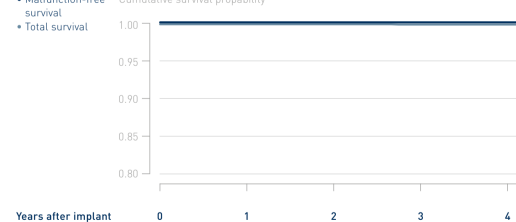
Lumax 740 DX

Product Versions	VR-T
NBG Codes	VVE-VDDR
Maximum Energy J	40
US Market Release	May 2012
CE Market Release	Nov 2011
Worldwide Distributed Devices	4570
Registered U.S. Implants	2230
Estimated Active U.S. Implants	1640
U.S. Normal Battery Depletions	1

	Quantity	Rate
U.S. Confirmed Malfunctions	_ 1	0.04%
Therapy Compromised	_ 1	0.04%
Therapy Available	_ 0	0.00%

• Malfunction-free

Cumu	Ilative	survival	propa	ability



Years after implant	0	1	2	3	4	5
Total [%]	100.0	100.0	100.0	99.9	99.9	99.9
CI [±%]		0.1	0.1	0.1	0.1	0.1
Malfunction-Free [%]	100.0	100.0	100.0	100.0	100.0	99.9
CI [±%]		0.1	0.1	0.1	0.1	0.1

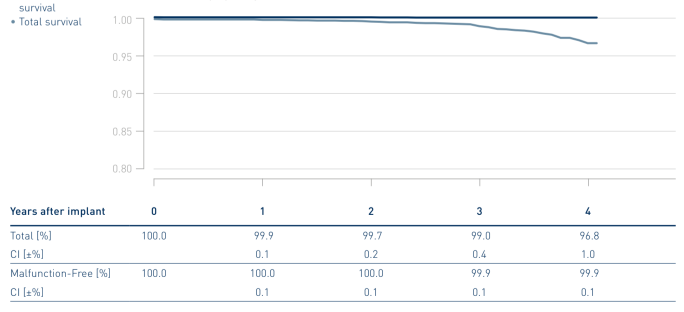
Т

Ilesto 7

Product Versions	HF-T
NBG Codes	VVE-DDDRV
Maximum Energy J	40
US Market Release	Sep 2013
CE Market Release	Jun 2013
Worldwide Distributed Devices	5330
Registered U.S. Implants	3840
Estimated Active U.S. Implants	2620
U.S. Normal Battery Depletions	51

	Quantity	Rate
U.S. Confirmed Malfunctions	3	0.08%
Therapy Compromised	2	0.05%
Therapy Available	1	0.03%

• Malfunction-free Cumulative survival propability



Ilesto 7 DF4

Product Versions	HF-T
NBG Codes	VVE-DDDRV
Maximum Energy J	40
US Market Release	Jul 2014
CE Market Release	Jun 2013
Worldwide Distributed Devices	2400
Registered U.S. Implants	968
Estimated Active U.S. Implants	710
U.S. Normal Battery Depletions	4

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.10%
Therapy Compromised	1	0.10%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability

survival

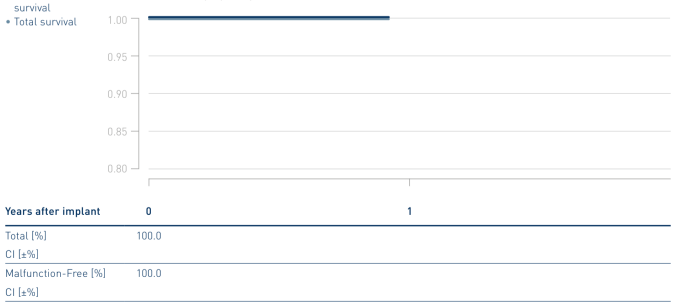
 Total survival 	1.00 -				
	0.95				
	0.70				
	0.90 —				
	0.85 —				
	0.00				
	0.80				
Years after impla	nt O	1	2	3	
	III 0		2	5	
Total [%]	100.0	99.9	99.8	99.6	
CI [±%]		0.1	0.1	0.1	
Malfunction-Free	[%] 100.0	100.0	99.9	99.9	
CI [±%]		0.1	0.1	0.1	
01[1.0]		0.1	0.1	0.1	

Ilivia 7 DF4

Product Versions NBG Codes Maximum Energy J US Market Release CE Market Release Worldwide Distributed Devices Registered U.S. Implants	VDE-DDDRV 40
Registered U.S. Implants	
Estimated Active U.S. Implants U.S. Normal Battery Depletions	

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability



Iperia 7

Product Versions	HF-T
NBG Codes	VDE-DDDRV
Maximum Energy J	_ 40
US Market Release	_ Apr 2016
CE Market Release	Dec 2014
Worldwide Distributed Devices	2710
Registered U.S. Implants	_ 889
Estimated Active U.S. Implants	_ 790
U.S. Normal Battery Depletions	_ 0

	Quantity	Rate
U.S. Confirmed Malfunctions	_ 0	0.00%
Therapy Compromised	_ 0	0.00%
Therapy Available	_ 0	0.00%

• Malfunction-free Cumulative survival propability

survival

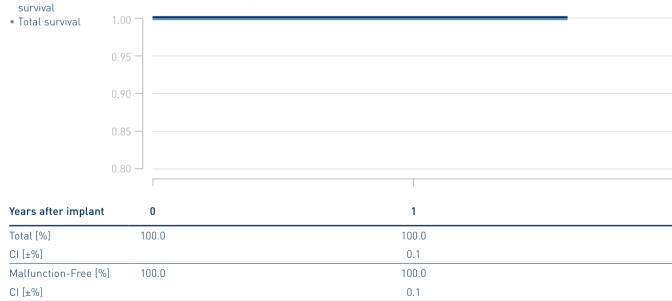
 Total survival 	1.00		
	0.05		
	0.95 -		
	0.90 -		
	0.85 -		
	0.80		
Years after impla	nt O	1	
Total [%]	100.0	100.0	
CI [±%]		0.1	
	[0/] 100.0		
Malfunction-Free	[%] 100.0	100.0	
CI [±%]		0.1	

Iperia 7 DF4

Product Versions	_HF-T
NBG Codes	VDE-DDDRV
Maximum Energy J	_ 40
US Market Release	_Apr 2016
CE Market Release	_Dec 2014
Worldwide Distributed Devices	_5640
Registered U.S. Implants	_1240
Estimated Active U.S. Implants	_1060
U.S. Normal Battery Depletions	_3

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
Therapy Compromised	_ 0	0.00%
Therapy Available	0	0.00%

• Malfunction-free Cumulative survival propability



Itrevia 7

Product Versions	HF-T
NBG Codes	VDE-DDDRV
Maximum Energy J	40
US Market Release	_ Mar 2015
CE Market Release	Dec 2014
Worldwide Distributed Devices	4600
Registered U.S. Implants	3040
Estimated Active U.S. Implants	2480
U.S. Normal Battery Depletions	5

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.03%
Therapy Compromised	0	0.00%
Therapy Available	1	0.03%

• Malfunction-free Cumulative survival propability

survivalTotal survival					
	0.95 -				
	0.90 —				
	0.85 —				
	0.80				
Years after implant	0	1	2		
Total [%]	100.0	99.9	99.8		
CI [±%]		0.1	0.1		
Malfunction-Free [%	b] 100.0	100.0	100.0		
CI [±%]		0.1	0.1		

Itrevia 7 DF4

Product Versions NBG Codes	_HF-T, HF-T QP _VDE-DDDRV
Maximum Energy J	40
US Market Release	Mar 2015
CE Market Release	_Dec 2014
Worldwide Distributed Devices	_ 5800
Registered U.S. Implants	_3210
Estimated Active U.S. Implants	_2560
U.S. Normal Battery Depletions	_ 5
y	

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.03%
Therapy Compromised	0	0.00%
Therapy Available	1	0.03%

• Malfunction-free Cumulative survival propability

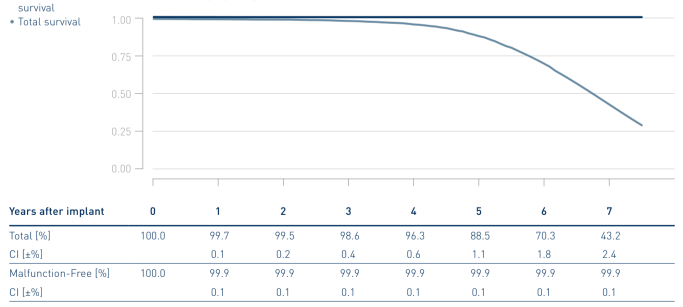
survival

survival • Total survival	1.00			=
	0.95 —			
	0.90 —			
	0.85 —			
	0.80			
Years after impla	int O	1	2	
Total [%]	100.0	99.9	99.8	
CI [±%]		0.1	0.1	
Malfunction-Free	[%] 100.0	100.0	100.0	
CI [±%]		0.1	0.1	

Product Versions	_HF, HF-T VVE-DDDRV	,
NBG Codes Maximum Energy J	40 40	/
US Market Release		
CE Market Release	Dec 2006	
Worldwide Distributed Devices	20 800	
Registered U.S. Implants	_5310	
Estimated Active U.S. Implants	_ 722	
U.S. Normal Battery Depletions	_1086	
	Quantity	Rate

U.S. Confirmed Malfunctions	4	0.08%
Therapy Compromised	2	0.04%
Therapy Available	2	0.04%

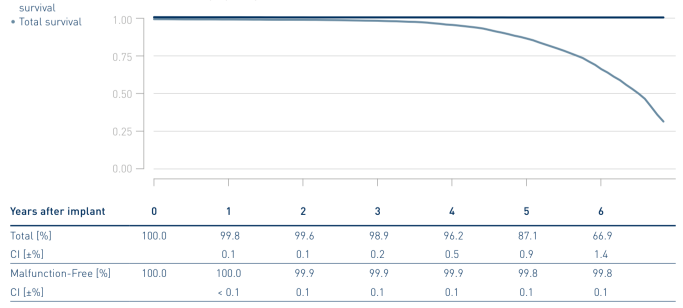
• Malfunction-free Cumulative survival propability



Product Versions	HF-T	
NBG Codes	VVE-DDDRV	
Maximum Energy J	40	
US Market Release	May 2009	
CE Market Release	Jun 2008	
Worldwide Distributed Devices	24 900	
Registered U.S. Implants	8660	
Estimated Active U.S. Implants	1920	
U.S. Normal Battery Depletions	2025	
	Quantity	Rate
II.C. Confirment Malfun attance	11	0 1 0 0/

U.S. Confirmed Malfunctions	11	0.13%
Therapy Compromised	5	0.06%
Therapy Available	6	0.07%

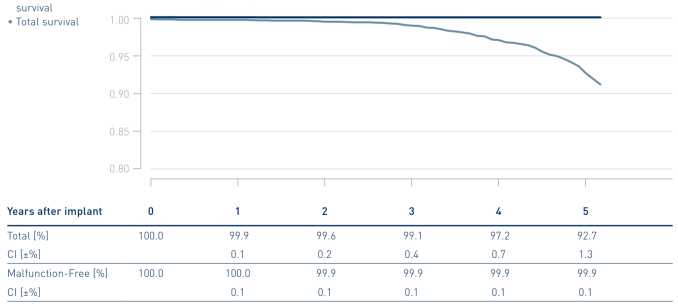
• Malfunction-free Cumulative survival propability



Product Versions	_HF-T
NBG Codes	_VVE-DDDRV
Maximum Energy J	_ 40
US Market Release	_Sep 2012
CE Market Release	_Apr 2012
Worldwide Distributed Devices	_7050
Registered U.S. Implants	_3410
Estimated Active U.S. Implants	_1950
U.S. Normal Battery Depletions	_ 135

	Quantity	Rate
U.S. Confirmed Malfunctions	_ 2	0.06%
Therapy Compromised	_ 0	0.00%
Therapy Available	2	0.06%

• Malfunction-free Cumulative survival propability



Methodology for Lead Survival Estimates Based on Returned Product Analysis and Complaint Information

5. Methodology for Lead Survival Estimates Based on Returned Product Analysis and Complaint Information

5.1 Cumulative Lead Survival Probability

This report has been prepared in accordance with ISO 5841-2:2014(E) applying actuarial analysis for the calculation of lead survival probabilities based on returned product analysis. Lead Survival Estimates given in this report are considered to be generally representative for worldwide performance of BIOTRONIK's pacing and ICD leads.

The Cumulative Survival Probability for leads is an estimate based on the percentage of devices that remain implanted and in service at various points of the product's service time in the absence of concurrent events such as morbidity. The Lead Survival Estimate over time is displayed in cumulative survival curves (Kaplan-Meier). The product's performance is evaluated in discrete one-month intervals. The survival probability for each month is given by the number of leads that remain implanted and active through this month divided by the number of leads that were actively implanted at the start of the interval. The cumulative survival probability for any period is given by multiplying all survival probabilities of previous months.

