Implantable Cardioverter Defibrillators Registry v1.08

	A. Participant A	Administration
Field Name:	Participant ID	Seq No: 1000
Short Name:	PartId	Core: Yes
Status:	New	Harvested: Yes
Format:	Integer	
Data Source:	Automatic	
Parent Element:		
Parent Value:		
Missing Data:	lllegal	
Valid Range:	1-999999	
Usual Range:		
•	Participant ID	
Definition:	-NCDR Participant is defined as one	signed to each Participant by the ACC-NCDR. An ACC entity that signs a Participation Agreement with the le to the harvest, and gets back one Outcomes Report
		o harvest must be in one data submission file. If one han one file (e.g. at two sites), then the data must be on file for the harvest.
		ngle purchased software, and enter cases into one orted into different data submisison files, one for each
Selections:		
	Participant Name	Seq No: 1010
Short Name:		Core: Yes
Status:		Harvested: Yes
	Text (100)	
Data Source:	Automatic	
Parent Element:		
Parent Value:		
Missing Data:	llegal	
Valid Range:		
Usual Range:	Destining of News	
•	Participant Name	
Definition:		acility where the implant procedure was performed. names with no abbreviations or variations in spelling
Selections:		
Field Name:	Medicare Provider Number	Seq No: 1015
Short Name:	MPN	Core: Yes
Status:	New	Harvested: Yes

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Format:	Text (6)	
Data Source:		
Parent Element:		
Parent Value:		
Missing Data:	Report	
Valid Range:		
Usual Range:		
Description:	Medicare Provider Number	
Definition:	Indicate the medicare provider number of the facility at implant.	which the patient received the
Selections:		
Field Name:	Participant NPI	Seq No: 1016
Short Name:	PartNPI	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (10)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	Report	
Valid Range:		
Usual Range:		
Description:	Participant National Provider Identifier	
Definitions		
Definition:	Indicate the hospital's (N)ational (P)rovider (I)dentifier. uniquely identify hospitals for Medicare billing purposes	
Selections:		
Selections:		
Selections:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission	
Selections: Field Name:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe	Seq No: 1020
Selections: Field Name: Short Name:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6)	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status: Format:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6)	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6)	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic Illegal	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic	Seq No: 1020 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic Illegal	Seq No: 1020 Core: Yes Harvested: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic Illegal Timeframe of Data Submission Indicate the timeframe of data included in the data subm	Seq No: 1020 Core: Yes Harvested: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic Illegal Timeframe of Data Submission Indicate the timeframe of data included in the data subm	Seq No: 1020 Core: Yes Harvested: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition:	uniquely identify hospitals for Medicare billing purposes Timeframe of Data Submission Timeframe New Text (6) Automatic Illegal Timeframe of Data Submission Indicate the timeframe of data included in the data subm 2005Q4 Transmission Number	Seq No: 1020 Core: Yes Harvested: Yes

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Format:	Integer	
Data Source:	-	
Parent Element:		
Parent Value:		
Missing Data:	Illegal	
-	1-999999999	
Usual Range:		
-	Transmission Number	
•		Illy inserted by the software. It identifies the
	number of times the software has created	d data submission files. The transmission number the data submission files are exported. The
Selections:		
Field Name:	Software Vendor Identifier	Seq No: 1050
Short Name:	Vendorld	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (15)	
Data Source:	Automatic	
Parent Element:		
Parent Value:		
Missing Data:	Illegal	
Valid Range:		
Usual Range:		
Description:	Software Vendor Identifier	
Definition:	the ACC) to identify software vendor. Ver	bon by mutual selection between the vendor and adors must use consistent name identification dentification must be approved by the ACC.
Selections:		
Field Name:	Vendor software version	Seq No: 1060
Short Name:	VendorVer	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (20)	
Data Source:	Automatic	
Parent Element:		
Parent Value:		
Missing Data:	Illegal	
Valid Range:		
Usual Range:		
Description:	Vendor software version	
Definition:		sion number identifying the software which created r controls the value in this field. Version passing at the ACC.
Selections:		

Implantable Cardioverter Defibrillators Registry v1.08

Field Name:	Registry Identifier	Seq No: 1070
Short Name:	RegistryId	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (20)	
Data Source:	Automatic	
Parent Element:		
Parent Value:		
Missing Data:	Illegal	
Valid Range:		
Usual Range:		
Description:	Registry Identifier	
Definition:		escribes which ACC data registry these records apply. e time the data is collected and the records are ma automatically by software.
Selections:		
Field Name:	Registry Version	Seq No: 1080
Short Name:	RegistryVer	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (10)	
Data Source:	Automatic	
Parent Element:		
Parent Value:		
Missing Data:	Illegal	
Valid Range:		
Usual Range:		
Description:	Registry Version	
Definition:	which each record conforms. It identified valid data for each field. It is the versi	n number of the Data Specifications/Dictionary, to ies which fields should have data, and what are the on implemented in the software at the time the data is . This is entered into the schema automatically by
Selections:		
Field Name:	Patient Population	Seq No: 1090
Short Name:	PatientPop	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (Categorical)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	Illegal	
Valid Range:		
Usual Range:		
Description:	Patient Population	

Implantable Cardioverter Defibrillators Registry v1.08

	registry.		ed in the data export and submitted tothe
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	All Patients	All patients regardless of insurance payor and ICD Indication.
	2	CMS Primary Prevention Patients	Patients with a Primary or Secondaryinsurance payor of"Medicare"andan ICD Indicationof"Primary Prevention".
Field Name:	Data Submiss	ion File Password	Seq No: 1100
Short Name:	Password		Core: Yes
Status:	New		Harvested: No
Format:	Text (20)		
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	No action		
Valid Range:			
Usual Range:			
-		ion File Password	
Description:	Data Submiss	ion File Password ACC assigned password that should be ap	plied to the data submission zip file.
Description:	Data Submiss Indicates the		plied to the data submission zip file.
Description: Definition:	Data Submiss Indicates the		plied to the data submission zip file. Seq No: 1110
Description: Definition: Selections:	Data Submiss Indicates the Auxiliary 0		· · ·
Description: Definition: Selections: Field Name:	Data Submiss Indicates the Auxiliary 0 Aux0		Seq No: 1110
Description: Definition: Selections: Field Name: Short Name: Status:	Data Submiss Indicates the Auxiliary 0 Aux0		Seq No: 1110 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status:	Data Submiss Indicates the A Auxiliary 0 Aux0 New Text (50)		Seq No: 1110 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format:	Data Submiss Indicates the A Auxiliary 0 Aux0 New Text (50)		Seq No: 1110 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source:	Data Submiss Indicates the Auxiliary 0 Aux0 New Text (50) Automatic		Seq No: 1110 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element:	Data Submiss Indicates the A Auxiliary 0 Aux0 New Text (50) Automatic		Seq No: 1110 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value:	Data Submiss Indicates the A Auxiliary 0 Aux0 New Text (50) Automatic		Seq No: 1110 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	Data Submiss Indicates the A Auxiliary 0 Aux0 New Text (50) Automatic No action		Seq No: 1110 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	Data Submiss Indicates the A Auxiliary 0 Aux0 New Text (50) Automatic No action Auxiliary 0	ACC assigned password that should be ap	Seq No: 1110 Core: Yes Harvested: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	Data Submiss Indicates the A Auxiliary 0 Aux0 New Text (50) Automatic No action Auxiliary 0 Not for partici	ACC assigned password that should be ap	Seq No: 1110 Core: Yes

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Implantable Cardioverter Defibrillators Registry v1.08 Data Dictionary - Full Specifications

B. Demographics Field Name: Patient Last Name Seq No: 2000 Short Name: Surname Core: Yes Harvested: Yes Status: New Format: Text (25) Data Source: Client Parent Element: **Parent Value:** Missing Data: Report Valid Range: **Usual Range: Description:** Patient Last Name Definition: Indicate the patient's last name. Selections: Field Name: Patient First Name Seq No: 2010 Short Name: GivenName Core: Yes Harvested: Yes Status: New Format: Text (25) Data Source: Client **Parent Element: Parent Value:** Missing Data: Report Valid Range: **Usual Range:** Description: Patient First Name Definition: Indicate the patient's first name. Selections: Field Name: Patient Middle Name Seq No: 2020 Short Name: MiddleName Core: Yes Status: New Harvested: Yes Format: Text (25) Data Source: Client Parent Element: **Parent Value:** Missing Data: Report Valid Range: **Usual Range: Description:** Patient Middle Name Definition: Indicate the patient's middle name or middle initial.

Selections:

Implantable Cardioverter Defibrillators Registry v1.08

Field Name:	Patient SSN	Seq No: 2030
Short Name:		Core: Yes
Short Name. Status:		Harvested: Yes
		Harvested: Yes
Format:		
Data Source:	Client	
Parent Element:		
Parent Value:	Deport	
Missing Data:	Report	
Valid Range:		
Usual Range:	Datiant SSN	
Description:		as Social Socurity Number (SSN). If the actions
	does not have a US assigned SSN, then I	es Social Security Number (SSN). If the patient eave the SSN blank.
Selections:		
Field Name:	Unique Patient ID	Seq No: 2040
Short Name:	PatientId	Core: Yes
Status:	New	Harvested: Yes
Format:	Integer	
Data Source:	Automatic	
Parent Element:		
Parent Value:		
Missing Data:	Illegal	
Valid Range:	1-999999999	
Usual Range:		
Description:	Unique Patient ID	
Definition:	uniquely identifies each patient. Once ass	Table ID like SSN or Medical Record Number) that igned to a patient at a health care facility, this will rent patient. If a patient returns to the same this same unique patient identifier.
Selections:		
Field Name:	Other ID	Seq No: 2045
Short Name:	OtherId	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (50)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	No action	
Valid Range:		
Usual Range:		
Description:	Other ID	
Definition:	An additional 'optional' patient identifier, s associated with the patient.	uch as medical record number, that can be

Implantable Cardioverter Defibrillators Registry v1.08

Selections:			
Field Name:	Patient DOB		Seq No: 2050
Short Name:	DOB Core: Yes		
Status:	New	Ha	arvested: Yes
Format:	Date (mm/dd/	уууу)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:		> 01/01/1850 and Patient DOB < Previous IC	D Date and Patient DOB < Previous
		nd Patient DOB < Admission Date	
Usual Range:			
•	Patient date c		
Definition:	Indicate the p	atient's date of birth.	
Selections:			
Field Name:	Gender		Seq No: 2060
Short Name:	Gender		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:			
Usual Range:			
Description:	Gender		
Definition:	Indicate the p	atient's gender at birth.	
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	ooung/oon		
	1	Male	
	2	Female	
Field Name:	Race		Seq No: 2070
Short Name:	Race		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	Text (Categorical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:	-		
Usual Range:			
Description:			
	-		

Implantable Cardioverter Defibrillators Registry v1.08

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	White	
	2	Black/African American	
	4	Asian	
	5	American Indian/Alaska Native	
	6	Native Hawaiian	
	7	Other	
Field Name:	Hispanic Ethn	icity	Seq No: 2075
Short Name:	•		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:			
Usual Range:			
Description:	Hispanic Ethn	icity	
Definition:	Indicate if the	patient is of hispanic ethnicity.	
Selections:	Coding/Sort	Coding/Sort Selection(Choose one) Explanation	
	0	No	
	1	Yes	
Field Name:	Auxiliary 1		Seq No: 2080
Short Name:	-		Core: Yes
	Aux1	Ha	
Short Name: Status:	Aux1	Ha	Core: Yes
Short Name: Status:	Aux1 New Text (50)	Ha	Core: Yes
Short Name: Status: Format: Data Source:	Aux1 New Text (50)	Ha	Core: Yes
Short Name: Status: Format: Data Source:	Aux1 New Text (50)	Ha	Core: Yes
Short Name: Status: Format: Data Source: Parent Element:	Aux1 New Text (50) Client	Ha	Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value:	Aux1 New Text (50) Client	Ha	Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data:	Aux1 New Text (50) Client	H	Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range:	Aux1 New Text (50) Client No action	H	Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	Aux1 New Text (50) Client No action Auxiliary 2 For participan	Hat use only. A 50 character text field that may ent or admission.	Core: Yes arvested: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	Aux1 New Text (50) Client No action Auxiliary 2 For participan	t use only. A 50 character text field that may	Core: Yes arvested: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition:	Aux1 New Text (50) Client No action Auxiliary 2 For participan about the pati	t use only. A 50 character text field that may	Core: Yes arvested: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Usual Range: Description: Definition:	Aux1 New Text (50) Client No action Auxiliary 2 For participan about the pati	t use only. A 50 character text field that may	Core: Yes arvested: Yes be used to collect additional information

Implantable Cardioverter Defibrillators Registry v1.08

Format:	Text (50)
Data Source:	Client
Parent Element:	
Parent Value:	
Missing Data:	No action
Valid Range:	
Usual Range:	
Description:	Auxiliary 2
Definition:	For participant use only. A 50 character text field that may be used to collect additional information about the patient or admission.
Selections:	

Implantable Cardioverter Defibrillators Registry v1.08

	C. Ac	Imission	
Field Name:	Admission Date	Seq No: 3000	
Short Name:	AdmitDate	Core: Yes	
Status:	New	Harvested: Yes	
Format:	Date (mm/dd/yyyy)		
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	-		
Valid Range:	Admission Date > Previous ICD Date and Admission Date > PrevCABGDate and Admission Date <= Date of Implant		
Usual Range:			
Description:	Admission Date		
	Indicate the date on which the pati	ent was admitted to the hospital for the current stay.	
Selections:			
Field Name:	Date of Implant	Seq No: 3010	
Short Name:	ImplantDate	Core: Yes	
Status:	New	Harvested: Yes	
Format:	Date (mm/dd/yyyy)		
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	lllegal		
Valid Range:	Date of Implant >= Admission Date and Date of Implant <= Date of Discharge		
Usual Range:			
Description:	ICD Implant Date		
Definition:	Indicate the date of the ICD implar	ıt.	
		Ds were implanted/explanted during a single admission, D implant. For clarification, see Sequence Numbers 3507,	
Selections:			
Field Name:	Insurance Payor-Primary	Seq No: 3020	
Short Name:	PayorPrim	Core: Yes	
Status:	New	Harvested: Yes	
Format:	Text (Categorical)		
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Illegal		
Valid Range:			

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Usual Range:

Description: Insurance Payor-Primary

Definition: Indicate the patient's primary insurance payor.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	Government	* RYHOLP HQAVUHHUV VR SDALHQAV ZKR DUH FRYHUHGE \JRYHOLP HQAVUHLP EXUAHG FDUH ,QVKH 8 6 WALV LQFOXGHV 0 HQEDUH 0 HQEDLG LQFOXGQJ DOOMAAM IHGHUDO 0 HQEDLG WSH SURJUDP V 7 UKLDUH WAH 9 HAMUDQV \$ GPLQLAADMARQ + HDOMK 3 ODQ DQG)HGHUDQ P SOR\HHTV,QAXUDQFH
	2	Commercial	Commercial refers to all indemnity (fee- for-service) carriers and Preferred Provider Organizations (PPOs) (e.g. Blue Cross/Blue Shield).
	3	НМО	HMO refers to a Health Maintenance Organization (HMO) characterized by coverage that provides health services for members on a pre-paid basis.
	4	Non-U.S. Insurance	Non-US Insurance refers to individuals who reside in and have health insurance in another country.
	5	None/Self Pay	None/Self Pay refers to individuals with no or limited health insurance; thus, the individual is the payor regardless of ability to pay.

Seq No: 3025

Harvested: Yes

Core: Yes

Field Name: Government Type-Primary

Short Name: GovTypePrim

Status: New

Format: Text (Categorical)

Data Source: Client

Parent Element: Insurance Payor-Primary(3020)

Parent Value: Government

Missing Data: Illegal

Valid Range:

Usual Range:

Description: Government Insurance Type-Primary

Definition: Indicate the type of insurance if the patient's primary insurance payor is Government.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	Medicare	
	2	Medicaid	
	3	TriCare	
	4	Veteran's Administration Health Plan	
	5	Federal Employee Insurance	

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Field Name: Insurance Payor-Secondary	Seq No: 3027
Short Name: PayorSecond	Core: Yes
Status: New	Harvested: Yes
Format: Text (Categorical)	
Data Source: Client	
Parent Element:	
Parent Value:	
Missing Data: Illegal	
Valid Range:	
Usual Range:	

Description: Insurance Payor-Secondary

Definition: Indicate the patient's secondary insurance payor.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	Government	* RYHOLP HQWUHHU/WRSDWL+QWZKRDUH FRYHUHGE\JRYHOLP HQWUHLP EXUMHGFDUH ,QWRH86 WAL/LQFOXGH/0HGFDUH 0HGLFDLG LQFOXGQJDOOMMAHIHGHUDO 0HGLFDLGWSHSURJUDP V7UL&DUHWAH 9HMAUDQV\$GPLQLMUMURQ+HDOMR3ODQDQG)HGHUDQ(PSOR\HHTV,QMXUDQFH
	2	Commercial	Commercial refers to all indemnity (fee- for-service) carriers and Preferred Provider Organizations (PPOs) (e.g. Blue Cross/Blue Shield).
	3	НМО	HMO refers to a Health Maintenance Organization (HMO) characterized by coverage that provides health services for members on a pre-paid basis.
	4	Non-U.S. Insurance	Non-US Insurance refers to individuals who reside in and have health insurance in another country.
	5	None/Self Pay	None/Self Pay refers to individuals with no or limited health insurance; thus, the individual is the payor regardless of ability to pay.

Field Name: Government Type-Secondary Short Name: GovTypeSecond Status: New Format: Text (Categorical) Data Source: Client Parent Element: Insurance Payor-Secondary(3027) Parent Value: Government Missing Data: Illegal Valid Range: Usual Range: Seq No: 3029 Core: Yes Harvested: Yes

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Implantable Cardioverter Defibrillators Registry v1.08

Definition:	Indicate the ty	pe of insurance if the patient's secondary in	
	Coding/Sort		Explanation
	1	Medicare	
	2	Medicaid	
	3	TriCare	
	4	Veteran's Administration Health Plan	
	5	Federal Employee Insurance	
Field Name:	Reason for A	dmission	Seq No: 3030
Short Name:	AdmissionRea	ason	Core: Yes
Status:	New	F	larvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	•		
Valid Range:			
Usual Range:			
Description			
-	Reason for A		
Definition:	Indicate the p	dmission rimary reason the patient was hospitalized fo	or this admission.
-	Indicate the p		or this admission. Explanation
Definition:	Indicate the p	rimary reason the patient was hospitalized fo	
Definition:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized for Selection(Choose one)	Explanation Admitted for ICD implantation.
Definition:	Indicate the p Coding/Sort 1	rimary reason the patient was hospitalized fo Selection(Choose one) Admitted for this Procedure	ExplanationAdmitted for ICD implantation.Admitted for management of heart failur other than implantation of an ICD.
Definition:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized fo Selection(Choose one) Admitted for this Procedure Cardiac-CHF	ExplanationAdmitted for ICD implantation.Admitted for management of heart failur other than implantation of an ICD.Admitted for a cardiac reason other than heart failure or implantation of an ICD.
Definition: Selections:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other	ExplanationAdmitted for ICD implantation.Admitted for management of heart failur other than implantation of an ICD.Admitted for a cardiac reason other than heart failure or implantation of an ICD.Admitted for a non-cardiac reason other than implantation of an ICD.
Definition: Selections: Field Name:	Indicate the p Coding/Sort 1 2 3 4 Auxiliary 3	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other	ExplanationAdmitted for ICD implantation.Admitted for management of heart failur other than implantation of an ICD.Admitted for a cardiac reason other than heart failure or implantation of an ICD.Admitted for a non-cardiac reason other than implantation of an ICD.Admitted for a non-cardiac reason other than implantation of an ICD.Seq No: 3040
Definition: Selections: Field Name: Short Name:	Indicate the p Coding/Sort 1 2 3 4 Auxiliary 3 Aux3	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failur other than implantation of an ICD. Admitted for a cardiac reason other than heart failure or implantation of an ICD. Admitted for a non-cardiac reason other than implantation of an ICD. Seq No: 3040 Core: Yes
Definition: Selections: Field Name: Short Name: Status:	Indicate the p Coding/Sort 1 2 3 4 Auxiliary 3 Aux3 New	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	ExplanationAdmitted for ICD implantation.Admitted for management of heart failur other than implantation of an ICD.Admitted for a cardiac reason other than heart failure or implantation of an ICD.Admitted for a non-cardiac reason other than implantation of an ICD.Admitted for a non-cardiac reason other than implantation of an ICD.Seq No: 3040
Definition: Selections: Field Name: Short Name: Status:	Indicate the p Coding/Sort 1 2 3 4 Auxiliary 3 Aux3 New Text (50)	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failur other than implantation of an ICD. Admitted for a cardiac reason other that heart failure or implantation of an ICD. Admitted for a non-cardiac reason other than implantation of an ICD. Seq No: 3040 Core: Yes
Definition: Selections: Field Name: Short Name: Status: Format: Data Source:	Indicate the p Coding/Sort 1 2 3 4 Auxiliary 3 Aux3 New Text (50) Client	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failur other than implantation of an ICD. Admitted for a cardiac reason other than heart failure or implantation of an ICD. Admitted for a non-cardiac reason other than implantation of an ICD. Seq No: 3040 Core: Yes
Definition: Selections: Field Name: Short Name: Status: Format: Data Source:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failur other than implantation of an ICD. Admitted for a cardiac reason other that heart failure or implantation of an ICD. Admitted for a non-cardiac reason other than implantation of an ICD. Seq No: 3040 Core: Yes
Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failur other than implantation of an ICD. Admitted for a cardiac reason other that heart failure or implantation of an ICD. Admitted for a non-cardiac reason other than implantation of an ICD. Seq No: 3040 Core: Yes
Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failu other than implantation of an ICD. Admitted for a cardiac reason other tha heart failure or implantation of an ICD. Admitted for a non-cardiac reason othe than implantation of an ICD. Seq No: 3040 Core: Yes
Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failu other than implantation of an ICD. Admitted for a cardiac reason other tha heart failure or implantation of an ICD. Admitted for a non-cardiac reason othe than implantation of an ICD. Seq No: 3040 Core: Yes
Definition: Selections: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range:	Indicate the p Coding/Sort	rimary reason the patient was hospitalized for Selection(Choose one) Admitted for this Procedure Cardiac-CHF Cardiac-Other Non-Cardiac	Explanation Admitted for ICD implantation. Admitted for management of heart failu other than implantation of an ICD. Admitted for a cardiac reason other tha heart failure or implantation of an ICD. Admitted for a non-cardiac reason othe than implantation of an ICD. Seq No: 3040 Core: Yes

Implantable Cardioverter Defibrillators Registry v1.08

Selections:		
Field Name:	Auxiliary 4	Seq No: 3050
Short Name:	Aux4	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (50)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	No action	
Valid Range:		
Usual Range:		
Description:	Auxiliary 4	
Definition:	For participant use only. information about the participant	A 50 character text field that may be used to collect additional atient or admission.
Selections:		

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

D. History and Risk Factors			
Field Name:	Syncope		Seq No: 3060
Short Name:	Syncope		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:			
Usual Range:			
Description:	Syncope		
Definition:	related to ane	patient had a sudden loss of consciousness, sthesia) with spontaneous recovery as report ce syncope when supine.	
	Note: Patient	history is defined as any time prior to the date	e of implant.
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
Field Name:	Family Hx Su	dden Death	Seq No: 3070
Short Name:	FHSudDeath		Core: Yes

Missing Data: Report Valid Range:

Format: Text (Categorical)

vallu Kallye.

Parent Element: Parent Value:

Status: New

Data Source: Client

Usual Range:

Description: Family History of Sudden Cardiac Death

Definition: Indicate if the patient has a known family history (parent or sibling) of sudden cardiac death.

Sudden cardiac death is defined as a natural death due to cardiac causes heralded by abrupt loss of consciousness, occurring before 75 years of age. The time and mode of death are unexpected even though preexisting heart disease may have been known to be present. Traumatic death subsequently proven to be due to sudden loss of control due to a cardiac problem is included. Coding Exception: If the patient is adopted, or the family history is unavailable, code "No".

Harvested: Yes

Note: Patient history is defined as prior to the current admission.

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Implantable Cardioverter Defibrillators Registry v1.08

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
Field Name:	-		Seq No: 3080
Short Name:			Core: Yes
Status:			arvested: Yes
	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:	_		
Missing Data:	Report		
Valid Range:			
Usual Range:	.		
-	Congestive H	eart Failure patient has a history of congestive heart failu	
	following: 1. Paroxysma 2. Dyspnea or 3. Chest X-Ra 4. Pedal eden	es physician documentation of the CHF histor I nocturnal dyspnea (PND); n exertion (DOE) due to heart failure; or ny (CXR) showing pulmonary congestion; na or dyspnea treated with medical therapy for history is defined as any time prior to the date	or heart failure.
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
Field Name:	CHF Duration		Seq No: 3090
Short Name:			Core: Yes
Status:		н	arvested: Yes
	Text (Categor		
Data Source:	· •		
Parent Element:			
Parent Value:	, ,		
Missing Data:			
Valid Range:	·		
Usual Range:			
-	Congestive H	eart Failure Duration	
-	-	ne since the initial CHF diagnosis.	
		ludes any time prior to date of implant.	