At the time of implantation, the cumulative lead survival probability is 100 %. Even though they are analyzed as part of our quality system monitoring, leads that are found to be out of specification prior to or during the implantation procedure are removed from the statistics as they do not contribute to a patient's risk of being subject to a device malfunction or replacement during the device's service time. Because this report is provided to communicate information regarding product performance, it does not include data regarding medical complications such as erosion, infection or diaphragmatic stimulation.

Compared to pacemakers and ICDs, a considerable portion of leads with observed or suspected failures are not explanted and returned for laboratory analysis. This is primarily because it is much more difficult and risky to the patient to remove chronically implanted leads. In order to report a conservative measure of lead performance, unconfirmed reports of lead complications are therefore also included in the calculation of a lead's survival probability.

In order to be classified as a qualifying lead complication and thus contributing to the survival probability calculation the same way as a confirmed malfunction, the reported anomaly must have occurred at least 30 days post-implant. Otherwise, factors not related to the lead would likely be the root cause of the observed anomaly, (i.e., patientspecific conditions or implant techniques).

In order to minimize the effect of underreporting of lead malfunctions, BIOTRONIK additionally includes the long term performance post market study data if available.

5.2 Lead Data Acquisition

The following sections of this report provide performance data on BIOTRONIK's pacing and ICD leads and are based on the observation of BIOTRONIK's U.S. products through review of our device registration and tracking systems, and through the analyses of both, returned leads as well as reports of lead complications of non-returned leads. The prospective data from BIOTRONIK's post-approval studies is presented separately in chapters 8 and 9.

In order to be included in the population under observation a lead must be registered and implanted for at least one calendar day. The cut-off date for the data included in this report is December 31, 2017. The sample sizes of U.S. leads that are implanted and remain active as well as the total number of products distributed worldwide are provided for each lead family in this report.

Survival estimates are calculated for lead families having accumulated at least 10,000 cumulative implant months. Products no longer being distributed with less than 500 active implants may be excluded from this report.

ISO 5841-2:2014(E) describes a method for adjusting the device survival probability for underreported malfunctions and unrelated patient deaths that result in an overestimation. of the device's survival probability. The factor for U.S. underreporting of malfunctions of pacing and ICD leads is unknown as currently no systematic data is available that reveals this factor. Consequently, this factor remains unaccounted for in this report. Patient mortality is artificially elevated if the reported rate from our registration and tracking systems is below the annual mortality in clinical studies.

5.3 Returned Product Analysis

Information for the lead sections of this report is taken from the analysis of returned products. The outcome of this analysis is the basis for the final classification of the cause for explantation of the lead. Additionally, reports of lead complications not confirmed by laboratory analysis are taken into consideration. Both, leads with confirmed malfunctions as well as unconfirmed lead complications decrease a lead's total survival probability. Every lead and lead segment returned to BIOTRONIK is analyzed per our internal procedures and classified as within specification, damaged by external causes, or out of specification (malfunction) while implanted and in service.

Those leads found to be out of specification, are divided into the following categories as proposed by AdvaMed and ISO 5841-2:2014(E):

Conductor Fracture

Conductor break with complete or intermittent loss of continuity that could interrupt current flow (e.g. fractured conductors)

Crimps, Welds and Bonds

Any interruption in the conductor or lead body associated with a point of connection

Insulation Breach

Any lead insulation breach

Other

Includes specific proprietary lead mechanical attributes.

5.4 Lead Complications

A considerable portion of leads with observed or suspected failures are not explanted and returned for laboratory analysis. A clinical observation is considered a lead complication if a complaint, associated with at least one of the clinical manifestations listed below, is reported and where the nonreturned lead is:

- Verified by medical records to have been implanted and in-service, and
- Reported to have been removed from service,
- Modified surgically or electrically to remedy the malfunction, or
- Left in service based on medical judgment.

Complications for leads implanted greater than 30 days are reported as Qualifying lead complications, whereas complications occurring during the first 30 days are reported as Acute Lead Observations.

In accordance with the latest AdvaMed guidelines and ISO 5841-2:2014(E) such clinical observations are classified in the following categories:

Failure to Capture

Intermittent or complete failure to achieve cardiac stimulation at programmed output delivered outside of the cardiac refractory period. Sudden and significant increase in the pacing threshold value at which 2:1 safety margin can no longer be achieved.

Failure to Sense

Intermittent or complete loss of sensing or failure to detect intended intrinsic cardiac signals during nonrefractory periods at programmed sensitivity settings

Oversensing

Misinterpretation of cardiac or non-cardiac events as cardiac depolarization

Abnormal Pacing Impedance

Pacing impedance is typically considered abnormal if a measurement is < 200 ohms or > 3000 ohms

Abnormal Defibrillation Impedance

Defibrillation impedance is typically considered abnormal if a measurement is < 20 ohms or > 200 ohms. Including high or low shock impedance when attempting to deliver a shock

Insulation Breach

A disruption or break in lead insulation observed visually, electrically, or radiographically

Conductor Fracture

A mechanical break within the lead conductor 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

Extracardiac Stimulation

Clinical observation of inadvertent nerve/muscle stimulation other than cardiac muscle

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, chest pain, and tamponade

Other

Specific proprietary attributes of a lead such as sensors which affect a lead's ability to perform as designed or remain in service

In order to report a conservative measure of lead performance, qualifying lead complications are also included in the calculation of a lead's survival probability.

Acute Lead Observations may be subject to a number of factors, including patient-specific anatomy, clinical conditions and/or varying implant conditions/techniques. Therefore, acute lead observations are not included in lead survival probability.

5.5 Lead Product Performance Graphs and Data

The lead performance information is shown in each section in alphabetical order and by product name.

For each lead, the report provides:

- Product versions that contribute to the evaluation
- Types of leads
- Polarity
- Steroid
- CE and U.S. market release dates
- Worldwide quantity of products that have been distributed
- U.S. registered implants (number of products included in this report)
- Estimated active U.S. implants
- Number of U.S.qualifying complications
- Number of U.S. acute lead observations
- Number of U.S. confirmed malfunctions
- Number of U.S. leads or partial leads returned post-implant for analysis with a complaint

The survival plots provide:

Total Survival

The cumulative survival probability free of component malfunction or unconfirmed observation of an anomaly. Removals for clinical reasons unrelated to the device's performance (i.e., infections) are excluded.

Products or subgroups of products may become subject to advisory notifications that can significantly impact the overall product performance.

Current advisories are listed in chapter 9 of this report, however to date, BIOTRONIK has never had a pacing or ICD lead safety advisory notification, therefore no summary of lead advisories is provided.

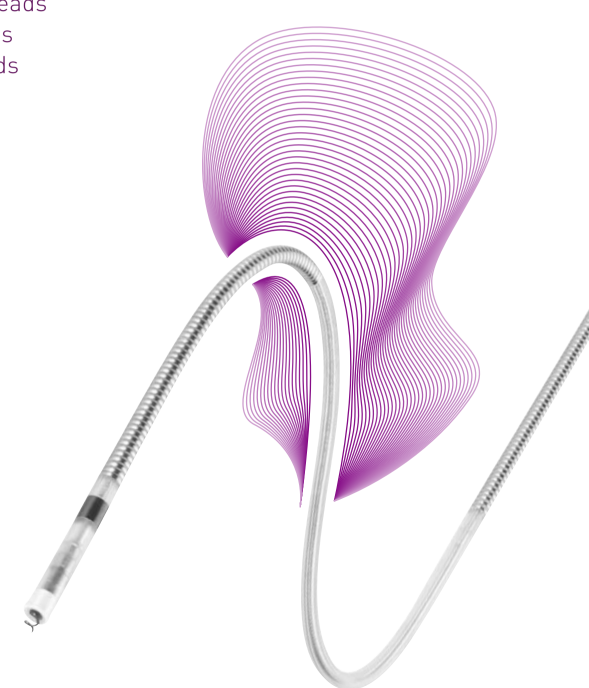
The cumulative survival data and the 95% confidence intervals according to the Greenwood's Formula¹ are shown in numerical form for the observed sample population.

1 Greenwood, M. The natural duration of cancer. Reports on Public Health and Medical Subjects 33, London: Her Majesty's Stationery Office, 1–26, 1926.

Performance of BIOTRONIK Leads

Based on Returned Products and Complaint Data Information

6.1 Pacing Leads6.2 ICD Leads6.3 CRT Leads



6.1 Pacing Leads

Arox

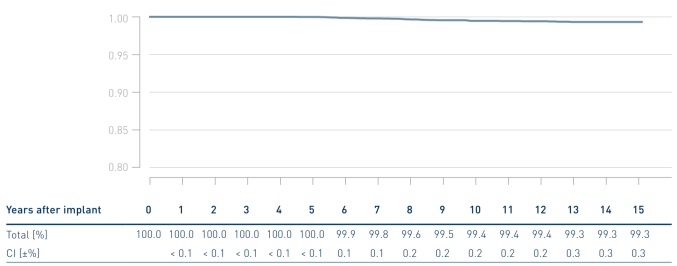
Product Versions	53-BP, 60-BP
Lead Type	straight, passive fixation
Polarity	bipolar
Steroid	no
U.S. Market Release	Sep 2002
CE Market Release	Jan 2002
Worldwide Distributed Devices	36 500
Registered U.S. Implants	8550
Estimated Active U.S. Implants	4470
U.S. Total Returned	19

	Quantity	Rate
U.S. Qualifying Complications	32	0.37%
Abnormal Pacing Impedance	10	0.12%
Conductor Fracture	2	0.02%
Failure to Capture	15	0.18%
Insulation Breach	2	0.02%
Oversensing	1	0.01%
Other	2	0.02%

	Quantity	Rate
U.S. Confirmed Malfunctions Insulation Breach	1 1	0.01% 0.01%
U.S. Acute Lead Observations	2	0.02% 0.02%

Total survival

Cumulative survival propability



Arox J

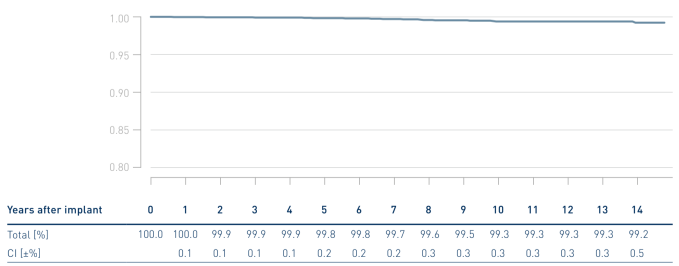
Product Versions	45-JBP, 53-JBP
Lead Type	J-shape, passive fixation
Polarity	bipolar
Steroid	no
U.S. Market Release	Sep 2002
CE Market Release	Jan 2002
Worldwide Distributed Devices	8 760
Registered U.S. Implants	3470
Estimated Active U.S. Implants	2 070
U.S. Total Returned	8

	Quantity	Rate
U.S. Qualifying Complications	16	0.46%
Abnormal Pacing Impedance	2	0.06%
Conductor Fracture	1	0.03%
Failure to Capture	10	0.29%
Lead Dislodgement	2	0.06%
Oversensing	1	0.03%

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
U.S. Acute Lead Observations	0	0.00%

• Total survival

Cumulative survival propability



Dextrus

Product Versions	4135, 4136, 4137
Lead Type	straight, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Apr 2007
CE Market Release	May 2007
Worldwide Distributed Devices	487 000
Registered U.S. Implants	385 000
Estimated Active U.S. Implants	259 000
U.S. Total Returned	2 2 2 2 6

	Quantity	Rate
U.S. Qualifying Complications	3 1 6 2	0.82%
Abnormal Pacing Impedance	256	0.07%
Cardiac Perforation	24	0.01%
Conductor Fracture	92	0.02%
Extracardiac Stimulation	18	0.00%
Failure to Capture	871	0.23%
Failure to Sense	128	0.03%
Insulation Breach	72	0.02%
Lead Dislodgement	496	0.13%
Oversensing	658	0.17%
Other	547	0.14%

	Quantity	Rate
U.S. Confirmed Malfunctions	300	0.08%
Conductor Fracture	116	0.03%
Insulation Breach	179	0.05%
Other	5	0.00%
U.S. Acute Lead Observations	1568	0.41%
Abnormal Pacing Impedance	31	0.01%
Cardiac Perforation	66	0.02%
Extracardiac Stimulation	15	0.00%
Failure to Capture	218	0.06%
Failure to Sense	60	0.02%
Insulation Breach		0.00%
Lead Dislodgement	631	0.16%
Oversensing	41	0.01%
Other	497	0.13%

• Total survival Cumulative survival propability 0.95 -0.80 -Τ Τ Years after implant 5 7 8 9 0 1 2 3 4 6 10 11 Total [%] 100.0 99.7 99.5 99.4 99.2 99.1 98.9 98.7 98.5 98.4 98.2 98.1 CI [±%] < 0.1 < 0.1 < 0.1 < 0.1 < 0.1 < 0.1 < 0.1 0.1 0.1 0.1 0.1