Implantable Cardioverter Defibrillators Registry v1.08

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	Within the past 3 months	
	2	3 to 9 months	
	3	Greater than 9 months	
Field Name:	Prior CHF Ho	spitalization	Seq No: 3095
Short Name:	PriorCHFHos	p	Core: Yes
Status:	New		Harvested: Yes
Format:	Text (Categor	rical)	
Data Source:	Client		
arent Element:	CHF(3080)		
Parent Value:	Yes		
Missing Data:	Report		
Valid Range:			
Usual Range:			
-	-	tive Heart Failure Hospitalization	
Definition	Indicate if the	patient has ever been hospitalized for	CHE prior to this admission Indicate the
20mmillion.		sociated with that hospitalization.	
	timeframe ass Note: This tim		on. The intent of this field is to capture
Selections:	timeframe ass Note: This tim hospitalization	sociated with that hospitalization. neframe does NOT include this admissions for CHF excluding the current admis	on. The intent of this field is to capture
	timeframe ass Note: This tim hospitalization	sociated with that hospitalization. neframe does NOT include this admissions for CHF excluding the current admis	on. The intent of this field is to capture sion.
	timeframe ass Note: This tim hospitalization Coding/Sort	sociated with that hospitalization. neframe does NOT include this admissions for CHF excluding the current admis Selection(Choose one)	on. The intent of this field is to capture sion.
	timeframe ass Note: This tim hospitalization Coding/Sort 0	sociated with that hospitalization. heframe does NOT include this admissions for CHF excluding the current admis Selection(Choose one) Not Hospitalized	on. The intent of this field is to capture sion.
Selections:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2	sociated with that hospitalization. neframe does NOT include this admissions for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months	on. The intent of this field is to capture sion.
Selections:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class -	sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months	on. The intent of this field is to capture sion. Explanation
Selections: Field Name:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class - NYHAclass	sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months	on. The intent of this field is to capture sion. Explanation Seq No: 3100
Selections: Field Name: Short Name: Status:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class - NYHAclass	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes
Selections: Field Name: Short Name: Status:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class NYHA Class NYHAclass New Text (Categor	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes
Selections: Field Name: Short Name: Status: Format:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class NYHA Class NYHAclass New Text (Categor	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class NYHA Class NYHAclass New Text (Categor	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: arent Element:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class - NYHA Class New Text (Categor Client	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: arent Element: Parent Value:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class - NYHA Class New Text (Categor Client	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: arent Element: Parent Value: Missing Data:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class - NYHA Class New Text (Categor Client	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes
Selections: Field Name: Short Name: Status: Format: Data Source: arent Element: Parent Value: Missing Data: Valid Range: Usual Range:	timeframe ass Note: This tim hospitalization Coding/Sort 0 1 2 NYHA Class - NYHA Class - NYHA class New Text (Categor Client Report	Sociated with that hospitalization. The frame does NOT include this admission for CHF excluding the current admis Selection(Choose one) Not Hospitalized Yes-Within 6 months Yes-Greater than 6 months - Current Status	on. The intent of this field is to capture sion. Explanation Seq No: 3100 Core: Yes

Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Coding/Sort	Selection(Choose one)	Explanation
-		· · · · · · · · · · · · · · · · · · ·
1	Class	Patient has cardiac disease but without resulting limitations of ordinary physical activity. Ordinary physical activity (e.g., walking several blocks or climbing stairs) does not cause undue fatigue, palpitation, dyspnea, or anginal pain. Limiting symptoms may occur with marked exertion.
2	Class II	Patient has cardiac disease resulting in slight limitation of ordinary physical activity. Patient is comfortable at rest. Ordinary physical activity such as walking more than two blocks or climbing more than one flight of stairs results in limiting symptoms (e.g., fatigue, palpitation, dyspnea, or anginal pain).
3	Class III	Patient has cardiac disease resulting in marked limitation of physical activity. Patient is comfortable at rest. Less than ordinary physical activity (e.g., walking one to two level blocks or climbing one flight of stairs) causes fatigue, palpitation, dyspnea, or anginal pain.
4	Class IV	Patient has dyspnea at rest that increases with any physical activity. Patient has cardiac disease resulting in inability to perform any physical activity without discomfort. Symptoms may be present even at rest. If any physical activity is undertaken, discomfort is increased.
Cardiac Arres	t	Seq No: 3110
Arrest		Core: Yes
		arvested: Yes
	ical)	
Client		
Poport		
Report		
Report		
·	t	
Cardiac Arres	t patient experienced cardiac arrest due to arr	hythmia
	3 4 Cardiac Arres Arrest New	Country soft Selection (choose one) 1 Class I 2 Class II 3 Class III 4 Class IV Cardiac Arrest Arrest New H Text (Categorical)

Note: Patient history is defined as any time prior to the date of implant.

Implantable Cardioverter Defibrillators Registry v1.08

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No Arrest	
	1	Brady Arrest	
	2	Tachy Arrest	
Field Name:	Brady Arrest I	Reason	Seq No: 3111
Short Name:	-		Core: Yes
Status:	•	н	arvested: Yes
Format:	Text (Categor		
Data Source:	· •	,	
Parent Element:	Cardiac Arres	t(3110)	
Parent Value:	Brady Arrest		
Missing Data:	Report		
Valid Range:			
Usual Range:			
Description:	Brady Arrest I	Reason	
Definition:	Indicate the re	eason(s) for the Brady Arrest.	
Selections:	Coding/Sort	Selection(Choose multiple)	Explanation
	1	Acute MI	
	2	Severe Electrolyte Disturbance	
	3	Drug Induced Arrhythmia	
	4	Sinus Node Dysfunction/AV Block	
	5	Unknown Etiology	
Field Name:	Tachy Arrest	Reason	Seq No: 3112
Short Name:	TachyArrest		Core: Yes
Status:	New	H	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:	Cardiac Arres	t(3110)	
Parent Value:	Tachy Arrest		
Missing Data:	Report		
Valid Range:			
Usual Range:			
	Tachy Arrest		
	1 12 4 41	eason(s) for the Tachy Arrest.	

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Selections:	Coding/Sort	Selection(Choose multiple)	Explanation
	1	Acute MI	
	2	Severe Electrolyte Disturbance	
	3	Drug Induced Arrhythmia	
	4	Primary VT/VF	
	5	Unknown Etiology	
Field Name:	Atrial Fibrillati	on/Atrial Flu	Seq No: 3120
Short Name:	Flutter		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:			
Usual Range:			
Description:	Atrial Fibrillati	on/Atrial Flutter	
Definition:	Indicate if the	patient has a documented history of atrial fib	rillation or flutter.
	Note: Patient	history is defined as any time prior to the date	e of implant.
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
Field Name:	Ventricular Ta	chycardia	Seq No: 3130
Short Name:	VT		Core: Yes
Status:	New	Ha	arvested: Yes
-	_ /-		
Format:	Text (Categor	ical)	
Format: Data Source:		ical)	
		ical)	
Data Source:		ical)	
Data Source: Parent Element:	Client	ical)	
Data Source: Parent Element: Parent Value:	Client	ical)	
Data Source: Parent Element: Parent Value: Missing Data:	Client	ical)	
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	Client		

Note: Patient history is defined as any time prior to the date of implant.

Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	No history of spontaneous ventricular tachycardia.
	1	Yes-VT, Non-Sustained	Three or more consecutive beats of ventricular origin, terminating spontaneously in less than 30 seconds.
	2	Yes-Monomorphic Sustained VT	VT greater than 30 seconds in duration and/or requiring termination due to hemodynamic compromise in less than 30 seconds.
	3	Yes-Polymorphic Sustained VT	VT with a constantly changing morphology lasting greater than 30 seconds in duration and/or requiring termination due to hemodynamic compromise in less than 30 seconds.

Seq No: 3140

Harvested: Yes

Core: Yes

Field Name: Sinus Node Function

Short Name: SinusNodeFn

Status: New

Format: Text (Categorical)

Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

Valid Range:

Usual Range:

Description: Sinus Node Function

Definition: Indicate if the patient's sinus node function was normal or abnormal.

Note: Timeframe includes any time prior to the date of implant.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	Normal	
	2	Abnormal	Abnormal - History of any pause longer than 3 seconds OR Tachy/Brady Syndrome.
Field Name:	Cardiac Trans	splant	Seq No: 3150
Short Name:	XplantPrev		Core: Yes
Status:	New	H	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:			

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Usual Range:

Description: Cardiac Transplant

Definition: Indicate if the patient had a history of cardiac transplant surgery.

Note: Patient history is defined as any time prior to the date of implant

	Note: Patient history is defined as any time prior to the date of implant.		
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
Field Name:	Non-Ischemic	Dilated Cardiomyopathy	Seq No: 3160
Short Name:	NIDilatedCard	ИМуо	Core: Yes
Status:	New	Н	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:			
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:			
Usual Range:			
Description:	Non-Ischemic	Dilated Cardiomyopathy	
Definition:		patient has a history of non-ischemic dilated	cardiomyopathy documented by heart
	failure and reduced systolic function.		
	Note: Patient history is defined as any time prior to the date of implant.		
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes-Within the past 3 months	
	2	Yes-3 to 9 months	
	3	Yes-Greater than 9 months	
Field Name:	Ischemic Hea	rt Disease	Seq No: 3180
Short Name:	IschemicHD		Core: Yes
Status:	New	н	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Element:			

Parent Value: Missing Data: Report

Valid Range:

Usual Range:

Description: Ischemic Heart Disease

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Definition: Indicate if the patient shows evidence of ischemic heart disease as documented by any of the following conditions:

-At least one major epicardial artery with more than 70% obstruction by coronary angiography. -Other Diagnostic Tests: History of myocardial infarction associated with wall motion abnormality as shown by echocardiography or other cardiac imaging. Stress testing diagnostic of coronary artery disease with or without imaging; ECG with evidence of MI; history of chest pain associated with cardiac enzyme abnormality.

Note:

1. Patient history is defined as any time prior to the date of implant.

2. At least one major epicardial artery with more than 70% obstruction by coronary angiography takes precedence over other diagnostic tests if ischemic heart disease has been documented.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes-At Least 1 Epicardial artery greater than 70% Obstruction	At least one epicardial artery greater than 70% obstruction (Angiography).
	2	Yes-Other Diagnostic Tests	History of myocardial infarction associated with wall motion abnormality as shown by echocardiography or other cardiac imaging. Stress testing diagnostic of coronary artery disease with or without imaging; ECG with evidence of MI; history of chest pain associated with cardiac enzyme abnormality.
Field Name:	Previous MI		Seq No: 3190

Core: Yes

Harvested: Yes

Short Name: PrevMITime Status: New

Format: Text (Categorical)

Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

Valid Range:

Usual Range:

Description: Previous Myocardial Infarction and timeframe

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Definition: Indicate if the patient had a MI prior to the device implant.

The patient had at least one documented STEMI or NSTEMI. This can be coded based on physician documentation or history noted in the medical record.

Definitions: NON ST ELEVATION MYOCARDIAL INFARCTION (NSTEMI)

The patient was hospitalized for a myocardial infarction documented in the medical record.

AT LEAST ONE OF THE FOLLOWING BIOCHEMICAL INDICATORS for detecting myocardial necrosis must be present (see below for a definition of Reference Control Limits):

1) Troponin T or I:

a) Maximal concentration of troponin T or I > the MI decision limit on at least one occasion during the first 24 hours after the index clinical event.

2) CK-MB:

a) Maximal value of CK-MB > 2 x the upper limit of normal on one occasion during the first hours after the index clinical event. OR

b) Maximal value of CK-MB, preferable CK-MB mass, > upper limit of normal on two successive samples.

3) Total CK:

a) In the absence of availability of a troponin or CK-MB assay, total CK > 2 x the upper limit of normal, or the B fraction of CK may be employed, but these last two biomarkers are considerably less satisfactory than CK-MB.

AND ONE OF THE FOLLOWING:

1) Either ST segment depression or T wave abnormalities; or

2) Ischemic symptoms in the presence or absence of chest discomfort. Ischemic symptoms may include:

a) unexplained nausea and vomiting; or

b) persistent shortness of breath secondary to left ventricular failure; or

c) unexplained weakness, dizziness, lightheadedness, or syncope.

ST ELEVATION MYOCARDIAL INFARCTION (STEMI)

Indicate whether the patient was hospitalized for an ST Elevation Myocardial Infarction (STEMI) documented in the medical record.

AT LEAST ONE OF THE FOLLOWING BIOCHEMICAL INDICATORS for detecting myocardial necrosis must be present (see below for a definition of Reference Control Limits):

1) Troponin T or I:

a) Maximal concentration of troponin T or I > the MI decision limit on at least one occasion during the first 24 hours after the index clinical event.
2) CK-MB:

a) Maximal value of CK-MB > 2 x the upper limit of normal on one occasion during the first hours after the index clinical event; OR

b) Maximal value of CK-MB, preferable CK-MB mass, > upper limit of normal on two successive samples.

3) Total CK

a) In the absence of availability of a troponin or CK-MB assay, total CK > 2 x the upper limit of normal, or the B fraction of CK may be employed, but these last two biomarkers are considerably less satisfactory than CK-MB.

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Definition: AND ONE OF THE FOLLOWING ECG CHANGES:

1) ST-segment elevation: New or presumed new ST segment elevation at the J point in two or more continguous leads with the cut-off points >=0.2 mV in leads V1, V2, or V3, or >=0.1 mV in other leads; OR

2) Development of any Q wave in leads V1 through V3, or the development of a Q-wave > or = to 30 ms (0.03s) in leads I, II, aVL, aVF, V4, V5, or V6. (Q wave changes must be present in any two continguous leads, and be > or = to 1mm in depth.)

Defining Reference Control Values (MI Diagnostic Limit and Upper Limit of Normal):

Reference values must be determined in each laboratory by studies using specific assays with appropriate quality control, as reported in peer-reviewed journals. Acceptable imprecision (coefficient of variation) at the 99th percentile for each assay should be defined as < or = to 10%. Each individual laboratory should confirm the range of reference values in their specific setting.

Note: If more than one MI occurred, code the most recent event.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes-Within 40 Days of ICD Implant	
	2	Yes-Greater than 40 Days prior to ICD Implant	
	3	Yes-Both Within 40 days/Greater than 40 days	

Field Name: Previous CABG Short Name: PrevCABG Status: New Format: Text (Categorical)

Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

Valid Range:

Usual Range:

Description: Previous Coronary Artery Bypass Graft

Definition: Indicate if the patient had Coronary Bypass Graft Surgery by any approach.