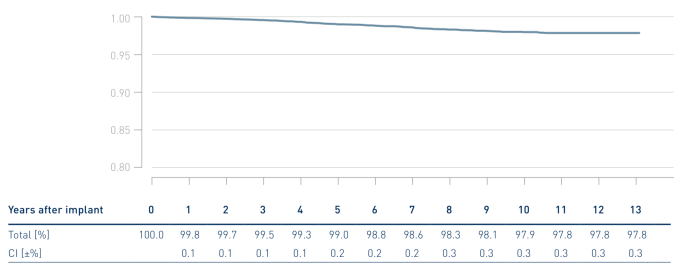
Selox JT

Product Versions	45, 53
Lead Type	J-shape, passive fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Nov 2004
CE Market Release	Nov 2004
Worldwide Distributed Devices	150 000
Registered U.S. Implants	16500
Estimated Active U.S. Implants	12400
U.S. Total Returned	115

	Quantity	Rate
U.S. Qualifying Complications	192	1.17%
Abnormal Pacing Impedance	25	0.15%
Cardiac Perforation	1	0.01%
Conductor Fracture	7	0.04%
Extracardiac Stimulation	1	0.01%
Failure to Capture	91	0.55%
Failure to Sense	8	0.05%
Insulation Breach	8	0.05%
Lead Dislodgement	30	0.18%
Oversensing	4	0.02%
Other	17	0.10%

	Quantity	Rate
U.S. Confirmed Malfunctions Insulation Breach	9 9	0.05% 0.05%
U.S. Acute Lead Observations	44	0.27%
Failure to Capture		0.05%
Lead Dislodgement	33	0.20%
Other	3	0.02%

• Total survival



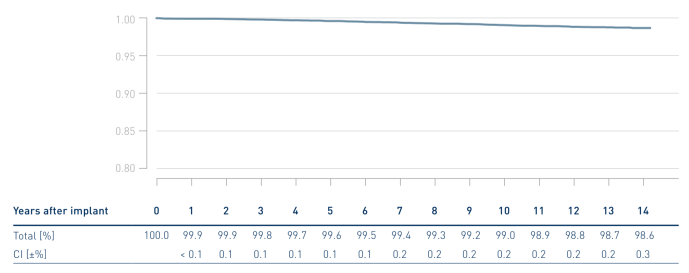
Selox SR

Product Versions	45, 53, 60
Lead Type	straight, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Mar 2004
CE Market Release	Feb 2004
Worldwide Distributed Devices	172000
Registered U.S. Implants	14400
Estimated Active U.S. Implants	7 130
U.S. Total Returned	61

	Quantity	Rate
U.S. Qualifying Complications	102	0.71%
Abnormal Pacing Impedance	4	0.03%
Conductor Fracture	9	0.06%
Extracardiac Stimulation	2	0.01%
Failure to Capture	40	0.28%
Failure to Sense	1	0.01%
Insulation Breach	6	0.04%
Lead Dislodgement	14	0.10%
Oversensing	12	0.08%
Other	14	0.10%

	Quantity	Rate
U.S. Confirmed Malfunctions	11	0.08%
Insulation Breach	11	0.08%
U.S. Acute Lead Observations	21	0.15%
Cardiac Perforation	1	0.01%
Failure to Capture	11	0.08%
Insulation Breach	1	0.01%
Lead Dislodgement	8	0.06%

• Total survival



Selox ST

Product Versions	53, 60
Lead Type	straight, passive fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Nov 2004
CE Market Release	Nov 2004
Worldwide Distributed Devices	375 000
Registered U.S. Implants	31800
Estimated Active U.S. Implants	22800
U.S. Total Returned	166

	Quantity	Rate
U.S. Qualifying Complications	512	1.61%
Abnormal Pacing Impedance	112	0.35%
Cardiac Perforation	3	0.01%
Conductor Fracture	52	0.16%
Extracardiac Stimulation	7	0.02%
Failure to Capture	248	0.78%
Failure to Sense	1	0.00%
Insulation Breach	35	0.11%
Lead Dislodgement	21	0.07%
Oversensing	10	0.03%
Other	23	0.07%

	Quantity	Rate
U.S. Confirmed Malfunctions	16	0.05%
Conductor Fracture	1	0.00%
Crimps, Welds and Bonds	1	0.00%
Insulation Breach	14	0.04%
U.S. Acute Lead Observations	48	0.15%
Abnormal Pacing Impedance	1	0.00%
Failure to Capture	20	0.06%
Lead Dislodgement	21	0.07%
Other	6	0.02%

• Total survival

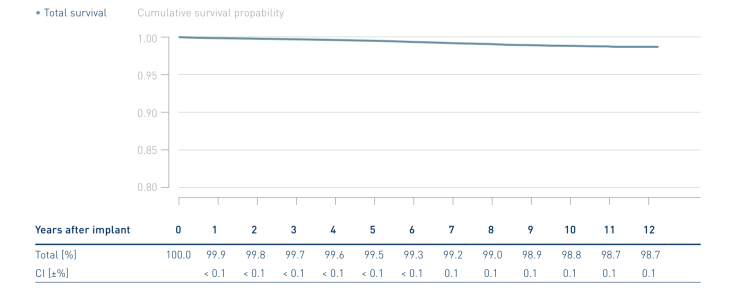


Setrox S

Product Versions	_ 45, 53, 60
Lead Type	_ straight, active fixation
Polarity	_ bipolar
Steroid	_ yes
U.S. Market Release	_ Apr 2006
CE Market Release	_ Mar 2006
Worldwide Distributed Devices	_ 668 000
Registered U.S. Implants	_ 245 000
Estimated Active U.S. Implants	_ 200 000
U.S. Total Returned	_ 1 528

	Quantity	Rate
U.S. Qualifying Complications	1244	0.51%
Abnormal Pacing Impedance	88	0.04%
Cardiac Perforation	8	0.00%
Conductor Fracture	58	0.02%
Extracardiac Stimulation	11	0.00%
Failure to Capture	444	0.18%
Failure to Sense	33	0.01%
Insulation Breach	64	0.03%
Lead Dislodgement	287	0.12%
Oversensing	156	0.06%
Other	95	0.04%

	Quantity	Rate
U.S. Confirmed Malfunctions	141	0.06%
Conductor Fracture	50	0.02%
Insulation Breach	89	0.04%
Other	2	0.00%
U.S. Acute Lead Observations	272	0.11%
Abnormal Pacing Impedance	1	0.00%
Cardiac Perforation	23	0.01%
Failure to Capture	36	0.01%
Failure to Sense	3	0.00%
Insulation Breach	4	0.00%
Lead Dislodgement	189	0.08%
Other	16	0.01%

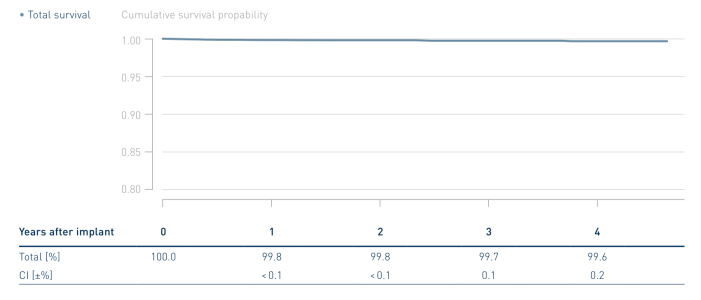


Siello S/Solia S

Product Versions	45, 53, 60
Lead Type	
Polarity	bipolar
Steroid	yes
U.S. Market Release	Jun 2016
CE Market Release	Jul 2009
Worldwide Distributed Devices	1023000
Registered U.S. Implants	73800
Estimated Active U.S. Implants	70800
U.S. Total Returned	266

	Quantity	Rate
U.S. Qualifying Complications	112	0.15%
Abnormal Pacing Impedance	3	0.00%
Cardiac Perforation	7	0.01%
Conductor Fracture	1	0.00%
Failure to Capture	37	0.05%
Failure to Sense	3	0.00%
Insulation Breach	1	0.00%
Lead Dislodgement	57	0.08%
Oversensing	2	0.00%
Other	1	0.00%

	Quantity	Rate
U.S. Confirmed Malfunctions	12	0.02%
Conductor Fracture	4	0.01%
Insulation Breach	8	0.01%
U.S. Acute Lead Observations	98	0 13%
Abnormal Pacing Impedance		0.00%
Cardiac Perforation		0.01%
Conductor Fracture	1	0.00%
Failure to Capture	26	0.04%
Failure to Sense	2	0.00%
Insulation Breach	1	0.00%
Lead Dislodgement	57	0.08%
Oversensing	2	0.00%
Other		0.00%



Tilda JT

Product Versions	45, 53
Lead Type	J-shape, passive fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Feb 2012
CE Market Release	Sep 2011
Worldwide Distributed Devices	17 400
Registered U.S. Implants	764
Estimated Active U.S. Implants	743
U.S. Total Returned	0

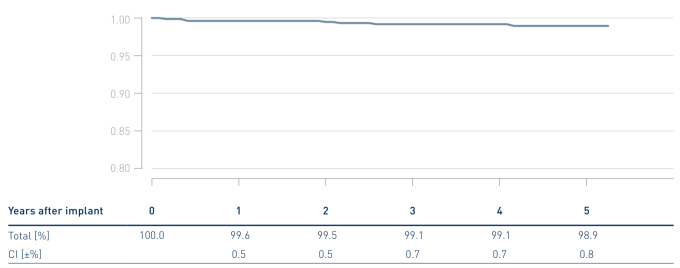
Rate 0.92% 0.26% 0.26% 0.39%

	Quantity
U.S. Qualifying Complications	7
Abnormal Pacing Impedance	2
Failure to Capture	2
Lead Dislodgement	3

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
U.S. Acute Lead Observations Lead Dislodgement	1 1	0.13% 0.13%

• Total survival

Cumulative survival propability

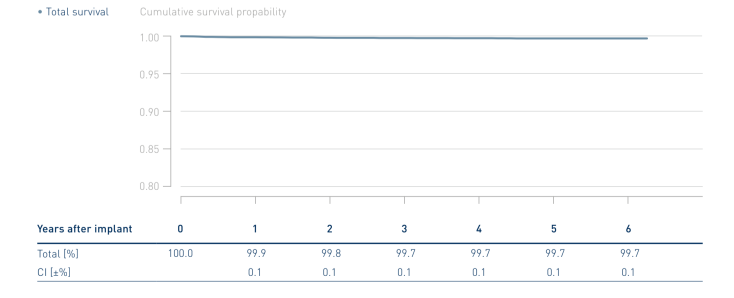


Tilda R

Product Versions Lead Type	45, 53, 60 straight, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Dec 2011
CE Market Release	Aug 2011
Worldwide Distributed Devices	42000
Registered U.S. Implants	9660
Estimated Active U.S. Implants	9 280
U.S. Total Returned	15

	Quantity	Rate
U.S. Qualifying Complications	27	0.28%
Abnormal Pacing Impedance	1	0.01%
Conductor Fracture	3	0.03%
Extracardiac Stimulation	1	0.01%
Failure to Capture	7	0.07%
Insulation Breach	2	0.02%
Lead Dislodgement		0.09%
Oversensing	1	0.01%
Other	3	0.03%

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.01%
Conductor Fracture	1	0.01%
U.S. Acute Lead Observations	9	0.09%
Failure to Capture	1	0.01%
Lead Dislodgement	8	0.08%



Tilda T

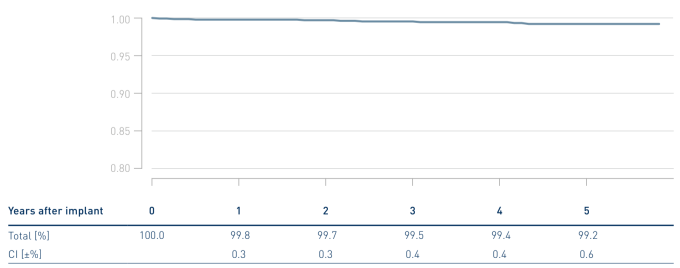
Product Versions	53, 60
Lead Type	straight, passive fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Dec 2011
CE Market Release	Aug 2011
Worldwide Distributed Devices	22 700
Registered U.S. Implants	1310
Estimated Active U.S. Implants	1260
U.S. Total Returned	1

	Quantity	Rate
U.S. Qualifying Complications		0.69%
Abnormal Pacing Impedance	3	0.23%
Conductor Fracture	1	0.08%
Insulation Breach	1	0.08%
Lead Dislodgement	4	0.31%

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
U.S. Acute Lead Observations	0	0.00%

• Total survival

Cumulative survival propability



Kentrox RV

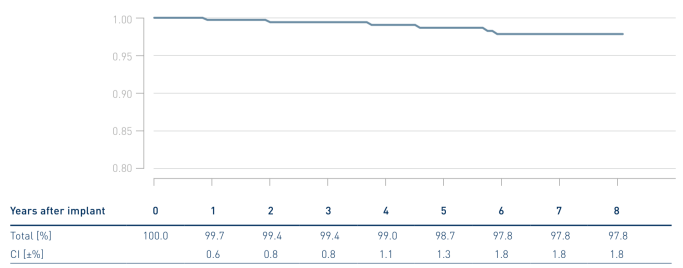
Product Versions	65, 75, -Steroid
Lead Type	single-coil, passive fixation
Polarity	bipolar
Steroid	yes/no
U.S. Market Release	Mar 2002 / Oct 2004
CE Market Release	Jan 2001 / Dec 2004
Worldwide Distributed Devices	5 4 9 0
Registered U.S. Implants	409
Estimated Active U.S. Implants	172
U.S. Total Returned	8