Note: Patient history is defined as any time prior to the current admission. Timeframe does NOT include the current admission. CABGs performed during this admission should be coded within Sequence Number 3590: CABG During this Admission.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
			A N A A A

Field Name: Previous CABG Date Short Name: PrevCABGDate

Seq No: 3210 Core: Yes

Seg No: 3200

Core: Yes Harvested: Yes

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Status:			arvested: Yes	
	Date (mm/dd/	уууу)		
Data Source:		0(0000)		
Parent Element:		G(3200)		
Parent Value:				
Missing Data:	Report			
•	Previous CABG Date > Patient DOB and Previous CABG Date < Admission Date			
Usual Range:				
-		onary Artery Bypass Graft Date		
Definition:	Indicate the d	ate of the most recent CABG. If month and/c	or day are not known enter 01.	
	Note: In the ca	ase of multiple CABGs prior to this admission	, indicate the most recent.	
Selections:				
Field Name:	Previous PCI		Seq No: 3220	
Short Name:	PrevPCI		Core: Yes	
Status:	New	Ha	arvested: Yes	
Format:	Text (Categor	ical)		
Data Source:	Client			
Parent Element:				
Parent Value:				
Missing Data:	Report			
Valid Range:				
Usual Range:				
Description:	Previous Perc	utaneous Coronary Intervention		
		patient had a previous percutaneous coronal alloon angioplasty, stent or other), performed		
	Note: Timeframe does NOT include the current admission. PCIs performed during this admission should be coded within Sequence Number 3610: PCI During this Admission.			
Selections:	Coding/Sort	Selection(Choose one)	Explanation	
	<u>^</u>	Νο		
	0	INU		
	0 1	Yes-Within the past 3 months		

Field Name: Previous Valvular Surgery Short Name: PrevValveSurg Status: New

Format: Text (Categorical)

Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

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Core: Yes Harvested: Yes

Seq No: 3230

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Valid Range:			
Usual Range:			
-	Previous Valve Surgery		
-	Indicate if the patient had a previous surgical replacement and/or repair of a cardiac valve, by any		
	approach.		
	Note: Patient	history is defined as any time prior to the date	e of implant.
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
Field Name:	Permanent Pa	acemaker	Seq No: 3240
Short Name:	PermPacema	ker	Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	• •	,	
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:	·		
Usual Range:			
-	Permanent Pa	acemaker	
•	Indicate if the	patient had a Pacemaker inserted prior to cu	rrent ICD implant. If yes indicate the type
	of pacemaker		inclution implant. If yes, indicate the type
Selections:	of pacemaker Coding/Sort		Explanation
Selections:	Coding/Sort		
Selections:		Selection(Choose one)	
Selections:	Coding/Sort	Selection(Choose one)	
Selections:	Coding/Sort	Selection(Choose one) No Yes-Atrial Chamber	
Selections:	Coding/Sort 0 1 2	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber	Explanation
	Coding/Sort 0 1 2 3 4	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber	Explanation Both atrial and ventricular chambers.
Field Name:	Coding/Sort 0 1 2 3 4 Previous ICD	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber	Explanation Both atrial and ventricular chambers. Seq No: 3250
Field Name: Short Name:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes
Field Name: Short Name: Status:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250
Field Name: Short Name: Status: Format:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New Text (Categor	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes
Field Name: Short Name: Status: Format: Data Source:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New Text (Categor	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes
Field Name: Short Name: Status: Format: Data Source: Parent Element:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New Text (Categor	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes
Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New Text (Categor Client	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes
Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New Text (Categor Client	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes
Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New Text (Categor Client	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes
Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	Coding/Sort 0 1 2 3 4 Previous ICD PrevICD New Text (Categor Client	Selection(Choose one) No Yes-Atrial Chamber Yes-Ventricular Chamber Yes-Dual Chamber Yes-Biventricular Ha	Explanation Both atrial and ventricular chambers. Seq No: 3250 Core: Yes

Implantable Cardioverter Defibrillators Registry v1.08

Definition:	Indicate if the patient had an ICD Implant procedure prior to this admission.			
		me does NOT include the current admission. : ICD Explant Device ID.	This device is coded within Sequence	
Selections:	Coding/Sort	Selection(Choose one)	Explanation	
	0	No		
	1	Yes-Single Chamber		
	2	Yes-Dual Chamber		
	3	Yes-Biventricular		
Field Name:	Previous ICD	Date	Seq No: 3260	
Short Name:	PrevICDDate		Core: Yes	
Status:	New	Ha	arvested: Yes	
Format:	Date (mm/dd/	уууу)		
Data Source:	Client			
Parent Element:	Previous ICD((3250)		
Parent Value:	Yes-Single Ch	namber; Yes-Dual Chamber; Yes-Biventricula	ir	
Missing Data:	-			
Valid Range:	Previous ICD Date > Patient DOB and Previous ICD Date < Admission Date			
Usual Range:				
Description:	Previous ICD Date			
Definition:	Indicate the date of the most recent previous ICD implant. If month and/or day are not known, enter 01.			
	Note: In the ca	ase of multiple implants prior to this admissio	n, code the most recent.	
Selections:				
Field Name:	Previous ICD	Reason	Seq No: 3280	
Short Name:	PrevICDReas	on	Core: Yes	
Status:	New	Ha	arvested: Yes	
Format:	Text (Categor	ical)		
Data Source:	Client			
Parent Element:	Previous ICD((3250)		
Parent Value:	Yes-Single Ch	namber; Yes-Dual Chamber; Yes-Biventricula	ar	
Missing Data:	Report			
Valid Range:				
Usual Range:				
Description:	Previous ICD	Reason		
Definition:	Indicate the p	Indicate the previous ICD indication (reason for ICD implant).		

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Implantable Cardioverter Defibrillators Registry v1.08

Selections:	Coding/Sort	Selection(Choose multiple)	Explanation
	1	Primary Prevention	Primary Prevention was the original indication for patients who were at risk for sudden death but had not suffered from a spontaneous life-threatening ventricular arrhythmia, syncope, or sudden cardiac death.
	2	Syncope with Inducible VT	Syncope: Sudden loss of consciousness with loss of postural tone, not related to anesthesia. Inducible VT refers to performance of electrophysiological testing with resulting induction of VT.
	3	Spontaneous Monomorphic Sustained VT	VT with a constant morphology greater than 30 seconds in duration and/or requiring termination due to hemodynamic compromise in less than 30 seconds.
	4	Spontaneous Polymorphic Sustained VT	VT with a constantly changing morphology lasting greater than 30 seconds in duration and/or requiring termination due to hemodynamic compromise in less than 30 seconds.
	5	Ventricular Fibrillation	Rapid, usually more than 300 bpm (cycle length 180 msec or less), grossly irregular ventricular rhythm with marked variability in cycle length, lack of discernible discreet QRS complex.
	6	Cardiac Arrest/Arrhythmia - Etiology Unknown	Sudden loss of consciousness requiring cardioversion or defibrillation to restore hemodynamic stability.
	7	Syncope and High Risk Characteristics	Syncope: Sudden loss of consciousness with loss of postural tone, not related to anesthesia, and High Risk Characteristics specific for non-ischemic dilated cardiomyopathy, or ischemic heart disease with significant ventricular dysfunction, hypertrophic cardiomyopathy, Brugada Syndrome, Long QT Syndrome.

Implantable Cardioverter Defibrillators Registry v1.08

Selections:	8	AFib	A cardiac arrythmia arising from the atrium with an atrial rate greater than 300 bpm and an irregular ventricular response in the presence of conduction. AF can be further characterized as: First detected AF Paroxysmal AF: AF is self-terminating within 7 days of recognized onset. Persistent AF: AF is not self-terminating within 7 days, or is terminated electrically or pharmacologically. Chronic AF: AF lasting more than 6 months.
Field Name:	Previous ICD	Implant Site	Seq No: 3290
Short Name:	PrevICDsite		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:	Previous ICD	(3250)	
Parent Value:	Yes-Single Ch	namber; Yes-Dual Chamber; Yes-Biventricula	ır
Missing Data:	Report		
Valid Range:			
Usual Range:			
•	Previous ICD	Implant Site	
Description: Definition:	Indicate the p	Implant Site revious ICD implant site.	
Description:	Indicate the p	•	Explanation
Description: Definition:	Indicate the p	revious ICD implant site.	Explanation
Description: Definition:	Indicate the p Coding/Sort	revious ICD implant site. Selection(Choose one)	Explanation
Description: Definition: Selections:	Indicate the p Coding/Sort	revious ICD implant site. Selection(Choose one) Pectoral Abdominal	Explanation Seq No: 3310
Description: Definition: Selections:	Indicate the p Coding/Sort 1 2 Cerebrovascu	revious ICD implant site. Selection(Choose one) Pectoral Abdominal	
Description: Definition: Selections: Field Name:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease	revious ICD implant site. Selection(Choose one) Pectoral Abdominal lar Disease	Seq No: 3310
Description: Definition: Selections: Field Name: Short Name: Status:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease	Seq No: 3310 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease New Text (Categor	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease	Seq No: 3310 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease New Text (Categor	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease	Seq No: 3310 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease New Text (Categor	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease	Seq No: 3310 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease New Text (Categor Client	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease	Seq No: 3310 Core: Yes
Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease New Text (Categor Client	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease	Seq No: 3310 Core: Yes
Description: Definition: Selections: Selections: Short Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease New Text (Categor Client Report	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease Haical)	Seq No: 3310 Core: Yes
Description: Definition: Selections: Selections: Short Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	Indicate the p Coding/Sort 1 2 Cerebrovascu CVDisease New Text (Categor Client	revious ICD implant site. Selection(Choose one) Pectoral Abdominal Ilar Disease Haical)	Seq No: 3310 Core: Yes

Implantable Cardioverter Defibrillators Registry v1.08

Definition:	 Indicate if the patient had cerebrovascular disease (CVD) prior to device implant, defined as any one of the following: Cerebrovascular Accident (CVA): Patient has a history of stroke, i.e., loss of neurological function with residual symptoms at least 72 hours after onset. Reversible Ischemic Neurologic Deficit (RIND): Patient has a history of loss of neurological function with symptoms at least 24 hours after onset but with complete return of function within 72 hours. Transient Ischemic Attack (TIA): Patient has a history of loss of neurological function that was abrupt in onset but with complete return of function within 24 hours. Non-invasive/invasive carotid test with greater than 75% occlusion. Previous carotid artery surgery. This does not include neurological disease processes such as metabolic and/or anoxic ischemic encephalopathy. 			
Selections:	Coding/Sort	· · · · ·	Explanation	
	0	No		
	1	Yes		
Field Name:	Chronic Lung	Disease	Seq No: 3320	
Short Name:	LungDisease		Core: Yes	
Status:	New	Ha	arvested: Yes	
Format:	Text (Categor	ical)		
Data Source:	Client			
Parent Element:				
Parent Value:				
Missing Data:	Report			
Valid Range:				
Usual Range:				
Description:	Chronic Lung	Disease		
Definition:	chronic obstru	Indicate if the patient has a documented history of chronic lung disease prior to this admission (i.e. chronic obstructive pulmonary disease, emphysema, asthma, chronic bronchitis), or has been or is currently being treated with pharmocologic therapy.		
Selections:	Coding/Sort	Selection(Choose one)	Explanation	
	0	No		
	1	Yes		
Field Name:	Diabetes		Seq No: 3330	
Short Name:	Diabetes		Core: Yes	
Status:	New	Ha	arvested: Yes	
Format:	Text (Categor	ical)		
Data Source:	Client			
Parent Element:				
Parent Value:				
Missing Data:	Report			
Valid Range:				
Usual Range:				

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Description:		patient has a history of diabetes, regardless	of duration of disease or need for		
Demition.		gents. It does not include gestational diabetes			
	Note: Patient	Note: Patient history is defined as any time prior to the date of implant.			
Selections:	Coding/Sort	Selection(Choose one)	Explanation		
	0	No			
	1	Yes			
Field Name:	Hypertension		Seq No: 3340		
Short Name:	Hypertension		Core: Yes		
Status:	New	Ha	arvested: Yes		
Format:	Text (Categor	ical)			
Data Source:	Client				
Parent Element:					
Parent Value:					
Missing Data:	Report				
Valid Range:					
Usual Range:					
Description:	Hypertension				
Definition:		patient has a history of hypertension defined			
		hypertension diagnosed and treated with medi			
	 Blood pressure greater than 140 systolic or 90 diastolic on at least 2 occasions. Currently on antihypertensive pharmacologic therapy. 				
Colootionos	Note: Patient history is defined as any time prior to the date of implant.				
Selections:		······································	e of implant.		
	Coding/Sort		Explanation		
	Coding/Sort				
	Coding/Sort	Selection(Choose one)			
Field Name:	0	Selection(Choose one) No Yes			
Field Name: Short Name:	0 1 Renal Failure	Selection(Choose one) No Yes	Explanation		
	0 1 Renal Failure- Dialysis	Selection(Choose one) No Yes Dialysis	Explanation Seq No: 3350		
Short Name: Status:	0 1 Renal Failure- Dialysis	Selection(Choose one) No Yes -Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status:	0 1 Renal Failure- Dialysis New Text (Categor	Selection(Choose one) No Yes -Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status: Format:	0 1 Renal Failure Dialysis New Text (Categor Client	Selection(Choose one) No Yes -Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status: Format: Data Source:	0 1 Renal Failure- Dialysis New Text (Categor Client	Selection(Choose one) No Yes -Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status: Format: Data Source: Parent Element:	0 1 Renal Failure Dialysis New Text (Categor Client	Selection(Choose one) No Yes -Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status: Format: Data Source: Parent Element: Parent Value:	0 1 Renal Failure- Dialysis New Text (Categor Client Report	Selection(Choose one) No Yes -Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data:	0 1 Renal Failure- Dialysis New Text (Categor Client Report	Selection(Choose one) No Yes -Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	0 1 Renal Failure- Dialysis New Text (Categor Client Report	Selection(Choose one) No Yes Dialysis Ha	Explanation Seq No: 3350 Core: Yes		
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	0 1 Renal Failure Dialysis New Text (Categor Client Report Report	Selection(Choose one) No Yes Dialysis Ha	Explanation Seq No: 3350 Core: Yes arvested: Yes		

Note: Patient history is defined as any time prior to the date of implant.