	Quantity	Rate
U.S. Qualifying Complications	8	1.96%
Conductor Fracture	1	0.24%
Failure to Capture	2	0.49%
Insulation Breach	1	0.24%
Oversensing	4	0.98%

	Quantity	Rate
U.S. Confirmed Malfunctions	2	0.49%
Conductor Fracture	1	0.24%
Insulation Breach	1	0.24%
U.S. Acute Lead Observations	0	0.00%

• Total survival

Cumulative survival propability



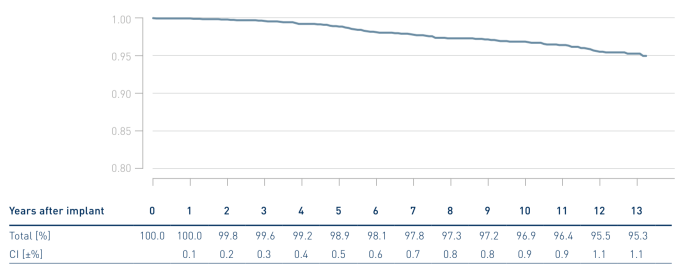
Kentrox SL-S

Product Versions	65/16, 18 -Steroid
Lead Type	dual-coil, active fixation
Polarity	bipolar
Steroid	yes/no
U.S. Market Release	Oct 2004
CE Market Release	Jun 2004
Worldwide Distributed Devices	8 730
Registered U.S. Implants	2440
Estimated Active U.S. Implants	1250
U.S. Total Returned	41

	Quantity	Rate
U.S. Qualifying Complications	57	2.34%
Abnormal Defibrillation Impedance	1	0.04%
Abnormal Pacing Impedance	3	0.12%
Conductor Fracture	4	0.16%
Failure to Capture	3	0.12%
Failure to Sense	1	0.04%
Insulation Breach	3	0.12%
Lead Dislodgement	3	0.12%
Oversensing	37	1.52%
Other	2	0.08%

	Quantity	Rate
U.S. Confirmed Malfunctions	14	0.57%
Insulation Breach	14	0.57%
U.S. Acute Lead Observations	2	0.08%
Insulation Breach	1	0.04%
Oversensing	1	0.04%

• Total survival

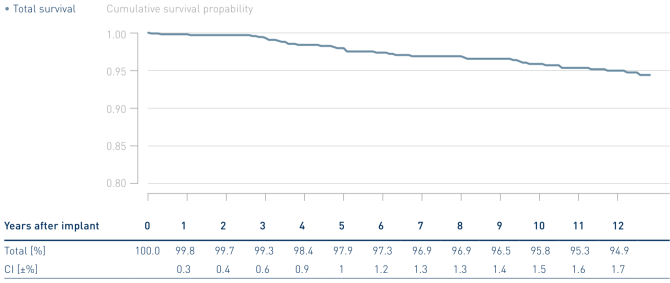


Kentrox SL

Product Versions Lead Type	65, 75, 100, -Steroid dual coil, passive fixation
Polarity	bipolar
Steroid	
U.S. Market Release	Oct 2004
CE Market Release	Dec 2003 / Dec 2004
Worldwide Distributed Devices	8480
Registered U.S. Implants	1020
Estimated Active U.S. Implants	538
U.S. Total Returned	19

	Quantity	Rate
U.S. Qualifying Complications	31	3.06%
Abnormal Defibrillation Impedance	1	0.10%
Abnormal Pacing Impedance	3	0.30%
Conductor Fracture	3	0.30%
Failure to Capture	1	0.10%
Insulation Breach	6	0.59%
Oversensing	15	1.48%
Other	2	0.20%

	Quantity	Rate
U.S. Confirmed Malfunctions	5	0.49%
Insulation Breach	5	0.49%
U.S. Acute Lead Observations	0	0.00%



Linox S

Product Versions Lead Type	
Polarity	
Steroid	yes
U.S. Market Release	Feb 2007
CE Market Release	Mar 2007
Worldwide Distributed Devices	32 500
Registered U.S. Implants	2 500
Estimated Active U.S. Implants	1690
U.S. Total Returned	77

	Quantity	Rate
U.S. Qualifying Complications	65	2.61%
Abnormal Defibrillation Impedance	6	0.24%
Abnormal Pacing Impedance	3	0.12%
Conductor Fracture	5	0.20%
Failure to Capture	7	0.28%
Insulation Breach	4	0.16%
Oversensing	34	1.36%
Other	6	0.24%

	Quantity	Rate
U.S. Confirmed Malfunctions Conductor Fracture Insulation Breach		1.61% 0.20% 1.40%
U.S. Acute Lead Observations Lead Dislodgement Other		0.08% 0.04% 0.04%

• Total survival Cumulative survival propability 0.95 -1 0.90 · 0.85 -0.80 — Т T Years after implant 0 1 2 3 4 5 6 7 8 9 10 Total [%] 100.0 99.8 99.2 98.8 98.0 97.1 96.3 95.4 94.8 94.5 93.7 CI [±%] 0.2 0.4 0.5 0.6 0.7 0.8 0.9 1 1.1 1.4

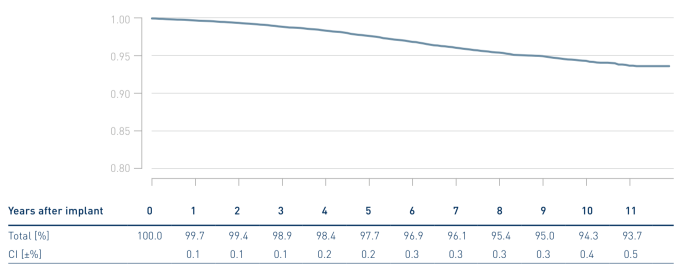
Linox SD

Product Versions	60/16, 65/16, 65/18, 75/18 dual-coil, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Apr 2006
CE Market Release	Aug 2006
Worldwide Distributed Devices	55100
Registered U.S. Implants	22300
Estimated Active U.S. Implants	14500
U.S. Total Returned	468

	Quantity	Rate
U.S. Qualifying Complications	686	3.08%
Abnormal Defibrillation Impedance	52	0.23%
Abnormal Pacing Impedance	47	0.21%
Cardiac Perforation	3	0.01%
Conductor Fracture	72	0.32%
Failure to Capture	66	0.30%
Failure to Sense	11	0.05%
Insulation Breach	57	0.26%
Lead Dislodgement	31	0.14%
Oversensing	303	1.36%
Other	44	0.20%

	Quantity	Rate
U.S. Confirmed Malfunctions	188	0.84%
Conductor Fracture	26	0.12%
Insulation Breach	161	0.72%
Other	1	0.00%
U.S. Acute Lead Observations	11	0.05%
Abnormal Pacing Impedance	1	0.00%
Cardiac Perforation	1	0.00%
Failure to Capture	1	0.00%
Lead Dislodgement	6	0.03%
Oversensing	1	0.00%
Other	1	0.00%

• Total survival



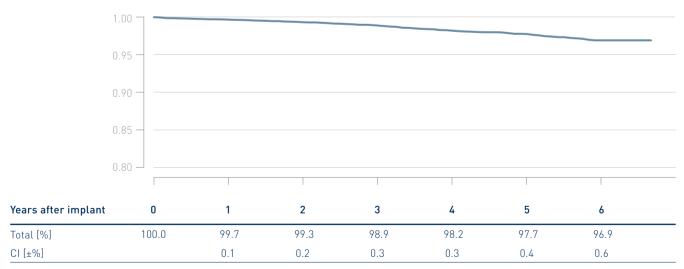
Linox^{smart} S

Product Versions	60, 65, 75 single-coil, active fixation
Polarity	5
Steroid	yes
U.S. Market Release	Aug 2011
CE Market Release	Dec 2010
Worldwide Distributed Devices	46 800
Registered U.S. Implants	7650
Estimated Active U.S. Implants	6460
U.S. Total Returned	145

	Quantity	Rate
U.S. Qualifying Complications	101	1.32%
Abnormal Defibrillation Impedance	5	0.07%
Abnormal Pacing Impedance	5	0.07%
Cardiac Perforation	1	0.01%
Conductor Fracture	9	0.12%
Failure to Capture	15	0.20%
Failure to Sense	4	0.05%
Insulation Breach	3	0.04%
Lead Dislodgement	14	0.18%
Oversensing	39	0.51%
Other	6	0.08%

	Quantity	Rate
U.S. Confirmed Malfunctions	47	0.61%
Conductor Fracture	7	0.09%
Insulation Breach	40	0.52%
U.S. Acute Lead Observations	11	0.14%
Abnormal Pacing Impedance	1	0.01%
Cardiac Perforation	1	0.01%
Lead Dislodgement	8	0.10%
Other	1	0.01%

• Total survival



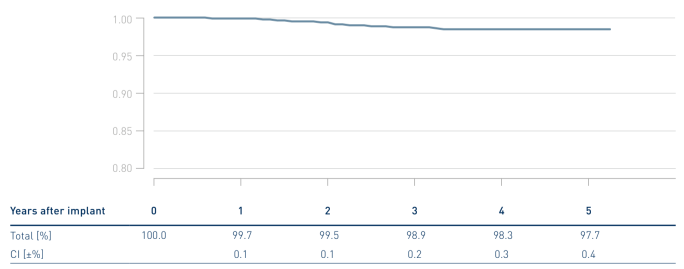
Linox^{smart} S DX

Product Versions Lead Type Polarity	65/15, 65/17 single-coil, active fixation bipolar
Steroid	
U.S. Market Release	
CE Market Release	Mar 2010
Worldwide Distributed Devices	35 700
Registered U.S. Implants	16300
Estimated Active U.S. Implants	14900
U.S. Total Returned	260

	Quantity	Rate
U.S. Qualifying Complications	113	0.70%
Abnormal Defibrillation Impedance	9	0.06%
Abnormal Pacing Impedance	3	0.02%
Conductor Fracture	16	0.10%
Failure to Capture	11	0.07%
Failure to Sense	7	0.04%
Insulation Breach	2	0.01%
Lead Dislodgement	28	0.17%
Oversensing	33	0.20%
Other	4	0.02%

	Quantity	Rate
U.S. Confirmed Malfunctions	54	0.33%
Conductor Fracture	2	0.01%
Insulation Breach	52	0.32%
U.S. Acute Lead Observations	39	0.24%
Cardiac Perforation	4	0.02%
Failure to Capture	9	0.06%
Lead Dislodgement	17	0.10%
Oversensing	2	0.01%
Other	7	0.04%

• Total survival



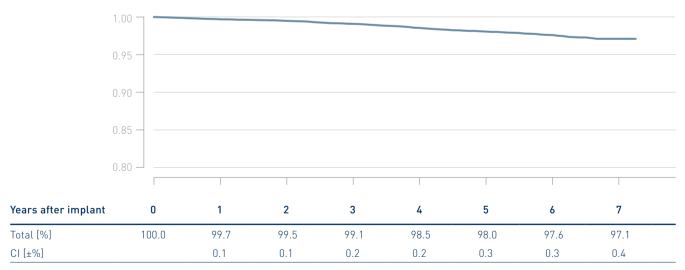
Linox^{smart} SD

Product Versions Lead Type	60/16, 65/16, 65/18, 75/18 dual-coil, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Jan 2011
CE Market Release	Oct 2009
Worldwide Distributed Devices	55 400
Registered U.S. Implants	13300
Estimated Active U.S. Implants	10900
U.S. Total Returned	215

	Quantity	Rate
U.S. Qualifying Complications	199	1.50%
Abnormal Defibrillation Impedance	15	0.11%
Abnormal Pacing Impedance	6	0.05%
Conductor Fracture	24	0.18%
Extracardiac Stimulation	1	0.01%
Failure to Capture	17	0.13%
Failure to Sense	4	0.03%
Insulation Breach	7	0.05%
Lead Dislodgement	23	0.17%
Oversensing	96	0.73%
Other	6	0.05%

	Quantity	Rate
U.S. Confirmed Malfunctions	52	0.39%
Conductor Fracture	4	0.03%
Insulation Breach	47	0.36%
Other	1	0.01%
U.S. Acute Lead Observations	29	0.22%
Abnormal Defibrillation Impedance	1	0.01%
Cardiac Perforation	2	0.02%
Failure to Capture	4	0.03%
Insulation Breach	1	0.01%
Lead Dislodgement	12	0.09%
Oversensing	2	0.02%
Other	7	0.05%

• Total survival



Linox^{smart} TD

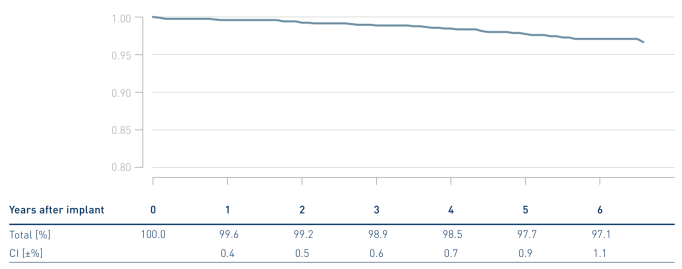
Product Versions	65/16, 65/18, 75/18
Lead Type	dual-coil, passive fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Jan 2011
CE Market Release	Oct 2009
Worldwide Distributed Devices	7740
Registered U.S. Implants	1280
Estimated Active U.S. Implants	1040
U.S. Total Returned	19