Implantable Cardioverter Defibrillators Registry v1.08 Data Dictionary - Full Specifications

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	

Implantable Cardioverter Defibrillators Registry v1.08

E. Diagnostic Studies						
Field Name:	Ejection Fract	ion Assessed	Seq No: 3360			
Short Name:	EFDone		Core: Yes			
Status:	New		Harvested: Yes			
Format:	Text (Categor	ical)				
Data Source:	Client					
Parent Element:						
Parent Value:						
Missing Data:	Report					
Valid Range:						
Usual Range:						
Description:	Ejection Fract	ion Assessed				
Definition:	Indicate if the patient's ejection fraction was assessed before or during the EP lab visit via invasive (i.e. LV gram) or non-invasive testing (i.e. Echo).					
Selections:	Coding/Sort	Selection(Choose c	one) Explanation			
	0	No				
	1	Yes				
Field Name:	EF %		Seq No: 3370			
Short Name:	EFPercent		Core: Yes			
Status:	New		Harvested: Yes			
Format:	Integer					
Data Source:	Client					
Parent Element:	Ejection Fraction Assessed(3360)					
Parent Value:	· · · ·					
Missing Data:	Report					
Valid Range:						
Usual Range:	10-65					
Description:	Ejection Fraction %					
Definition:	Indicate the lowest Ejection Fraction percent that led to the decision to implant the ICD. The Ejectic Fraction percent is the percentage of blood that has emptied from the ventricle at the end of the contraction.					
Selections:						
Field Name:	EF Timeframe		Seq No: 3380			
Short Name:	EFTimeframe		Core: Yes			
Status:			Harvested: Yes			
Format:	Text (Categorical)					
Data Source:						
Parent Element:	Ejection Fraction Assessed(3360)					
Parent Value:						
		Report				

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Valid Range:

Usual Range:

Description: Ejection Fraction Timeframe

Definition: Indicate the timeframe of the Ejection Fraction percent that led to the decision to implant the ICD.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	0 to 1 month	0 to 1 month (up to 30 days)
	2	1 to 2 months	1 to 2 months (31-60 days)
	3	2 to 3 months	2 to 3 months (61-90 days)
	4	3 to 6 months	3 to 6 months (91-180 days)
	5	6 to 12 months	6 to 12 months (181-365 days)
	6	Greater than 12 months	Greater than 12 months (366 days and greater)

Seq No: 3390

Harvested: Yes

Core: Yes

Field Name: Electrophysiology Study Done

Short Name: EPStudy

Status: New

Format: Text (Categorical)

Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

Valid Range:

Usual Range:

Description: Electrophysiology Study Done

Definition: Indicate if the patient had an EP Study prior to the ICD implant.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	

Field Name:EPS TimeframeSeq No: 3400Short Name:EPSTimeframeCore: YesStatus:NewHarvested: YesFormat:Text (Categorical)Data Source:ClientParent Element:Electrophysiology Study Done(3390)Parent Value:YesMissing Data:ReportValid Range:Usual Range:Description:Electrophysiology Study Timeframe

Definition: Indicate the timeframe of the most recent Electrophysiology study.

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Implantable Cardioverter Defibrillators Registry v1.08

Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) 0 No Arrythmias Induced 1 VT Induced 2 Non-sustained VT 3 Sustained Monomorphic 4 Sustained Polymorphic 5 Ventricular Flutter Induced 6 Ventricular Fibrillation Induced 7 Results Unattainable	Selections:	Coding/Sort	Selection(Choose one)	Explanation
3 2 to 3 months 2 to 3 months 61-90 days) 4 3 to 6 months 3 to 6 months (91-180 days) 5 6 to 12 months 61 to 12 months 6 to 12 months (91-180 days) 6 Greater than 12 months Greater than 12 months Greater than 12 months (91-180 days) 7 Greater than 12 months Greater than 12 months Greater than 12 months (91-180 days) 7 Response Status: Nor Yes Status: Status: <td></td> <td>1</td> <td>0 to 1 month</td> <td>0 to 1 month (up to 30 days)</td>		1	0 to 1 month	0 to 1 month (up to 30 days)
4 3 to 6 months 3 to 6 months (91-180 days) 5 6 to 12 months 6 to 12 months (91-180 days) 6 Greater than 12 months Greater than 12 months (91-180 days) 6 Greater than 12 months Greater than 12 months (91-180 days) 6 Greater than 12 months Greater than 12 months (366 days and greater) Field Name: EPS Findings Seq No: 3410 Short Name: EPS Findings Format: Text (Categorical) Data Surce: Client Parent Value: Yes Pormat: Text (Categorical) Data Surce: Client Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced Induced" is selected, no other selections may be chosen. Select		2	1 to 2 months	1 to 2 months (31-60 days)
5 6 to 12 months 6 to 12 months (181-365 days) 6 Greater than 12 months Greater than 12 months greater) Greater than 12 months Greater than 12 months (366 days and greater) Short Name: EPS Findings Seq No: 3410 Short Name: EPSFindings Core: Yes Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client arent Element: Electrophysiology Study Done(3390) Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced 1 VT Induced 1 1 VT Induced 1 1 VT Induced 1 3 Sustained Monomorphic 1 1 1 1 1 4 Sustained Polymorphic 1<		3	2 to 3 months	2 to 3 months (61-90 days)
6 Greater than 12 months Greater than 12 months (366 days and greater) Field Name: EPS Findings Seq No: 3410 Short Name: EPS Findings Core: Yes Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced		4	3 to 6 months	3 to 6 months (91-180 days)
Field Name: EPS Findings Seq No: 3410 Short Name: EPSFindings Core: Yes Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client Parent Element: Electrophysiology Study Done(3390) Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced		5	6 to 12 months	6 to 12 months (181-365 days)
Short Name EPSFindings Core: Yes Status: New Harvested: Yes Status: Collent Parent Element: Electrophysiology Study Done(3390) Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Usual Range: EPS Findings Detrinition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced		6	Greater than 12 months	· · · ·
Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client Parent Element: Electrophysiology Study Done(3390) Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Selections: Coding/Sort Selection(Choose multiple) 0 No Arrythmias Induced 1 VT Induced 2 Non-sustained VT 3 Sustained Polymorphic 4 Sustained Polymorphic 5 Ventricular Fibrillation Induced 6 Ventricular Fibrillation Induced 7 Results Unattainable Obtained or located. Status Short Name: QRSDuration Status: New Harvested: Yes Format: Integer Data Source: Client Client Parent Value: Value	Field Name:	EPS Findings		Seq No: 3410
Format: Text (Categorical) Data Source: Client Parent Element: Electrophysiology Study Done(3390) Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Usual Range: Description: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced	Short Name:	EPSFindings		Core: Yes
Data Source: Client Parent Element: Electrophysiology Study Done(3390) Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced	Status:	New	H	larvested: Yes
Parent Element: Electrophysiology Study Done(3390) Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Indicate' is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced 1 VT Induced 2 Non-sustained VT 3 Sustained Monomorphic 4 Sustained Polymorphic 5 Ventricular Flutter Induced 6 Ventricular Flutter Induced 7 Results Unattainable 5 Ventricular Florillation Induced 7 Results Unattainable 5 Status: New Harvested: Yes 5 Status: New Yes 5 Status	Format:	Text (Categor	rical)	
Parent Value: Yes Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced 1 VT Induced 2 Non-sustained VT 3 Sustained Monomorphic 4 Sustained Polymorphic 5 Ventricular Fibrillation Induced 7 Results Unattainable The results of the EP Study could not b obtained or located. Field Name: QRS Duration Status: New Harvested: Yes Format Integer Data Source: Client Parent Value: Value	Data Source:	Client		
Missing Data: Report Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced 1 VT Induced 2 Non-sustained VT 3 Sustained Monomorphic 4 Sustained Polymorphic 5 Ventricular Flutter Induced 6 Ventricular Fibrillation Induced 7 Results Unattainable 7 Results Unattainable 5 Seq No: 3420 Short Name: QRS Duration Status: New Harvested: Yes Format: Integer Data Source: Client Parent Value:	Parent Element:	Electrophysio	logy Study Done(3390)	
Valid Range: Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced	Parent Value:	Yes		
Usual Range: Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced	Missing Data:	Report		
Description: EPS Findings Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced 1 1 1 VT Induced 1 1 2 Non-sustained VT 1 1 3 Sustained Monomorphic 1 1 4 Sustained Polymorphic 1 1 5 Ventricular Flutter Induced 1 1 6 Ventricular Fibrillation Induced 1 1 7 Results Unattainable The results of the EP Study could not b obtained or located. Short Name: QRS Duration Seq No: 3420 Short Name: New Harvested: Yes Format: Integer Data Source: Client Parent Value: Venture Parent Value: Venture	Valid Range:	-		
Definition: Indicate the findings associated with the most recent Electrophysiology Study. If "No Arrythmias Induced" is selected, no other selections may be chosen. Selections: Coding/Sort Selection(Choose multiple) Explanation 0 No Arrythmias Induced 1 1 1 VT Induced 1 1 2 Non-sustained VT 1 1 3 Sustained Monomorphic 1 1 4 Sustained Polymorphic 1 1 5 Ventricular Flutter Induced 1 1 6 Ventricular Fibrillation Induced 1 1 7 Results Unattainable The results of the EP Study could not b obtained or located. Short Name: QRS Duration Seq No: 3420 Short Name: QRSDuration Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Value: Value: Value: Value:	Usual Range:			
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2 Non-sustained VT 3 Sustained Monomorphic 4 Sustained Polymorphic 5 Ventricular Flutter Induced 6 Ventricular Fibrillation Induced 7 Results Unattainable The results of the EP Study could not be obtained or located. Steq No: 3420 Short Name: QRS Duration Steq No: 3420 Short Name: QRSDuration Core: Yes Status: New Harvested: Yes Parent Element: Parent Value:	Definition:	Indicate the finduced" is se	ndings associated with the most recent Elec elected, no other selections may be chosen.	
3 Sustained Monomorphic 4 Sustained Polymorphic 5 Ventricular Flutter Induced 6 Ventricular Fibrillation Induced 7 Results Unattainable The results of the EP Study could not be obtained or located. Seq No: 3420 Short Name: QRS Duration Seq No: 3420 Short Name: QRSDuration Status: New Harvested: Yes Data Source: Client Parent Value:	Definition:	Indicate the finduced" is see Coding/Sort	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple)	
4 Sustained Polymorphic 5 Ventricular Flutter Induced 6 Ventricular Fibrillation Induced 7 Results Unattainable 7 Results Unattainable 7 Results Unattainable Status: QRS Duration Status: New Format: Integer Data Source: Client Parent Value: Value:	Definition:	Indicate the finduced" is see Coding/Sort	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced	
5 Ventricular Flutter Induced 6 Ventricular Fibrillation Induced 7 Results Unattainable 7 Results Unattainable The results of the EP Study could not b obtained or located. Field Name: QRS Duration Seq No: 3420 Short Name: QRS Duration Status: New Harvested: Yes Format: Integer Data Source: Client Parent Value:	Definition:	Indicate the finduced" is set Coding/Sort 0 1	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced	
6 Ventricular Fibrillation Induced 7 Results Unattainable 8 Results One and Seq No: 3420 Core: Yes Format: Integer Data Source: Client Parent Element: Parent Value:	Definition:	Indicate the final Induced" is set Coding/Sort 0 1 2	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT	
7 Results Unattainable The results of the EP Study could not be obtained or located. Field Name: QRS Duration Seq No: 3420 Short Name: QRSDuration Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: Parent Value:	Definition:	Indicate the finduced" is set Coding/Sort 0 1 2 3	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic	
Field Name: QRS Duration Seq No: 3420 Short Name: QRSDuration Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: Parent Value:	Definition:	Indicate the final induced is set Coding/Sort 0 1 2 3 4	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic	
Short Name: QRSDurationCore: YesStatus: NewHarvested: YesFormat: IntegerJata Source: ClientParent Element:Parent Value:	Definition:	Indicate the final Induced" is set Coding/Sort 0 1 2 3 4 5	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced	
Short Name: QRSDurationCore: YesStatus: NewHarvested: YesFormat: IntegerJata Source: ClientParent Element:Parent Value:	Definition:	Indicate the final Induced" is set Coding/Sort 0 1 2 3 4 5 6	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced	Explanation Image: second se
Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: Parent Value:	Definition: Selections:	Indicate the final Induced" is set Coding/Sort 0 1 2 3 4 5 6 7	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced Results Unattainable	Explanation
Format: Integer Data Source: Client Parent Element: Parent Value:	Definition: Selections: Field Name:	Indicate the final Induced" is set Coding/Sort 0 1 2 3 4 5 6 7 QRS Duration	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced Results Unattainable	Explanation Explanation The results of the EP Study could not b obtained or located. Seq No: 3420
Data Source: Client Parent Element: Parent Value:	Definition: Selections: Field Name: Short Name:	Indicate the fin Induced" is set Coding/Sort 0 1 2 3 4 5 6 7 QRS Duration QRS Duration	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced Results Unattainable	Explanation Explanation The results of the EP Study could not b obtained or located. Seq No: 3420 Core: Yes
Parent Element: Parent Value:	Definition: Selections: Field Name: Short Name: Status:	Indicate the finduced" is set Coding/Sort 0 1 2 3 4 5 6 7 QRS Duration QRSDuration New	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced Results Unattainable	Explanation Explanation The results of the EP Study could not b obtained or located. Seq No: 3420 Core: Yes
Parent Value:	Definition: Selections: Field Name: Short Name: Status: Format:	Indicate the finduced" is set Coding/Sort 0 1 2 3 4 5 6 7 QRS Duration QRSDuration New Integer	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced Results Unattainable	Explanation Explanation The results of the EP Study could not b obtained or located. Seq No: 3420 Core: Yes
	Definition: Selections: Field Name: Short Name: Status: Format: Data Source:	Indicate the fil Induced" is set Coding/Sort 0 1 2 3 4 5 6 7 QRS Duration QRS Duration New Integer Client	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced Results Unattainable	Explanation Explanation The results of the EP Study could not b obtained or located. Seq No: 3420 Core: Yes
	Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element:	Indicate the finduced" is set Coding/Sort 0 1 2 3 4 5 6 7 QRS Duration QRSDuration New Integer Client	ndings associated with the most recent Elec elected, no other selections may be chosen. Selection(Choose multiple) No Arrythmias Induced VT Induced Non-sustained VT Sustained Monomorphic Sustained Polymorphic Ventricular Flutter Induced Ventricular Fibrillation Induced Results Unattainable	Explanation Explanation The results of the EP Study could not b obtained or located. Seq No: 3420 Core: Yes