	Quantity	Rate
U.S. Qualifying Complications	25	1.96%
Abnormal Defibrillation Impedance	4	0.31%
Abnormal Pacing Impedance	2	0.16%
Conductor Fracture	1	0.08%
Failure to Capture	6	0.47%
Insulation Breach	2	0.16%
Lead Dislodgement	5	0.39%
Oversensing	7	0.55%

	Quantity	Rate
U.S. Confirmed Malfunctions Insulation Breach	1 1	0.08% 0.08%
U.S. Acute Lead Observations	3	0.24% 0.24%



Cumulative survival propability



Linox T

• Total survival

Product Versions	65, 75
Lead Type	single-coil, passive fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Feb 2007
CE Market Release	Mar 2007
Worldwide Distributed Devices	2 280
Registered U.S. Implants	322
Estimated Active U.S. Implants	225
U.S. Total Returned	4

	Quantity	Rate
U.S. Qualifying Complications	13	4.04%
Abnormal Pacing Impedance	2	0.62%
Failure to Capture	3	0.93%
Insulation Breach	1	0.31%
Oversensing	6	1.86%
Other	1	0.31%

Cumulative survival propability

	Quantity	Rate
U.S. Confirmed Malfunctions Conductor Fracture Insulation Breach	1	0.93% 0.31% 0.62%
U.S. Acute Lead Observations Other	2 1 1	0.82% 0.31% 0.31%

0.95 -0.85 -Τ Years after implant 0 1 2 3 4 5 6 7 Total [%] 100.0 100.0 99.3 99.0 97.1 96.3 95.4 94.6 CI [±%] < 0.1 0.9 1.2 2 2.3 2.5 2.8

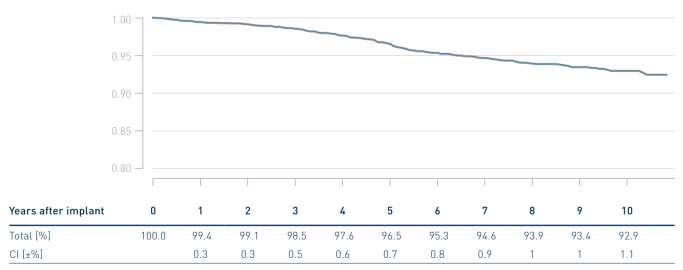
Linox TD

Product Versions Lead Type	65/16, 75/16, 100/16, 100/18 dual-coil, passive fixation
	bipolar
Steroid	yes
U.S. Market Release	Oct 2006
CE Market Release	Oct 2006
Worldwide Distributed Devices	14600
Registered U.S. Implants	3060
Estimated Active U.S. Implants	2040
U.S. Total Returned	77

	Quantity	Rate
U.S. Qualifying Complications	118	3.87%
Abnormal Defibrillation Impedance	10	0.33%
Abnormal Pacing Impedance	12	0.39%
Conductor Fracture	15	0.49%
Failure to Capture	18	0.59%
Failure to Sense	4	0.13%
Insulation Breach	13	0.43%
Lead Dislodgement	4	0.13%
Oversensing	39	1.28%
Other	3	0.10%

	Quantity	Rate
U.S. Confirmed Malfunctions Conductor Fracture	37	1.21% 0.20%
Insulation Breach	31	1.02%
U.S. Acute Lead Observations	3	0.10%
Failure to Capture	1	0.03%
Lead Dislodgement	2	0.07%

• Total survival



Plexa S

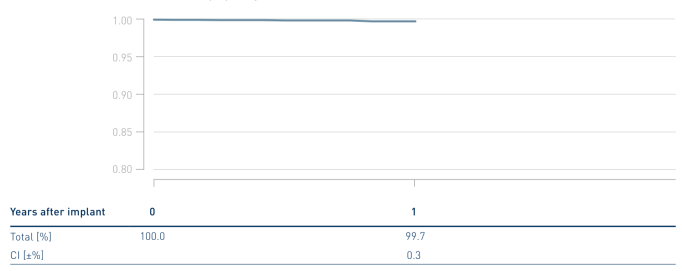
Product Versions	65, 75 single-coil, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Mar 2017
CE Market Release	Feb 2017
Worldwide Distributed Devices	27 100
Registered U.S. Implants	3 520
Estimated Active U.S. Implants	3400
U.S. Total Returned	15

	Quantity	Rate
U.S. Qualifying Complications	4	0.11%
Cardiac Perforation	1	0.03%
Lead Dislodgement	3	0.09%

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
U.S. Acute Lead Observations Failure to Capture	6 1	0.17% 0.03%
Lead Dislodgement	5	0.14%

• Total survival

Cumulative survival propability



Plexa S DX

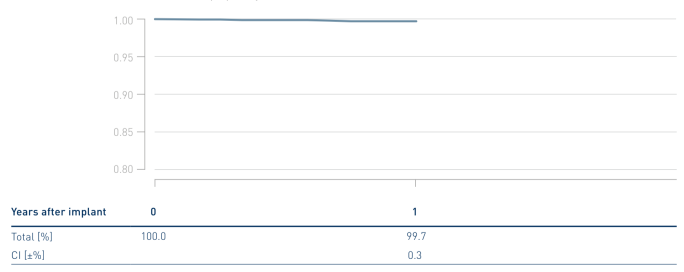
Product Versions Lead Type	65/15, 65/17 single-coil, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Mar 2017
CE Market Release	Feb 2017
Worldwide Distributed Devices	9 480
Registered U.S. Implants	3660
Estimated Active U.S. Implants	3560
U.S. Total Returned	20

	Quantity	Rate
U.S. Qualifying Complications	3	0.08%
Failure to Capture	1	0.03%
Lead Dislodgement	1	0.03%
Oversensing	1	0.03%

	Quantity	Rate
U.S. Confirmed Malfunctions	4 4	0.11% 0.11%
U.S. Acute Lead Observations	2	0.05%

• Total survival

Cumulative survival propability



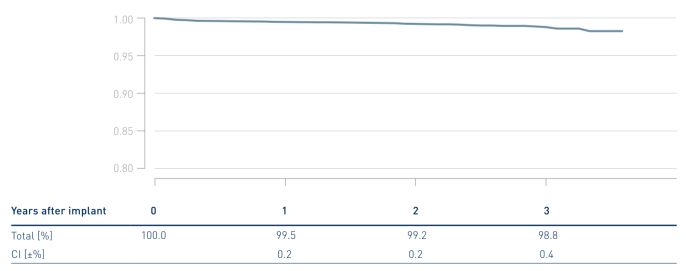
Protego S

Product Versions	60, 65, 75
Lead Type	single-coil, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Jul 2014
CE Market Release	Feb 2014
Worldwide Distributed Devices	53 900
Registered U.S. Implants	8 150
Estimated Active U.S. Implants	7450
U.S. Total Returned	68

	Quantity	Rate
U.S. Qualifying Complications	54	0.66%
Abnormal Pacing Impedance	1	0.01%
Cardiac Perforation	1	0.01%
Conductor Fracture	3	0.04%
Extracardiac Stimulation	1	0.01%
Failure to Capture	10	0.12%
Failure to Sense	1	0.01%
Lead Dislodgement	22	0.27%
Oversensing	11	0.13%
Other	4	0.05%

	Quantity	Rate
U.S. Confirmed Malfunctions	12	0.15%
Conductor Fracture	2	0.02%
Insulation Breach	10	0.12%
U.S. Acute Lead Observations	28	0.34%
Cardiac Perforation	2	0.02%
Extracardiac Stimulation	1	0.01%
Failure to Capture	3	0.04%
Lead Dislodgement	13	0.16%
Other		0.11%

• Total survival



Protego SD

• Total survival

Product Versions	60/16, 65/16, 65/18, 75/18 dual-coil, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Jul 2014
CE Market Release	May 2013
Worldwide Distributed Devices	18 500
Registered U.S. Implants	3420
Estimated Active U.S. Implants	3140
U.S. Total Returned	27

	Quantity	Rate
U.S. Qualifying Complications	17	0.50%
Abnormal Pacing Impedance	1	0.03%
Conductor Fracture	2	0.06%
Failure to Capture	3	0.09%
Insulation Breach	1	0.03%
Lead Dislodgement	4	0.12%
Oversensing	6	0.18%

Cumulative survival propability

	Quantity	Rate
U.S. Confirmed Malfunctions Insulation Breach	5	0.15% 0.15%
U.S. Acute Lead Observations Lead Dislodgement Other	3 2 1	0.09% 0.06% 0.03%

0.95 -0.85 -Years after implant 1 2 3 0 Total [%] 100.0 99.8 99.5 99.0 CI [±%] 0.1 0.3 0.5

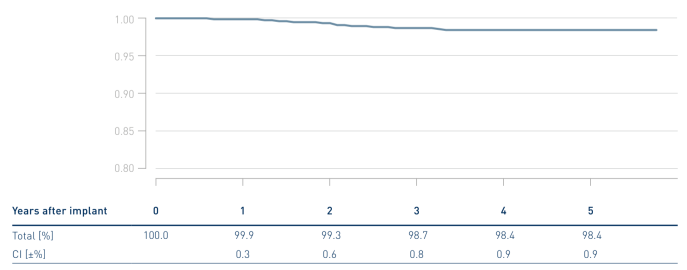
Vigila 2CR

Product Versions Lead Type	
Polarity	bipolar
Steroid	yes
U.S. Market Release	Feb 2012
CE Market Release	Oct 2011
Worldwide Distributed Devices	3010
Registered U.S. Implants	799
Estimated Active U.S. Implants	728
U.S. Total Returned	11

	Quantity	Rate		Quantity	Rate
U.S. Qualifying Complications	9	1.13%	U.S. Confirmed Malfunctions	3	0.38%
Conductor Fracture	1	0.13%	Insulation Breach	3	0.38%
Lead Dislodgement	3	0.38%			
Oversensing	5	0.63%	U.S. Acute Lead Observations	0	0.00%

• Total survival

Cumulative survival propability



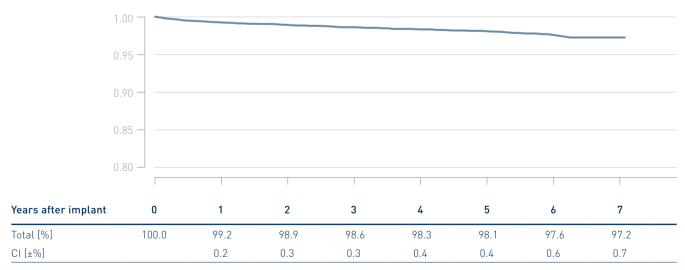
Corox OTW-L

Product Versions	_ 75, 85
Lead Type	dual-curve fixation
Polarity	_ bipolar
Steroid	_ yes
U.S. Market Release	_ Jan 2011
CE Market Release	Dec 2009
Worldwide Distributed Devices	_ 31800
Registered U.S. Implants	_ 6 2 9 0
Estimated Active U.S. Implants	_ 5120
U.S. Total Returned	69

	Quantity	Rate
U.S. Qualifying Complications	98	1.56%
Abnormal Pacing Impedance	1	0.02%
Conductor Fracture	4	0.06%
Extracardiac Stimulation	16	0.25%
Failure to Capture	38	0.60%
Failure to Sense	1	0.02%
Insulation Breach	1	0.02%
Lead Dislodgement	30	0.48%
Oversensing	1	0.02%
Other	6	0.10%

	Quantity	Rate
U.S. Confirmed Malfunctions	4	0.06%
Conductor Fracture	3	0.05%
Insulation Breach	1	0.02%
U.S. Acute Lead Observations	21	0.33%
Extracardiac Stimulation	6	0.10%
Failure to Capture	2	0.03%
Lead Dislodgement	10	0.16%
Other	3	0.05%

• Total survival

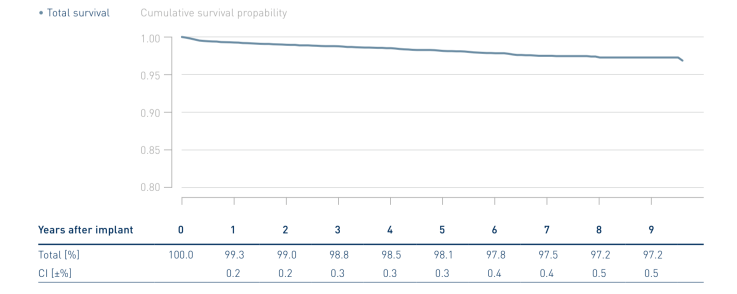


Corox OTW-S

Product Versions	75, 85
Lead Type	thread fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	May 2008
CE Market Release	Dec 2006
Worldwide Distributed Devices	26500
Registered U.S. Implants	8220
Estimated Active U.S. Implants	5790
U.S. Total Returned	122

	Quantity	Rate
U.S. Qualifying Complications	131	1.60%
Abnormal Pacing Impedance	6	0.07%
Conductor Fracture	4	0.05%
Extracardiac Stimulation	13	0.16%
Failure to Capture		0.41%
Insulation Breach	4	0.05%
Lead Dislodgement	53	0.65%
Oversensing	2	0.02%
Other	15	0.18%

	Quantity	Rate
U.S. Confirmed Malfunctions	11	0.13%
Conductor Fracture	6	0.07%
Insulation Breach	4	0.05%
Other	1	0.01%
U.S. Acute Lead Observations	33	0.40%
Cardiac Perforation	1	0.01%
Extracardiac Stimulation	5	0.06%
Failure to Capture	6	0.07%
Lead Dislodgement	20	0.24%
Other	1	0.01%



Corox OTW

Product Versions	75, 85
Lead Type	helix fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	May 2008
CE Market Release	Dec 2006
Worldwide Distributed Devices	28 800
Registered U.S. Implants	4 1 4 0
Estimated Active U.S. Implants	2670
U.S. Total Returned	74

	Quantity	Rate
U.S. Qualifying Complications	101	2.44%
Abnormal Pacing Impedance	2	0.05%
Conductor Fracture	3	0.07%
Extracardiac Stimulation		0.19%
Failure to Capture	36	0.87%
Insulation Breach	2	0.05%
Lead Dislodgement		0.90%
Oversensing	2	0.05%
Other	11	0.27%

	Quantity	Rate
U.S. Confirmed Malfunctions Conductor Fracture Insulation Breach		0.39% 0.36% 0.02%
U.S. Acute Lead Observations Lead Dislodgement Other	9 7 2	0.22% 0.17% 0.05%

• Total survival Cumulative survival propability 0.95 -0.85 -Τ Т Τ Years after implant 1 2 3 4 5 6 7 8 9 0 Total [%] 100.0 99.5 98.8 98.1 97.7 97.1 96.7 96.1 95.9 95.9 CI [±%] 0.2 0.4 0.5 0.5 0.6 0.7 0.7 0.8 0.8

Corox OTW

Product Versions	_ 75, 85
Lead Type	helix fixation
Polarity	_ unipolar
Steroid	_yes
U.S. Market Release	_ Aug 2006
CE Market Release	_ Apr 2004
Worldwide Distributed Devices	_ 10400
Registered U.S. Implants	_ 1 4 3 0
Estimated Active U.S. Implants	_ 686
U.S. Total Returned	26

	Quantity	Rate
U.S. Qualifying Complications	39	2.74%
Extracardiac Stimulation	7	0.49%
Failure to Capture	14	0.98%
Insulation Breach	2	0.14%
Lead Dislodgement	10	0.70%
Oversensing	1	0.07%
Other	5	0.35%

	Quantity	Rate
U.S. Confirmed Malfunctions	2	0.14%
Insulation Breach	2	0.14%
U.S. Acute Lead Observations	4	0.28%
Failure to Capture	3	0.21%
Lead Dislodgement	1	0.07%

0.85 -Τ Years after implant 2 3 4 5 6 7 8 9 0 1 10 11 Total [%] 100.0 99.2 99.0 98.7 98.0 97.7 97.0 96.1 95.9 95.5 98.4 96.8 CI [±%] 0.5 0.6 0.6 0.7 0.8 0.9 1.1 1.1 1.3 1.3 1.4

• Total survival Cumulative survival propability

Sentus OTW QP L

Product Versions	_ 75, 75/49, 85, 85/49 dual-curve fixation
Lead Type Polarity	_ quadripolar
Steroid	_ yes
U.S. Market Release	_ May 2017
CE Market Release	_ Dec 2014
Worldwide Distributed Devices	_ 44 200
Registered U.S. Implants	_ 3 720
Estimated Active U.S. Implants	_ 3 290
U.S. Total Returned	_ 24

	Quantity	Rate
U.S. Qualifying Complications	23	0.62%
Extracardiac Stimulation	2	0.05%
Failure to Capture	5	0.13%
Lead Dislodgement	15	0.40%
Other	1	0.03%

	Quantity	Rate
U.S. Confirmed Malfunctions Conductor Fracture	1	0.03% 0.03%
	I	0.0370
U.S. Acute Lead Observations	8	0.22%
Extracardiac Stimulation	2	0.05%
Lead Dislodgement	5	0.13%
Oversensing	1	0.03%

• Total survival Cumulative survival propability 0.95 -0.85 -0.80 Years after implant 1 0 Total [%] 100.0 98.7 CI [±%] 0.7

Sentus OTW QP S

Product Versions	75, 75/49, 85, 85/49
Lead Type	thread fixation
Polarity	_ quadripolar
Steroid	yes
U.S. Market Release	May 2017
CE Market Release	Dec 2014
Worldwide Distributed Devices	11900
Registered U.S. Implants	1420
Estimated Active U.S. Implants	1170
U.S. Total Returned	38

	Quantity	Rate
U.S. Qualifying Complications	10	0.71%
Extracardiac Stimulation	1	0.07%
Failure to Capture	1	0.07%
Lead Dislodgement		0.56%

	Quantity	Rate
U.S. Confirmed Malfunctions		0.14% 0.14%
U.S. Acute Lead Observations	19	1.34%
Extracardiac Stimulation	1	0.07%
Failure to Capture	3	0.21%
Lead Dislodgement	15	1.06%

• Total survival Cumulative survival propability 0.95 -0.85 -0.80 Years after implant 1 0 Total [%] 100.0 98.8 CI [±%] 0.7

Methodology for Lead Survival Estimates Based on Clinical Studies

7.1 Introduction

7.2 BIOTRONIK's Clinical Studies

7.3 Lead Complications

7. Methodology for Lead Survival Estimates Based on Clinical Studies

7.1 Introduction

All leads and lead segments returned to BIOTRONIK are thoroughly analyzed to determine whether or not they meet BIOTRONIK's long term quality standards.

Although analysis of returned product is an excellent method for gaining insight into lead failure mechanisms, this data relies on the return of explanted leads. For the majority of complications the lead is not received for analysis as challenging clinical environments may not allow for the return, e.g. the extraction of an implanted lead may not be possible.

BIOTRONIK includes all reported chronic complications in the calculation of the survival estimates as described in chapter 6, i.e. reports with returned and without returned products.

However, BIOTRONIK can only report events in the survival estimates if those events were reported to BIOTRONIK. In order to eliminate possible biased survival estimates due to underreporting, BIOTRONIK performs clinical surveillance studies with active follow-up's under FDA guidance yielding the most reliable lead performance data.

In the following chapter BIOTRONIK shows – in addition to the survival data based on returned product analysis and chronic complication information – the lead performance data from clinical trials. These studies are designed to record clinical observations representative of the total clinical experience.

7.2 BIOTRONIK's Clinical Studies

7.2.1 GALAXY and CELESTIAL

BIOTRONIK'S GALAXY and CELESTIAL Registries are prospective, nonrandomized, observational studies. The key purpose of these registries is to confirm the long-term safety and reliability of BIOTRONIK leads as used in conjunction with a BIOTRONIK ICD (GALAXY) or CRT (CELESTIAL) system. All devices in the registries are legally marketed and available to physicians according to approved FDA indications for use. GALAXY and CELESTIAL Registries are registered on clinicaltrials.gov under NCT00836589 and NCT00810264 respectively.

The evaluation of safety for GALAXY is based on the analysis of BIOTRONIK Linox ICD lead-related adverse events. The evaluation of safety for CELESTIAL is based on the analysis of BIOTRONIK Corox LV pacing lead-related adverse events. However, many CELESTIAL patients also have a Linox ICD lead implanted and the Linox clinical studies data in this report represents combined data from the GALAXY and CELESTIAL registries. Both registries are designed to continue for a 5 year follow-up duration per patient. As of the January 2018 PPR, incremental updates to Linox data originate from the CELESTIAL Registry, as the GALAXY Registry is complete. To ensure a sufficiently large and representative source of data, participating clinical sites must meet pre-specified selection criteria.

Every effort is made to ensure participants are representative of the range of clinical environments in which BIOTRONIK's cardiac rhythm products are used. Patients will be seen for routine follow-up visits per their institutional standard of care until they can no longer be followed (e.g., death, lost to follow-up, etc.). However, to ensure regular patient status assessments are completed, follow-up windows consistent with typical care practices have been established with a minimum of once every six months follow-up reguirement.

During each follow up at a study center the following steps are required during the follow-up visit:

- Interrogate programmed parameters
- Determine lead electrical parameters
- Evaluate device diagnostics and programmed parameters to ensure the device is providing appropriate therapy
- Determine if there are any reportable lead-related, pulse generator-related or implant procedure-related adverse events. If there are, complete an adverse event electronic case report form (eCRF)
- Complete all appropriate eCRFs

This active surveillance model ensures a robust dataset for effectively monitoring product performance.

Patient Enrollment Criteria

To support the objectives of these registries, participants are required to meet the following inclusion criteria prior to enrollment:

- Successfully implanted BIOTRONIK ICD (GALAXY) or BIOTRONIK CRT (CELESTIAL) system, including the study lead
- Able to understand the nature of the registry and give informed consent
- Available for follow-up visits on a regular basis at the study site
- Age greater than or equal to 18 years

Each site must inform BIOTRONIK whenever a lead complication has

occurred or when a patient is no longer participating.

All leads that experience a complication and are subsequently explanted and returned to BIOTRONIK undergo root-cause analyses. Product performance is analyzed as a function of time using the survival analysis method. Root causes for any failures, regardless of the incidence rates, are investigated.

7.2.2 SIELLO Clinical Study

BIOTRONIK's SIELLO Clinical Study is a prospective, non-randomized, combined Pre-Market Study and Post-Approval Registry designed to demonstrate the safety and effectiveness of the Siello pacing lead as used in conjunction with any market-released BIOTRONIK pacemaker device. The SIELLO Clinical Study is registered on clinicaltrials.gov under NCT01791127.

For the Pre-Market Study, the evaluation of safety is based on the analysis of Siello lead-related adverse events through a follow-up time of 12 months post-implant, while the evaluation of effectiveness is based on analysis of the success rate of the implanted system including one or two Siello leads to sense and deliver pacing at 12 months post-implant.

For the Post-Approval Registry, the evaluation of long-term safety is based on the analysis of Siello lead-related adverse events through a follow-up time of 5 years post-implant. To ensure a sufficiently large and representative source of data, participating clinical sites must meet pre-specified selection criteria.

Every effort is made to ensure participants are representative of the range of clinical environments in which BIOTRONIK's cardiac rhythm products are used. Patients will be seen for routine follow-up visits until they can no longer be followed (e.g., death, lost to follow-up, etc.). However, to ensure regular patient status assessments are completed, a study follow-up schedule consistent with typical care practices has been established, with required follow-ups at 3 and 6 months post-implant and every 6 months thereafter.

During each study follow-up visit the following steps are required:

- Interrogate programmed parameters
- Record electrical parameters of the implanted leads. Perform all pacing threshold measurements at 0.4 ms or 0.5 ms pulse width when feasible
- Evaluate device diagnostics, electrical parameters and programmed parameters to ensure the device is correctly pacing and sensing
- Determine if there are any leadrelated, pulse generator-related or procedure related adverse events. If any are recorded, complete the Adverse Event eCRF
- Complete all appropriate eCRFs

This active surveillance model ensures a robust dataset for effectively monitoring product performance.

Patient Enrollment Criteria

To support the objectives of the study, participants are required to meet the following inclusion criteria prior to enrollment:

- Candidate for de novo implantation of a market-released BIOTRONIK pacemaker system, including one or two Siello leads. Candidate meets recommendation for pacemaker system implant put forth by guidelines of relevant professional societies
- Able to understand the nature of the study and provide informed consent
- Available for follow-up visits on a regular basis at the investigational site for the expected 5 years of follow-up
- Age greater than or equal to 18 years

Each site must inform BIOTRONIK whenever a lead complication has

occurred or when a patient is no longer participating.

All leads that experience a complication and are subsequently explanted and returned to BIOTRONIK undergo root-cause analyses. Product performance is analyzed as a function of time using the survival analysis method. Root causes for any failures, regardless of the incidence rates, are investigated.

7.2.3 Protego Post-Approval Registry

BIOTRONIK's Protego Post-Approval Registry is a prospective, single-arm, non-randomized registry to confirm the long-term safety and reliability of the Protego DF4 ICD lead when used in conjunction with a BIOTRONIK DF4 compatible ICD or CRT-D pulse generator. The Protego DF4 Post-Approval Registry is registered on clinicaltrials.gov under NCT02243696.

The evaluation of safety will be based on the analysis of Protego DF4 lead related or header related adverse events through a follow-up time of 5 years post-implant. To ensure a sufficiently large and representative source of data, participating clinical sites must meet pre-specified selection criteria.

Every effort is made to ensure participants are representative of the range of clinical environments in which BIOTRONIK's cardiac rhythm products are used. Patients will be seen for routine follow-up visits per their institutional standard of care until they can no longer be followed (e.g., death, lost to follow-up, etc.). However, to ensure regular patient status assessments are completed, a study follow-up schedule consistent with typical care practices has been established, with required follow-ups at 3 and 6 months post-implant, and every 6 months thereafter.