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Implantable Cardioverter Defibrillators Registry v1.08

Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: PR Interval Attainable Seq No: 3429 Short Name: PRIntervalAtt Core: Yes Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client Parent Element: Parent Value: Missing Data: Report Valid Range: Usual Range: Usual Range: Usual Range: Coding/Sort Selections: Coding/Sort Selection(Choose one) Explanation Core: Yes Status: New Harvested: Yes Format: Itely Field Name: PR Interval Selections: Field Name: PR Interval Selections: Field Name: PR Interval Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Value Field Name: PR Interval Selection: Field Name: PR Interval Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Value: Yes Missing Data: Report Valid Range: Description: PR Interval Core: Yes Status: New Harvested inte in milliseconds from onset of P wave to onset of QRS complexed in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Seq No: 3440 Short Name: AV Conduction Seq No: 2440 Seq No: Yes Seq	•	QRS Duration Indicate the patient's QRS duration in milliseconds from simultaneous (preferably 3 or more) ECG leads, including I, II, and VI, from the onset to the termination of the QRS.		
Field Name: PR Interval Attainable Seq No: 3429 Short Name: PRIntervalAtt Core: Yes Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client Parent Value: Missing Data: Missing Data: Report Valid Range: Usual Range: Description: PR Interval Attainable Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No	Soloctions		the most recent EKG findings prior to the IC	D implant.
Short Name: PRIntervalAtt Core: Yes Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client Parent Element: Parent Value: Missing Data: Report Valid Range: Usual Range: Description: PR Interval Attainable Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No			toinabla	See Net 2420
Status: New Harvested: Yes Format: Text (Categorical) Data Source: Client Parent Element: Parent Value: Missing Data: Report Valid Range: Usual Range: Usual Range: Description: PR Interval Attainable Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No 1 Yes Field Name: PR Interval Seq No: 3430 Short Name: PR Interval Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: PR Interval Attainable(3429) Parent Value: Yes Missing Data: Report Valid Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Selections:			lamable	•
Format: Text (Categorical) Data Source: Client Parent Element: Parent Value: Missing Data: Report Valid Range: Usual Range: Description: PR Interval Attainable Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No			н	
Data Source: Client Parent Element: Parent Yalue: Missing Data: Report Valid Range: Usual Range: Description: Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) 0 No 1 Yes Field Name: PR Interval Status: No 1 Yes Status: Ne Harvested: Yes Status: Ne Parent Element: PR Interval Attainable(3429) Parent Serption: Yes Missing Data: Report Valid Range: 40-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: <th></th> <th></th> <th></th> <th></th>				
Parent Element: Parent Value: Missing Data: Missing Data: Report Valid Range: Usual Range: Description: Price of the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) 0 No 1 Yes Field Name: PR Interval Seq No: 3430 Short Name: Short Name: PR Interval Seq No: 3430 Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the most recent EKG findings prior to the ICD implant. Selections: Selections: Field Name: AV Conduction Seq No: 3440				
Parent Value: Missing Data: Report Valid Range: Usual Range: Description: PR Interval Attainable Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No				
Missing Data: Report Valid Range: Usual Range: Description: PR Interval Attainable Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No				
Valid Range: Usual Range: Description: PR Interval Attainable Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No				
Usual Range Description PR Interval Attainable Definition: Indicate the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No	•	•		
Description PR Interval Attainable Definition Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No	-			
Definition: Indicate if the patient's PR Interval was attainable. No should be answered when a patient has Afib, a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No	•		tainable	
a greater than 1st Degree Heart block or has a Ventricular Paced rhythm. Selections: Coding/Sort Selection(Choose one) Explanation 0 No	•			uld be answered when a patient has Afib
Country Soft Selection (choose one) Expandion 0 No	Dominion			
1 Yes Field Name: PR Interval Seq No: 3430 Short Name: PRInterval Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: PR Interval Attainable(3429) Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct Core: Yes	Selections:	Coding/Sort	Selection(Choose one)	Explanation
Field Name: PR Interval Seq No: 3430 Short Name: PRInterval Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: PR Interval Attainable(3429) Parent Element: Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Short Name: AV Conduct Core: Yes		0	No	
Short Name: PRInterval Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: PR Interval Attainable(3429) Parent Value: Yes Missing Data: Report Yalid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct Core: Yes		1	Yes	
Short Name: PRInterval Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client Parent Element: PR Interval Attainable(3429) Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Seq No: 3440 Short Name: AVConduction	Field Name:	PR Interval		Seg No: 3430
Format: Integer Data Source: Client Parent Element: PR Interval Attainable(3429) Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct	Short Name:	PRInterval		•
Data Source: Client Parent Element: PR Interval Attainable(3429) Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Seq No: 3440 Short Name: AV Conduct	Status:	New	Ha	arvested: Yes
Parent Element: PR Interval Attainable(3429) Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct	Format:	Integer		
Parent Value: Yes Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct Core: Yes	Data Source:	Client		
Missing Data: Report Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Seq No: 3440 Short Name: AVConduct Core: Yes	Parent Element:	PR Interval At	tainable(3429)	
Valid Range: 10-600 Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct	Parent Value:	Yes		
Usual Range: 40-350 Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct	Missing Data:	Report		
Description: PR Interval Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: Field Name: AV Conduction Short Name: AVConduct	Valid Range:	10-600		
Definition: Indicate the patient's longest measured time in milliseconds from onset of P wave to onset of QRS complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections:	Usual Range:	40-350		
complex in any given ECG lead. Note: Indicate the most recent EKG findings prior to the ICD implant. Selections: Field Name: AV Conduction Short Name: AVConduct Core: Yes	Description:			
Selections: Seq No: 3440 Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct Core: Yes		PR Interval		
Selections: Seq No: 3440 Field Name: AV Conduction Seq No: 3440 Short Name: AVConduct Core: Yes	Definition:	Indicate the page		s from onset of P wave to onset of QRS
Field Name: AV ConductionSeq No: 3440Short Name: AVConductCore: Yes	Definition:	Indicate the pacture complex in an	y given ECG lead.	
Short Name: AVConduct Core: Yes		Indicate the p complex in an Note: Indicate	y given ECG lead.	
	Selections:	Indicate the p complex in an Note: Indicate	y given ECG lead. the most recent EKG findings prior to the IC	D implant.
	Selections: Field Name:	Indicate the p complex in an Note: Indicate	y given ECG lead. the most recent EKG findings prior to the IC	D implant. Seq No: 3440

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Format: Text (Categorical) Data Source: Client Parent Element: **Parent Value:** Missing Data: Report Valid Range:

Usual Range:

Description: AV Conduction

Definition: Indicate the patient's Atrioventricular Conduction rhythm.

Note: Indicate the EKG findings leading to the decision to implant the ICD.

Selections: Coding/Sort Selection(Choose one) Explanation 1 Normal 2 Abnormal-1st Degree Heart Block Only 3 Abnormal-Heart Block 2nd or 3rd Degree (not paced) 4 Paced(any) Seq No: 3450 Core: Yes

Harvested: Yes

Field Name: Intraventricular Conduction

Short Name: IVConduct

Status: New

Format: Text (Categorical)

Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

Valid Range:

Usual Range:

Description: Intraventricular Conduction

Definition: Indicate the patient's Intraventricular Conduction.

Note: Indicate the EKG findings leading to the decision to implant the ICD.

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	Normal	
	2	Abnormal-Left Anterior Fascicular Block	
	3	Abnormal-Left Posterior Fascicular Block	
	4	Abnormal-LBBB	
	5	Abnormal-RBBB	
	6	Abnormal-Intraventricular Conduction Delay, Nonspecific	
	7	Paced	
	8	Abnormal-Bifascicular Block (RBBB Plus LAF)	
	9	Abnormal-Bifascicular Block (RBBB Plus LPF)	
Field Name:	Creatinine Le	vel	Seq No: 3460
Short Name:	Creatinine		Core: Yes

Harvested: Yes Status: New Format: Decimal (2,1) eg. 99.9 Data Source: Client Parent Element: Parent Value: Missing Data: Report Valid Range: 0.1-30 Usual Range: 0.1-9 **Description:** Creatinine Level Definition: Indicate the patient's most recent preoperative Creatinine level prior to the ICD implant. The creatinine level is measured in mg/dL. Selections: Field Name: BUN Level Seq No: 3470 Short Name: BUNLevel Core: Yes Status: New Harvested: Yes Format: Integer Data Source: Client **Parent Element:** Parent Value: Missing Data: Report Valid Range: 1-150 Usual Range: 1-80

Description: Blood Urea Nitrogen Level

Definition: Indicate the patient's most recent preoperative BUN (Blood Urea Nitrogen) level prior to the ICD implant. The BUN level is measured in mg/dL.

Selections:

Implantable Cardioverter Defibrillators Registry v1.08

Field Name:	Sodium Level		Seq No: 3480
Short Name:	NaLevel		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Integer		
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:	80-160		
Usual Range:	110-150		
Description:	Sodium Level		
Definition:	Indicate the pa level is measu	atient's most recent preoperative Sodium leve ured in mEq/L.	el prior to the ICD implant. The Sodium
Selections:			
Field Name:	BNP Drawn		Seq No: 3485
Short Name:	BNPDrawn		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Text (Categor	ical)	
Data Source:	Client		
Parent Element:			
Parent Value:			
Missing Data:	Report		
Valid Range:			
Usual Range:			
Description:	BNP Drawn		
Definition:	Indicate if the implant.	patient had a preoperative BNP (B-type Natr	iuretic Peptide) drawn prior to the ICD
Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
Field Name:	BNP Level		Seq No: 3490
Short Name:	BNPLevel		Core: Yes
Status:	New	Ha	arvested: Yes
Format:	Integer		
Data Source:	Client		
Parent Element:	BNP Drawn(3	485)	
Parent Value:	Yes		
Missing Data:	Report		
Valid Range:	1-5000		
Usual Range:	1-2000		
Description:	BNP Level		

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Implantable Cardioverter Defibrillators Registry v1.08

Definition:	Indicate the patient's most recent preoperat ICD implant. The BNP is measured in pg/m	ive BNP (B-type Natriuretic Peptide) prior to the IL.
Selections:		
Field Name:	Systolic BP	Seq No: 3500
Short Name:	SystolicBP	Core: Yes
Status:	New	Harvested: Yes
Format:	Integer	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	Report	
Valid Range:	60-250	
Usual Range:	70-200	
Description:	Systolic Blood Pressure	
Definition:	Indicate the patient's systolic blood pressure in mm-Hg.	e on day of implant prior to sedation. Measured
Selections:		

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Implantable Cardioverter Defibrillators Registry v1.08 Data Dictionary - Full Specifications

F. ICD Procedure(s)

Seq No: 3505

Harvested: Yes

Core: Yes

Field Name: ICD Indication

Short Name: ICDIndication

Status: New

Format: Text (Categorical)

Data Source: Client

Parent Element:

Parent Value:

Missing Data: Illegal

Valid Range:

Usual Range:

Description: ICD Indication

Definition: Indicate the reason for the first ICD implantation in this patient's lifetime.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	1	Primary Prevention	Primary prevention is an indication for patients who are at risk for sudden death but have not yet suffered from a spontaneous life-threatening ventricular arrhythmia, syncope, or sudden cardiac death. (This includes patients who have never experienced syncope or cardiac arrest but have inducible ventricular tachycardia during electrophysiologic testing for risk stratification.)
	2		Secondary prevention is an indication for patients who have already experienced a spontaneous life-threatening ventricular arrhythmia, a cardiac arrest, or unexplained syncope with workup suggesting a high probability that a ventricular tachyarrythmia was the cause of the syncope.