During each study visit, the following are required:

- Interrogate programmed
 parameters
- Record electrical parameters of the implanted leads. Perform all pacing threshold measurements at 0.4 ms or 0.5 ms pulse width when feasible
- Evaluate device diagnostics and programmed parameters to ensure the device is providing appropriate therapy
- Determine if there are any leadrelated, pulse generator-related or procedure related adverse events. If any are recorded, complete the Adverse Event eCRF
- Complete all appropriate eCRFs

This active surveillance model ensures a robust dataset for effectively monitoring product performance.

Patient Enrollment Criteria

To support the objectives of the study, participants are required to meet the following inclusion criteria prior to enrollment:

- Candidate for implant or successfully implanted with a BIOTRONIK ICD or BIOTRONIK CRT-D system, including a Protego lead
- Able to understand the nature of the registry and give informed consent
- Available for follow-up visits on a regular basis at the study site
- Age greater than or equal to 18 years

Each site must inform BIOTRONIK whenever a lead complication has occurred or when a patient is no longer participating.

All leads that experience a complication and are subsequently explanted and returned to BIOTRONIK undergo root-cause analyses. Product performance is analyzed as a function of time using the survival analysis method. Root causes for any failures, regardless of the incidence rates, are investigated.

7.2.4 QP ExCELs

BIOTRONIK'S QP ExCELS Clinical Study is a combined Pre-Market and Post-Approval, non-randomized, multi-center registry designed to confirm the safety and efficacy of BIOTRONIK'S Sentus QP leads in a clinical investigation to support regulatory approval as well as a longterm post-approval evaluation of the devices in the United States. The QP ExCELs Clinical Study is registered on clinicaltrials.gov under NCT02290028.

For the Pre-Market Study, the evaluation of safety is based on the analysis of Sentus QP lead-related adverse events through a follow-up time of 6 months post-implant, while the evaluation of effectiveness is based on analysis on the percentage of subjects with an acceptable LV pacing threshold in the permanently programmed vector at 3-months postimplant.

For the Post-Approval Study, the evaluation of safety will be based on the analysis of Sentus QP lead-related adverse events through a follow-up time of 5 years post-implant. To ensure a sufficiently large and representative source of data, participating clinical sites must meet pre-specified selection criteria.

Every effort is made to ensure participants are representative of the range of clinical environments in which BIOTRONIK's cardiac rhythm products are used. Patients will be seen for routine follow-up visits per their institutional standard of care until they can no longer be followed (e.g., death, lost to follow-up, etc.). However, to ensure regular patient status assessments are completed, a study follow-up schedule consistent with typical care practices has been established, which required follow-ups at discharge/wound check, 3 and 6 months post-implant, and every 6 months thereafter

Patient Enrollment Criteria

To support the objectives of the study, participants are required to meet the

following inclusion criteria prior to enrollment:

- Standard CRT-D indication according to clinical routine
- De novo implantation or upgrade from existing ICD or pacemaker implant (with no prior attempt at LV lead placement) utilizing a BIOTRONIK CRT-D system with IS4 LV port and Sentus QP LV lead
- Patient is able and willing to complete all routine study visits at the investigational site through 5 years of follow-up
- Patient is able to understand the nature of the clinical investigation and provide written informed consent
- Patient accepts Home Monitoring concept
- Age greater than or equal to 18 years

Each site must inform BIOTRONIK whenever a lead complication has occurred or when a patient is no longer participating.

All leads that experience a complication and are subsequently explanted and returned to BIOTRONIK undergo root-cause analyses. Product performance is analyzed as a function of time using the survival analysis method. Root causes for any failures, regardless of the incidence rates, are investigated.

7.3 Lead Complications

The data presented characterizes chronic lead performance by estimating lead-related complication free survival probabilities. Following industry practice, for analysis purposes, the complication criteria, which align with the AdvaMed "Industry Guidance for Uniform Reporting of Clinical Performance of Cardiac Rhythm Management Pulse Generators and Leads", are defined below.

7.3.1 GALAXY and CELESTIAL

All reported lead-related adverse events within the GALAXY and CELESTIAL Registries are classified by the reporting investigator and are adjudicated by an independent event adjudication committee. A lead related complication is considered to have occurred if a clinical observation happens after successful implantation. is classified with at least one of the following event classifications and at least one of the following clinical actions is made. Any Clinical Event without a related Clinical Action is not considered a Qualifying Complication. Events with an onset date 30 days or less after the implant are acute observations and are listed separately.

Event Classifications

- Failure to Capture
- Failure to Sense/undersensing
- Oversensing
- Abnormal Pacing Impedance (based on lead model, but normal range is typically 200- 2,000 ohms)
- Abnormal Defibrillation Impedance (based on lead model, but normal range is 25 – 150 ohms)
- Insulation Breach
- Conductor Fracture, confirmed electrically, visually or radiographically
- Extracardiac Stimulation
- Cardiac Perforation
- Lead Dislodgement

Clinical Actions

- Lead surgically abandoned/capped
- Lead electrically abandoned
- Lead explanted
- Lead replaced
- Lead conductor taken out of service
- Lead use continued based on medical judgment despite a known clinical performance issue
- Other lead-related surgery

7.3.2 SIELLO

All reported lead-related adverse events within the SIELLO Clinical Study are classified by the reporting investigator and are adjudicated by an independent event adjudication committee. A lead related complication is considered to have occurred if a clinical observation happens after successful implantation and is classified with at least one of the following event classifications. Events with an onset date 30 days or less after the implant are acute observations and are listed separately.

Event Classifications

- Failure to Capture
- Failure to Sense/undersensing
- Oversensing
- Abnormal Pacing Impedance (based on lead model, but normal range is typically 200 - 2,000 Ohm)
- Insulation Breach
- Conductor Fracture, confirmed electrically, visually or radiographically
- Extracardiac Stimulation
- Cardiac Perforation
- Lead Dislodgement

7.3.3 Protego

All reported lead-related adverse events within the Protego Registry are classified by the reporting investigator and are adjudicated by an independent event adjudication committee. A lead related complication is considered to have occurred if a clinical observation happens after successful implantation and is classified with at least one of the following event classifications. Events with an onset date 30 days or less after the implant are acute observations and are listed separately.

Event Classifications

- Failure to Capture
- Failure to Sense/undersensing
- Oversensing

- Abnormal Pacing Impedance (based on lead model, but normal range is typically 200 - 2,000 Ohm)
- Abnormal Defibrillation Impedance (based on lead model, but normal range is 25 to 150 Ohm)
- Insulation Breach
- Conductor Fracture, confirmed electrically, visually or radiographically
- Extracardiac Stimulation
- Cardiac Perforation
- Lead Dislodgement

7.3.4 QP ExCELs

All reported lead-related adverse events within the QP ExCELs registry are classified by the reporting investigator and are adjudicated by an independent event adjudication committee. A lead related complication is considered to have occurred if a clinical observation happens after successful implantation and is classified with at least one of the following event classifications. Events with an onset date 30 days or less after the implant are acute observations and are listed separately.

Event Classifications

- Failure to Capture
- Failure to Sense/undersensing
- Oversensing
- Abnormal Pacing Impedance (based on lead model, but normal range is typically 200 - 2,000 Ohm)
- Insulation Breach
- Conductor Fracture, confirmed electrically, visually or radiographically
- Extracardiac Stimulation
- Cardiac Perforation
- Lead Dislodgement

7.4 Lead Product Performance Graphs and Data

The clinical data presented on the following page is intended to show the long term clinical performance of leads based on clinical studies. The same analysis methods as described in chapter 6 are applied.

Returned Product Analysis Results

Although the returned product analysis data is not used to generate the survival estimates for the clinical data, it provides valuable insight into the causes of lead malfunction. Following the same approach as for complaint data, a malfunction is reported as described in section 6.3 of this report.

Performance of BIOTRONIK Leads Based on Clinical Study Data

8.1 Performance of Pacing Leads8.2 Performance of ICD Leads

8.1 Performance of Pacing Leads

Siello S / Solia S Study Data

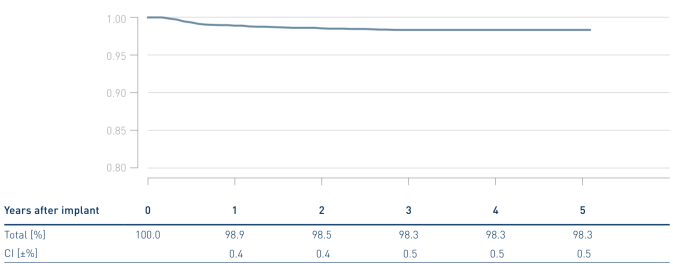
Product Versions Lead Type Polarity	45, 53, 60 straight, active fixation bipolar
Steroid	yes
U.S. Market Release	Jun 2016
CE Market Release	Jul 2009
Worldwide Distributed Devices	1023000
Registered U.S. Implants	3240

	Quantity	Rate
U.S. Qualifying Complications	51	1.57%
Abnormal Pacing Impedance	2	0.06%
Cardiac Perforation	3	0.09%
Conductor Fracture	1	0.03%
Failure to Capture	23	0.71%
Failure to Sense (undersensing)	11	0.34%
Lead Dislodgement		0.28%
Oversensing	1	0.03%
Other	1	0.03%

	Quantity	Rate
U.S. Confirmed Malfunctions	2	0.06%
Conductor Fracture	1	0.03%
Insulation Breach	1	0.03%
U.S. Acute Lead Observations	26	0.80%
Cardiac Perforation	8	0.25%
Extracardiac Stimulation	2	0.06%
Failure to Capture	6	0.19%
Failure to Sense (undersensing)	5	0.15%
Lead Dislodgement	5	0.15%

• Total survival

Cumulative survival propability



Linox SD Study Data

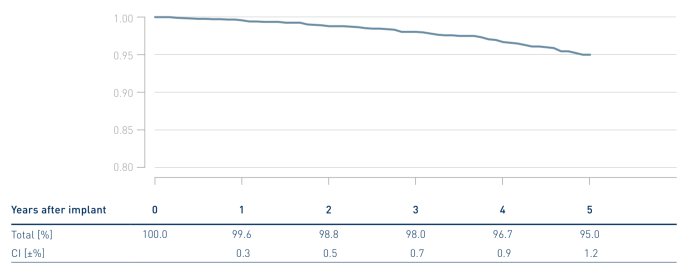
Product Versions	_ 60/16, 65/16, 65/18, 75/18
Lead Type	_ dual-coil, active fixation
Polarity	_ bipolar
Steroid	_ yes
U.S. Market Release	_ Apr 2006
CE Market Release	_ Aug 2006
Worldwide Distributed Devices	_ 55 100
Registered U.S. Implants	_ 2 272

	Quantity	Rate
U.S. Qualifying Complications	67	2.95%
Abnormal Defibrillation Impedance	4	0.18%
Abnormal Pacing Impedance	9	0.40%
Cardiac Perforation	1	0.04%
Conductor Fracture	10	0.44%
Failure to Capture	7	0.31%
Failure to Sense (undersensing)	3	0.13%
Insulation Breach	13	0.57%
Lead Dislodgement	3	0.13%
Oversensing	17	0.75%

	Quantity	Rate
U.S. Confirmed Malfunctions	23	1.01%
Conductor Fracture	3	0.13%
Insulation Breach	20	0.88%
U.S. Acute Lead Observations	9	0.40%
Cardiac Perforation	4	0.18%
Conductor Fracture	1	0.04%
Failure to Capture	2	0.09%
Lead Dislodgement	1	0.04%
Other	1	0.04%

• Total survival

Cumulative survival propability



Linox^{smart} SD Study Data

Product Versions	60/16, 65/16, 65/18, 75/18
Lead Type	dual-coil, active fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	Jan 2011
CE Market Release	Oct 2009
Worldwide Distributed Devices	55 400
Registered U.S. Implants	736

	Quantity	Rate
U.S. Qualifying Complications	27	3.67%
Abnormal Defibrillation Impedance	2	0.27%
Abnormal Pacing Impedance	1	0.14%
Conductor Fracture	3	0.41%
Failure to Capture	2	0.27%
Insulation Breach	4	0.54%
Lead Dislodgement	6	0.82%
Oversensing	9	1.22%

	Quantity	Rate
U.S. Confirmed Malfunctions	7	0.95%
Insulation Breach	7	0.95%
U.S. Acute Lead Observations	2	0.27%
Lead Dislodgement	2	0.27%

• Total survival Cumulative survival propability 0.85 -Years after implant 1 2 3 4 5 0 Total [%] 100.0 99.1 98.1 96.3 95.2 94.1 CI [±%] 0.8 1.1 1.7 2.0 2.3

114

Protego S Study Data

Product Versions	_ 60, 65, 75
Lead Type	_single-coil, active fixation
Polarity	_ bipolar
Steroid	_ yes
U.S. Market Release	_ Jul 2014
CE Market Release	_ Feb 2014
Worldwide Distributed Devices	_ 53 900
Leads registered in study	_ 1 090

	Quantity	Rate
U.S. Qualifying Complications	10	0.92%
Conductor Fracture	2	0.18%
Failure to Capture	1	0.09%
Failure to Sense	1	0.09%
Lead Dislodgement	2	0.18%
Oversensing	4	0.37%

Cumulative survival propability

• Total survival

	Quantity	Rate
U.S. Confirmed Malfunctions	7	0.64% 0.18%
Insulation Breach	5	0.46%
U.S. Acute Lead Observations	4	0.37%
Cardiac Perforation	3	0.28%
Lead Dislodgement	1	0.09%

0.85 -Т Years after implant 1 2 3 4 0 Total [%] 100.0 99.5 99.3 99.0 98.7 CI [±%] 0.5 0.6 0.7 0.9

115

Protego SD Study Data

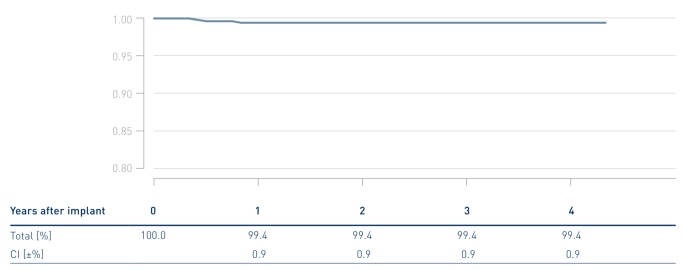
Product Versions	_ 60/16, 65/16, 65/18, 75/18
Lead Type	_ dual-coil, active fixation
Polarity	_ bipolar
Steroid	_ yes
U.S. Market Release	_ Jul 2014
CE Market Release	_ May 2013
Worldwide Distributed Devices	_ 18 500
Leads registered in study	_ 533

	Quantity	Rate
U.S. Qualifying Complications	3	0.56%
Abnormal Defibrillation Impedance	1	0.19%
Conductor Fracture	1	0.19%
Failure to Capture	1	0.19%

	Quantity	Rate
U.S. Confirmed Malfunctions	1 1	0.19% 0.19%
U.S. Acute Lead Observations	2	0.38% 0.38%

• Total survival

Cumulative survival propability



Corox OTW Study Data

Product Versions	75, 85
Lead Type	helix fixation
Polarity	bipolar
Steroid	yes
U.S. Market Release	May 2008
CE Market Release	Dec 2006
Worldwide Distributed Devices	28800
Leads registered in study	_ 696

	Quantity	Rate
U.S. Qualifying Complications	35	5.03%
Abnormal Pacing Impedance	6	0.86%
Conductor Fracture	5	0.72%
Extracardiac Stimulation	3	0.43%
Failure to Capture	5	0.72%
Lead Dislodgement	16	2.30%

	Quantity	Rate
U.S. Confirmed Malfunctions	6	0.86%
Conductor Fracture	6	0.86%
U.S. Acute Lead Observations	4	0.57%
Extracardiac Stimulation	1	0.14%
Lead Dislodgement	3	0.43%

Cumulative survival propability 0.95 -0.85 -Years after implant 1 2 3 4 5 0 Total [%] 100.0 98.2 96.9 95.2 94.2 93.0 1.4 CI [±%] 1.1 1.8 2.1 2.3

• Total survival

Corox OTW-L Study Data

Product Versions	_ 75, 85
Lead Type	_dual-curve fixation
Polarity	_ bipolar
Steroid	_yes
U.S. Market Release	_ Jan 2011
CE Market Release	_ Dec 2009
Worldwide Distributed Devices	_ 31 800
Leads registered in study	_ 699

	Quantity	Rate
U.S. Qualifying Complications	21	3.00%
Extracardiac Stimulation	4	0.57%
Failure to Capture	7	1.00%
Lead Dislodgement	10	1.43%

• Total survival

	Quantity	Rate
U.S. Confirmed Malfunctions	0	0.00%
U.S. Acute Lead Observations	4	0.57%
Extracardiac Stimulation	3	0.43%
Lead Dislodgement	1	0.14%

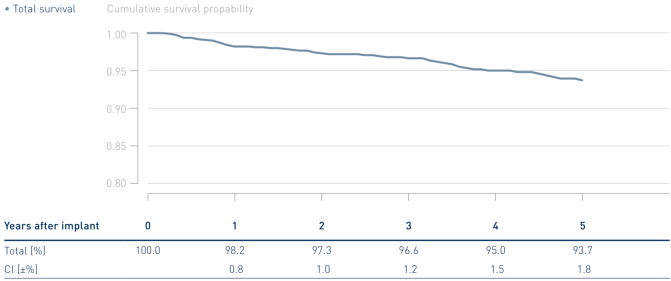
Cumulative survival propability 0.95 -0.85 -Years after implant 0 1 2 3 4 5 Total [%] 100.0 98.5 97.1 96.7 96.1 96.1 CI [±%] 1.0 1.4 1.5 1.7 1.7

Corox OTW-S Study Data

Product Versions	75,85
Lead Type	thread fixation
Polarity	_ bipolar
Steroid	yes
U.S. Market Release	_ May 2008
CE Market Release	Dec 2006
Worldwide Distributed Devices	26 500
Leads registered in study	_ 1 1 4 1

	Quantity	Rate
U.S. Qualifying Complications	48	4.21%
Abnormal Pacing Impedance	12	1.05%
Extracardiac Stimulation	9	0.79%
Failure to Capture		0.79%
Lead Dislodgement	18	1.58%

	Quantity	Rate
U.S. Confirmed Malfunctions		0.09% 0.09%
U.S. Acute Lead Observations	6	0.53%
Extracardiac Stimulation	1	0.09%
Failure to Capture	1	0.09%
Lead Dislodgement	4	0.35%



Cumulative survival propability

Sentus OTW QP S Study Data

Product Versions	_ 75, 75/49, 85, 85/49
Lead Type	_thread fixation
Polarity	_ quadripolar
Steroid	_yes
U.S. Market Release	_ May 2017
CE Market Release	_Dec 2014
Worldwide Distributed Devices	_ 11 900
Leads registered in study	_ 374

	Quantity	Rate
U.S. Qualifying Complications	14	3.74%
Conductor Fracture	1	0.27%
Extracardiac Stimulation	1	0.27%
Failure to Capture	2	0.53%
Lead Dislodgement	10	2.67%

	Quantity	Rate
U.S. Confirmed Malfunctions	2	0.53% 0.53%
U.S. Acute Lead Observations	8	2.14%
Cardiac Perforation	1	0.27%
Failure to Capture	1	0.27%
Lead Dislodgement	6	1.60%

0.85 -Years after implant 1 2 3 0 Total [%] 100.0 96.0 95.5 95.5 CI [±%] 2.2 2.4 2.4

• Total survival

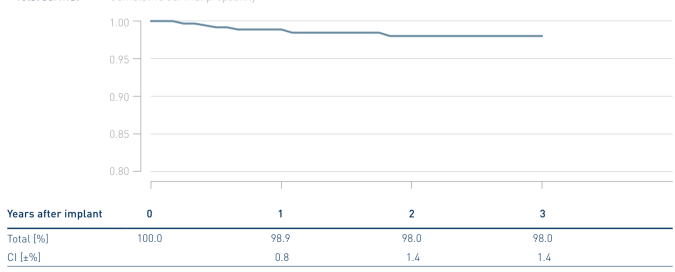
Cumulative survival propability

Sentus OTW QP L Study Data

Product Versions	75, 75/49, 85, 85/49
Lead Type	_dual-curve fixation
Polarity	_quadripolar
Steroid	_ yes
U.S. Market Release	_ May 2017
CE Market Release	_Dec 2014
Worldwide Distributed Devices	_ 44 200
Leads registered in study	_ 964

	Quantity	Rate
U.S. Qualifying Complications	12	1.24%
Abnormal Pacing Impedance	1	0.10%
Extracardiac Stimulation	2	0.21%
Failure to Capture	3	0.31%
Lead Dislodgement	6	0.62%

	Quantity	Rate
U.S. Confirmed Malfunctions	1	0.10% 0.10%
U.S. Acute Lead Observations		0.62%
Extracardiac Stimulation	2	0.21%
Failure to Capture	3	0.31%
Lead Dislodgement	1	0.10%



• Total survival Cumulative survival propability



9. Advisories

Stratos LV-T Potentially defective low voltage capacitors

84 devices world-wide, none in the United States

Status Update As of July 2018

- No reports of malfunctioning pacemakers associated with this capacitor lot were received.
- No reports of deaths or serious injuries were received associated with this advisory.
- An extended test program yielded no malfunctions of the capacitors under question.

Original communication July 2006

A limited number of our Stratos LV-T pacemakers may exhibit a device malfunction resulting in increased battery current and in loss of output, very likely simultaneously on all channels.

BIOTRONIK has identified a specific batch of low voltage capacitors from a single component supplier as the root cause. Please be assured that the capacitors from this batch are no longer being used in any BIOTRONIK device, and that no other pacemaker or ICD manufactured by BIOTRONIK is affected.

To date, no field reports related to this phenomenon were received.

The anomaly is limited to 84 devices world-wide. BIOTRONIK has taken immediate action to retrieve all potentially affected devices not yet delivered to hospitals. According to our records, 14 devices were delivered to hospitals and may have been implanted.

Based on preliminary data received from the component supplier, the projected rate of occurrence is expected to less than 1 out of 1000 devices.

Our records show that you are the implanting and/or follow-up physician for one or more patients with an affected device. We recommend the following actions:

- If Home Monitoring is **activated**, please check the Cardio Report and the pacing impedance history. A rapid and significant increase of the pacing impedance on one or more channels may indicate a possible loss of output. In this case we recommend to schedule a patient follow-up as soon as possible.
- If Home Monitoring is **not activated** or pacing impedance data from Cardio Reports are not available we recommend to schedule a patient follow-up as soon as possible.
- During a follow-up session
 please perform the "Battery Lead
 Telemetry" test. A significantly
 increased pacing impedance on one
 or more channels or a significantly
 increased battery current may
 indicate a possible loss of output.
- If no anomaly is detected please activate Home Monitoring. You may use regular follow-up intervals if Home Monitoring is used for remote monitoring between follow-ups.
- In any case we recommend to replace the device of a pacemakerdependent patient without underlying intrinsic rhythm.

We have tried to be as specific as possible with our recommendations. As always, individual circumstances and medical judgement determine decisions about patient care, the urgency of follow-ups and possible replacement of devices.

If a Stratos LV-T is replaced for the reason explained in this notification, BIOTRONIK will provide a replacement device consistent with the terms of our warranty. For activation of the warranty, please return the explanted device to BIOTRONIK within 30 days of the explantation.

X-Ray Identifiers for Pacemakers and ICDs

Pacemaker/ICD Product Versions	X-Ray ID
Cylos DR, DR-T, VR	RZ
Cylos 990 DR, DR-T, VR	FV
Edora 8 DR, DR-T	•
Entovis DR, DR-T, SR, SR-T	SF
Eluna 8 DR, DR-T	SF
Estella SR, SR-T, DR, DR-T	SF
Etrinsa 8 SR-T, DR-T, HF-T	SF
Evia DR, DR–T, SR, SR-T, HF, HF-T	SF
Iforia 7 VR-T DX, DR-T	NT
Ilesto 7 DR-T, HF-T, VR-T DX, VR-T, DR-T DF4	NT
Ilivia 7 VR-T, DR-T, VR-T DX, VR-T DF4, HF-T DF4	NK
Inventra 7 VR-T DX	AH
Iperia 7 VR-T, DR-T, VR-T DX, VR-T DF4, HF-T DF4	NH
Itrevia 7 VR-T, DR-T, VR-T DX, DR-T DF4, HF-T, HF-T DF4	NH
Lumax 340 DR-T, HF-T, VR-T	HR
Lumax 540 DR-T, HF-T, VR-T	SH
Lumax 740 DR-T, HF-T, VR-T, VR-T DX	RH
Philos DR, D, SLR, SR, S	LE
Philos DR-T	VV
Philos II DR, D, S, SLR, SR	ET
Philos II DR-T	KP
Stratos LV, LV-T	SV
Talos DR, D, SLR, SR, S	PV

Contacting BIOTRONIK

Regarding this Report

BIOTRONIK invites your suggestions and questions related to this Product Performance Report. Please send your comments to:

PPR Support Team

Phone +49 (0) 30 68905 1368 Fax +49 (0) 30 68905 96 1920 E-mail PPR@biotronik.com

Address

BIOTRONIK SE & Co. KG Attn: Quality Patient Safety Woermannkehre 1 12359 Berlin, Germany

Regarding Products

BIOTRONIK invites customers to call the following locations with suggestions, comments or specific questions related to BIOTRONIK products:

Worldwide CRM Product Support

Phone + 49 (0) 30 68905 1133 Fax + 49 (0) 30 68905 1960 Email product.support@biotronik.com

Address

BIOTRONIK SE & Co. KG Attn: Product Support Woermannkehre 1 12359 Berlin, Germany

Within the U.S.:

Phone (800) 284 6689 Fax (800) 387 2681 E-mail technical.services@biotronik.com

Address

BIOTRONIK, Inc. Attn: Technical Services 6024 Jean Road Lake Oswego, OR 97035