Field Name:Reason(s) for Re-implantationSeq No: 3506Short Name:ReImpReasonCore: YesStatus:NewHarvested: YesFormat:Text (Categorical)Harvested: YesData Source:ClientParent Element:Previous ICD(3250)Parent Value:Yes-Single Chamber; Yes-Dual Chamber; Yes-BiventricularMissing Data:Missing Data:ReportValid Range:Usual Range:Description:Reason(s) for Re-implantationKenter

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Definition: Indicate the reason(s) why device was re-implanted.

Notor	Applicable only	if Soguopo	Number 2250	Draviaua ICD in	"Voo"
note.	Applicable offi	/ II Sequence		: Previous ICD, is	5 165.

Selections:	Coding/Sort	Selection(Choose multiple)	Explanation
	1	End of Battery Life	The manufacturer's designation that the pulse generator battery has reached the end of its service life.
	2	Device Upgrage	Replacement of a pulse generator with a model with additional pacing capabilities such as an upgrade from a single to a dual chamber device, or the replacement of a non-CRT device with a CRT device.
	3	Device Infection	Replacement of a device because of an infection involving a previously implanted device
	4	Device Malfunction	'HMEH SHURUP DOFH RXWIGH PDQXIDFWUHUV GHUJQDWG VSHFILLFDWRQ WIDWFDQQRWEH UHVROHGZLWU UHSURJUDP PLQJ QHFHVVLWWQJLQWUH UHSODFHP HQWRI WUH GHMEH LQWUH SK\VLFLDQNV RSLQLRQ
	5	Device Under Manufacturer Advisory/Recalled	A device model recognized by the manufacturer as demonstrating a recurring performance failure resulting in an advisory letter to physicians. This may or may not reach the level of an FDA designated recall.

Field Name: Mult ICDs implanted during admit Short Name: MultipleICDs Status: New Format: Text (Categorical) Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

Valid Range:

Usual Range:

Description: Multiple ICDs implanted during this admission

Definition: Indicate if multiple ICD devices were implanted during the current admission.

Note: This field is meant to capture whether an ICD was implanted AND explanted during the current admission. Code "No" if the patient had only one implant during the current admission.

Seq No: 3507

Harvested: Yes

Core: Yes

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Field Name	Reason for Device Repl during admit	Seq No:	3508
Short Name	DevRepReason	Core:	Yes
Status	New	Harvested:	Yes
Format	Text (Categorical)		
Data Source:	Client		
Parent Element	Mult ICDs implanted during admit(3507)		
Parent Value:	Yes		
Missing Data	Report		
Valid Range			
Usual Range			
Description	Reason(s) for device replacement during this admission	on	
	The Rest of the second of the second state is the state of the second state of the sec		•

Definition: Indicate the reason(s) for multiple implants during the current admission.

Note: Applicable only if Sequence Number 3507: Multiple ICDs implated during this admission, is "Yes".

Selections:	Coding/Sort	Selection(Choose multiple)	Explanation
	Coung/Sort	Selection(Choose multiple)	Explanation
	1	Device Upgrade	Replacement of a pulse generator with a model with additional pacing capabilities such as an upgrade from a single to a dual chamber device, or the replacement of a non-CRT device with a CRT device.
	2	Device Infection	Replacement of a device because of an infection involving a previously implanted device
	3	Device Malfunction	'HMEH SHURUP DOFH RXWIGH P DOXIDFWIUHUN/GHMJQDWG VSHFILLFDWIRQ WIDWFDQQRWEH UHVROHGZLWK UHSURJUDP P IQJ QHFHMMUDWIQJIQ WIH UHSODFHP HQWRI WIH GHMEH IQ WIH SK/ VIFIDON/VRSIQIRQ
	4	Device Under Manufacturer Advisory/Recalled	A device model recognized by the manufacturer as demonstrating a recurring performance failure resulting in an advisory letter to physicians. This may or may not reach the level of an FDA designated recall.
Field Name:	Implant Opera	ator UPIN	Seq No: 3510
Short Name:	DrUpin		Core: Yes

Harvested: Yes

Field Name: Implant Operator UPIN Short Name: DrUpin Status: New

Format: Text (6) Data Source: Client

Parent Element:

Parent Value:

Missing Data: Report

Valid Range:

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Data Dictionary - Full Specifications

Usual Range:		
Description:	Implant Operator's Unique Physician Ic	entification Number
Definition:	assigned by CMS, are used to uniquely may contain any letter or number chara	ique (P)hysician (I)dentification (N)umber. UPINs, identify physicians for Medicare billing purposes and cter combination. The UPIN should be specified for the physician placing the leads. Implanting physician n.
Selections:		
Field Name:	Implant Operator NPI	Seq No: 3515
Short Name:	DrNPI	Core: Yes
Status:	New	Harvested: Yes
Format:	Text (10)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	Report	
Valid Range:		
Usual Range:		
•	Implant Operator's National Provider Id	
Definition:		vider (I)dentifier. NPIs, assigned by CMS, are used care billing purposes. The NPI should be specified not the physician placing the leads.
Selections:		
Field Name	Implant Operator First Name	Sea Net 2520
i ioia itailio.	Implant Operator First Name	Seq No: 3520
Short Name:	• •	Core: Yes
	DrGiven	-
Short Name: Status:	DrGiven	Core: Yes
Short Name: Status:	DrGiven New Text (50)	Core: Yes
Short Name: Status: Format:	DrGiven New Text (50)	Core: Yes
Short Name: Status: Format: Data Source:	DrGiven New Text (50)	Core: Yes
Short Name: Status: Format: Data Source: Parent Element:	DrGiven New Text (50) Automatic	Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range:	DrGiven New Text (50) Automatic	Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	DrGiven New Text (50) Automatic Report	Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	DrGiven New Text (50) Automatic Report Implant Operator's First Name	Core: Yes Harvested: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition:	DrGiven New Text (50) Automatic Report	Core: Yes Harvested: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	DrGiven New Text (50) Automatic Report Implant Operator's First Name	Core: Yes Harvested: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Usual Range: Description: Definition: Selections:	DrGiven New Text (50) Automatic Report Implant Operator's First Name Indicate the implant operator's first nam	Core: Yes Harvested: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name:	DrGiven New Text (50) Automatic Report Implant Operator's First Name Indicate the implant operator's first nam Implant Operator Middle Name DrMiddle	e. Seq No: 3525 Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Usual Range: Description: Definition: Selections: Field Name: Short Name:	DrGiven New Text (50) Automatic Report Implant Operator's First Name Indicate the implant operator's first nam Implant Operator Middle Name DrMiddle New	Core: Yes Harvested: Yes le. Seq No: 3525
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status: Format:	DrGiven New Text (50) Automatic Report Implant Operator's First Name Indicate the implant operator's first nam Implant Operator Middle Name DrMiddle New Text (50)	e. Seq No: 3525 Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Usual Range: Usual Range: Selections: Field Name: Short Name: Status: Format: Data Source:	DrGiven New Text (50) Automatic Report Implant Operator's First Name Indicate the implant operator's first nam Implant Operator Middle Name DrMiddle New Text (50)	e. Seq No: 3525 Core: Yes
Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status: Format:	DrGiven New Text (50) Automatic Report Implant Operator's First Name Indicate the implant operator's first nam Implant Operator Middle Name DrMiddle New Text (50)	e. Seq No: 3525 Core: Yes

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-	Implant Opera	tor's Middle Name nplant operator's middle name or middle ir	itial.	
Field Name:	Implant Opera	tor Last Name	Seq No: 3530	
Short Name:	DrSurname		Core: Yes	
Status:	New		Harvested: Yes	
Format:	Text (50)			
Data Source:	Automatic			
Parent Element:				
Parent Value:				
Missing Data:	Report			
Valid Range:				
Usual Range:				
Description:	Implant Opera	tor's Last Name		
Definition:	Indicate the in	plant operator's last name.		
Selections:				
Field Name:	ICD Type		Seq No: 3540	
Short Name:	ICDType		Core: Yes	
Status:	New Harvested: Yes		Harvested: Yes	
Format:	Text (Categor	Text (Categorical)		
Data Source:	Client			
Parent Element:				
Parent Value:				
Missing Data:	Report			
Valid Range:				
Usual Range:				
Description:	ICD Type			
		pe of ICD implanted.		
Selections:	Coding/Sort	Selection(Choose one)	Explanation	
	1	Single Chamber		
	2	Dual Chamber		
	3	Biventricular		
Field Name:	LV Lead Impla	antation Method	Seq No: 3550	
Short Name:	LeadMethod		Core: Yes	
Status:	New		Harvested: Yes	
Format:	Text (Categor	cal)		
Data Source:	Client			

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Parent Element:		10)				
Parent Value:						
Missing Data:	Report					
Valid Range:						
Usual Range:						
-	•	antation Method				
		nethod for implanting the LV lead.				
Selections:	Coding/Sort	Selection(Choose one)	Explanation			
	1	Coronary Sinus				
	2	Epicardial Lead				
	3	Other				
Field Name:	ICD Manufact	urer	Seq No: 3560			
Short Name:	ICDManu		Core: Yes			
Status:	New	Ha	arvested: No			
Format:	Text (100)					
Data Source:	Client					
Parent Element:						
Parent Value:						
Missing Data:	No action					
Valid Range:						
Usual Range:						
Description:	ICD Manufact	urer				
Definition:	Indicate the m	nanufacturer of the implanted or explanted IC	D.			
Selections:						
Field Name:	ICD Model Na	ame	Seq No: 3561			
Short Name:	ICDName		Core: Yes			
Status:	New	H	arvested: No			
Format:	Text (100)					
Data Source:	Client					
Parent Element:						
Parent Value:						
Missing Data:						
Valid Range:						
Usual Range:						
-	ICD Model Na					
		nodel name of the implanted or explanted ICE).			
Selections:						
Field Name:	ICD Model Nu	Imber	Seq No: 3562			
Short Name:	ICDNum		Core: Yes			
Status:	New	Ha	arvested: No			

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Format:	Text (100)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	No action	
Valid Range:		
Usual Range:		
Description:	ICD Model Number	
Definition:	Indicate the model number of the imp	planted or explanted ICD.
Selections:		
Field Name:	ICD Implant Device ID	Seq No: 3565
Short Name:	ICDImpIID	Core: Yes
Status:	New	Harvested: Yes
Format:	Integer (Categorical)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	Report	
Valid Range:		
Usual Range:		
Description:	ICD Implant Device ID	
Definition:	device. The ACC will assign a unique manufacturer, model and model num and added to the data entry tool as a software vendors certified by the ACC master list so that newly approved de	entification number associated with the implanted e identification number for each unique ICD device ber. The list of ICDs will be maintained by the ACC each new device receives FDA approval. Third party C will be required to download and import the ICD evices can be specified and submitted to the ACC. Only ified.Note: In the event of multiple implantations, code the current admission.
Selections:		
Field Name:	ICD Implant Serial Number	Seq No: 3566
Short Name:	ICDImplSerNo	Core: Yes
Status:	New	Harvested: Yes
	Text (100)	
Data Source:	Client	
Parent Element:		
Parent Value:		
Missing Data:	Report	
Valid Range:		
Usual Range:		
-	ICD Implant Serial Number	
	Indicate the ICD Device Serial Numb	er associated with the implanted device.
Selections:		

Implantable Cardioverter Defibrillators Registry v1.08

Field Name:	ICD Explant Device ID	Seq No: 3570		
Short Name:	ICDExpIID	Core: Yes		
Status:	New	Harvested: Yes		
Format:	Integer (Categorical)			
Data Source:	Client			
Parent Element:	Previous ICD(3250)			
Parent Value:	es-Single Chamber; Yes-Dual Chamber; Yes-Biventricular			
Missing Data:	Report			
Valid Range:				
Usual Range:				
Description:	ICD Explant Device ID			
	Indicate the unique ACC assigned identification number associated with the explanted device. The ACC will assign a unique identification number for each unique ICD device manufacturer, model and model number. The list of ICDs will be maintained by the ACC and added to the data entry tool as each new device receives FDA approval. Third party software vendors certified by the ACC will be required to download and import the ICD master list so that newly approved devices can be specified and submitted to the ACC. Only one ICD Explant Device can be specified. Note(1): Applicable only if Sequence Number 3250: Previous ICD, is "Yes". The intent of this field is to record the device in the patient at the time of admission for the current hospital stay. Note(2): This field is NOT to be used in the event of multiple implantations (Sequence Number 3507:Multiple ICDs implanted during this admission, is "Yes").			
Selections:				
	ICD Explant Serial Number	Seq No: 3571		
	ICDExplSerNo	Core: Yes		
Status:		Harvested: Yes		
	Text (100)			
Data Source:				
	Previous ICD(3250)	har: Vac Riventriaular		
Missing Data:	Yes-Single Chamber; Yes-Dual Cham			
Valid Range:	•			
Usual Range:				
-	ICD Explant Serial Number			
-		r associated with the explanted device.		
Selections:				

Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

		G. Adver	se Events				
Field Name:	Available Adv	erse Events		Seq No: 3575			
Short Name:	AvailAdvEver	ıt		Core: Yes			
Status:	New		Ha	arvested: Yes			
Format:	Text (Categor	ical)					
Data Source:	Automatic	Automatic					
Parent Element:							
Parent Value:							
Missing Data:	Report						
Valid Range:							
Usual Range:							
Description:	Available Eve	ents					
Definition:	data collection		ill be used to determine	e ACC website and imported into the IC e if the participant has the right adverse			
Selections:							
Field Name:	Adverse Ever	nts Exist		Seq No: 3580			
Short Name:	AdvEventExis	sts		Core: Yes			
Status:	New		Ha	arvested: Yes			
Format:	Text (Categor	[.] ical)					
Data Source:	Client						
Parent Element:							
Parent Value:							
Missing Data:	Illegal						
Valid Range:							
Usual Range:							
Description:	Adverse Ever	nts Exist					
Definition:	Indicate if the patient had any adverse events during or after the EP lab visit up until discharge. If "Yes" then complete the Adverse Events section.						
Selections:	Coding/Sort	Selection(C	Choose one)	Explanation			
	0	No					
	1	Yes					
Field Name:	Adverse Ever	nt		Seq No: 3581			
Short Name:	AdvEvent			Core: Yes			
Status:	New		Ha	arvested: Yes			
Format:	Text (Catego	rical)					
Data Courses	Client						
Data Source:		$t_{0} = E_{vict}(2580)$					
Parent Element:	Adverse Ever	IIS EXISI(3300)					
		115 EXISt(3560)					

Implantable Cardioverter Defibrillators Registry v1.08

Usual Range: Description: Adverse Event Definition: Indicate the Adverse Event that occurred during or after the EP lab visit up until discharge. The same adverse event can be repeated with a different adverse event date. Note: The initial set of Adverse Events that should be collected are documented in Appendix A of the data dictionary. These Adverse Events may be updated periodically by the ACC. When the Adverse Events have been updated, participants utilizing non-ACC software will be instructed to download and import the latest version of Adverse Event file into their data collection tool. Selections: Field Name: Adverse Event Date Seq No: 3583 Short Name: AdvEventDate Core: Yes	Valid Range:				
Definition: Indicate the Adverse Event that occurred during or after the EP lab visit up until discharge. The same adverse event can be repeated with a different adverse event date. Note: The initial set of Adverse Events that should be collected are documented in Appendix A of the data dictionary. These Adverse Events may be updated periodically by the ACC. When the Adverse Events have been updated, participants utilizing non-ACC software will be instructed to download and import the latest version of Adverse Event file into their data collection tool. Selections: Field Name: Adverse Event Date Seq No: 3583 Short Name: AdvEventDate Core: Yes	•				
The same adverse event can be repeated with a different adverse event date. Note: The initial set of Adverse Events that should be collected are documented in Appendix A of the data dictionary. These Adverse Events may be updated periodically by the ACC. When the Adverse Events have been updated, participants utilizing non-ACC software will be instructed to download and import the latest version of Adverse Event file into their data collection tool. Selections: Field Name: Adverse Event Date Seq No: 3583 Short Name: AdvEventDate Core: Yes	•				
A of the data dictionary. These Adverse Events may be updated periodically by the ACC. When the Adverse Events have been updated, participants utilizing non-ACC software will be instructed to download and import the latest version of Adverse Event file into their data collection tool. Selections: Field Name: Adverse Event Date Seq No: 3583 Short Name: AdvEventDate Core: Yes	Demition.				
Field Name: Adverse Event DateSeq No: 3583Short Name: AdvEventDateCore: Yes		A of the data dictionary. These Adverse Events may be updated periodically by the ACC. When the Adverse Events have been updated, participants utilizing non-ACC software will be instructed to download and import the latest version of Adverse Event file into their data			
Short Name: AdvEventDate Core: Yes	Selections:				
	Field Name:	Adverse Event Date	Seq No: 3583		
	Short Name:	AdvEventDate	Core: Yes		
Status: New Harvested: Yes	Status:	New	Harvested: Yes		
Format: Date (mm/dd/yyyy)	Format:	Date (mm/dd/yyyy)			
Data Source: Client	Data Source:	Client			
Parent Element:	Parent Element:				
Parent Value:	Parent Value:				
Missing Data: Report	Missing Data:	Report			
Valid Range: Adverse Event Date >= Date of Implant and Adverse Event Date <= Date of Discharge	Valid Range:	Adverse Event Date >= Date of Imp	lant and Adverse Event Date <= Date of Discharge		
Usual Range:	Usual Range:				
Description: Adverse Event Date	Description:	Adverse Event Date			
Definition: Indicate the date that the Adverse Event occurred.	Definition:	Indicate the date that the Adverse E	Event occurred.		
Selections:	Selections:				

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		H. Discharge			
Field Name:	CABG During	This Admission	Seq No: 3590		
Short Name:	CABGProc		Core: Yes		
Status:	New	H	arvested: Yes		
Format:	Text (Categor	ical)			
Data Source:	Client				
Parent Element:					
Parent Value:					
Missing Data:	Report				
Valid Range:					
Usual Range:					
Description:	CABG During	This Admission			
Definition:	Indicate if the patient had a CABG (Coronary Artery Bypass Graft Surgery) during the current admission.				
		e CABGs are performed during this admissic date of implant.	on, code the date of the CABG performed		
Selections:	Coding/Sort	Selection(Choose one)	Explanation		
	0	No			
	1	Yes			
Field Name:	CABG Date		Seq No: 3600		
Short Name:	CABGDate		Core: Yes		
Status:	New	H	arvested: Yes		
Format:	Date (mm/dd/	уууу)			
Data Source:	Client				
Parent Element:	CABG During	This Admission(3590)			
Parent Value:					
Missing Data:	Report				
Valid Range:	CABG Date >	= Admission Date and CABG Date <= Discha	arge Date		
Usual Range:					
Description:	CABG During	CABG During This Admission - Date			
	Indicate the date Coronary Artery Bypass Graft (CABG) Surgery was performed during the current				
Definition:	Indicate the data admission.	ate Coronary Aftery Bypass Graft (CABG) St	irgery was performed during the current		
Definition: Selections:	admission.	ate Coronary Artery Bypass Graft (CABG) St	irgery was performed during the current		
Selections:	admission.		Seq No: 3610		
Selections:	admission. PCI During Th				
Selections: Field Name:	admission. PCI During Th PCIProc	nis Admission	Seq No: 3610		
Selections: Field Name: Short Name: Status:	admission. PCI During Th PCIProc	his Admission	Seq No: 3610 Core: Yes		
Selections: Field Name: Short Name: Status:	admission. PCI During Th PCIProc New Text (Categor	his Admission	Seq No: 3610 Core: Yes		

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

Parent Value:							
Missing Data:	Report						
Valid Range:							
Usual Range:							
-	PCI During Th						
Definition.	indicate il trie	Indicate if the patient had a PCI during this admission.					
	Note: If multiple PCIs are performed during this admission, code the date of the PCI performed						
Ostastisus	closest to the	closest to the date of implant.					
Selections:	Coding/Sort	Coding/Sort Selection(Choose one) Explanation					
	0	No					
	1	Yes					
Field Name:	PCI Date		Seq No: 3620				
Short Name:	PCIDate		Core: Yes				
Status:	New	Ha	arvested: Yes				
Format:	Date (mm/dd/	уууу)					
Data Source:	Client						
Parent Element:	PCI During Th	nis Admission(3610)					
Parent Value:	Yes						
Missing Data:	Report						
Valid Range:	PCIDate >= A	dmission Date and PCIDate <= Discharge Da	ate				
Usual Range:							
Description:	PCI During This Admission - Date						
Definition:	Indicate the date PCI was performed during the current admission.						
Selections:							
Field Name:	Vital Status Seq No: 3630						
Short Name:	VitalStatus		Core: Yes				
Status:	New	Ha	arvested: Yes				
	· •	Text (Categorical)					
Data Source:	: Client						
Parent Element:							
Parent Value:							
Missing Data:	•						
Valid Range:							
Usual Range:							
Description:		notions ownized during the beautist story. If IN	as "indicate the source of depth				
Selections:		patient expired during the hospital stay. If "Ye					
Selections:	Coding/Sort	Selection(Choose one)	Explanation				
	1	Alive					
	2	Deceased-Cardiac Death					
	-						

3

Deceased-Non-Cardiac Death

Implantable Cardioverter Defibrillators Registry v1.08

Field Name:	Date of Death	This Admit	Seq No: 3640		
Short Name:	DeathDate		Core: Yes		
Status:	New		Harvested: Yes		
Format:	Date (mm/dd/	Date (mm/dd/yyyy)			
Data Source:	Client	Client			
Parent Element:	Vital Status(3	/ital Status(3630)			
Parent Value:	Deceased-Cardiac Death; Deceased-Non-Cardiac Death				
Missing Data:	Report	Report			
Valid Range:	Date of Death	This Admit >=Date of Implant and Date	e of Death This Admit <= Discharge Date		
Usual Range:					
Description:	Date of Death	This Admit			
Definition:	Indicate the d	ate the patient expired during this hosp	talization.		
Selections:					
Field Name:	Death in Lab		Seq No: 3645		
Short Name:	DeathInLab		Core: Yes		
Status:	New		Harvested: Yes		
Format:	Text (Categor	Text (Categorical)			
Data Source:					
	Client				
Data Source: Parent Element:	Client Vital Status(3		eath		
Data Source: Parent Element:	Client Vital Status(3 Deceased-Ca	630)	eath		
Data Source: Parent Element: Parent Value:	Client Vital Status(3 Deceased-Ca	630)	eath		
Data Source: Parent Element: Parent Value: Missing Data:	Client Vital Status(3 Deceased-Ca Report	630)	eath		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	Client Vital Status(3 Deceased-Ca Report	630)	eath		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	Client Vital Status(3 Deceased-Ca Report Death In Lab	630)			
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description:	Client Vital Status(3 Deceased-Ca Report Death In Lab Indicate if the	630) rdiac Death; Deceased-Non-Cardiac D			
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the	630) rdiac Death; Deceased-Non-Cardiac D patient's death occurred in the lab whe	re the device was implanted		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition:	Client Vital Status(3 Deceased-Ca Report Death In Lab Indicate if the Coding/Sort	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one)	re the device was implanted		
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Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes	re the device was implanted Explanation Seq No: 3650		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes arge	re the device was implanted Explanation Seq No: 3650 Core: Yes		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate New Date (mm/dd/	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes arge	re the device was implanted Explanation Seq No: 3650 Core: Yes		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status: Format:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate New Date (mm/dd/	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes arge	re the device was implanted Explanation Seq No: 3650 Core: Yes		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate New Date (mm/dd/	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes arge	re the device was implanted Explanation Seq No: 3650 Core: Yes		
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Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate New Date (mm/dd/ Client	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes arge	re the device was implanted Explanation Seq No: 3650 Core: Yes Harvested: Yes		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate New Date (mm/dd/ Client	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes arge	re the device was implanted Explanation Seq No: 3650 Core: Yes Harvested: Yes		
Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range: Description: Definition: Selections: Field Name: Short Name: Short Name: Status: Format: Data Source: Parent Element: Parent Value: Missing Data: Valid Range: Usual Range:	Client Vital Status(3) Deceased-Ca Report Death In Lab Indicate if the Coding/Sort 0 1 Date of Disch DischDate New Date (mm/dd/ Client	630) rdiac Death; Deceased-Non-Cardiac De patient's death occurred in the lab whe Selection(Choose one) No Yes arge yyyyy) te >= Date of Implant and Discharge Da	re the device was implanted Explanation Seq No: 3650 Core: Yes Harvested: Yes		

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Implantable Cardioverter Defibrillators Registry v1.08 Data Dictionary - Full Specifications

Selections:

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Implantable Cardioverter Defibrillators Registry v1.08

Data Dictionary - Full Specifications

I. Discharge Medications				
Field Name:	Discharge Medication	Seq No: 3660		
Short Name:	Medication	Core: Yes		
Status:	New	Harvested: Yes		
Format:	Text (Categorical)			
Data Source:	Automatic			
Parent Element:	Vital Status(3630)			
Parent Value:	Alive			
Missing Data:	Illegal			
Valid Range:				
Usual Range:				
Description:	Discharge Medication			
Definition:	Indicates the discharge medicatio	٦.		
	•	nedications that should be collected are documented in Apper		

Note: The initial set of discharge medications that should be collected are documented in Appendix B of the data dictionary. These medications may be updated periodically by the ACC. When the discharge medications have been updated, participants utilizing non-ACC software will be instructed to download and import the latest version of medication file into their data collection tool.

Selections:

Field Name: Discharge Medication Prescribed	Seq No: 3665	
Short Name: MedPresc	Core: Yes	
Status: New	Harvested: Yes	
Format: Text (Categorical)		
Data Source: Client		
Parent Element:		
Parent Value:		
Missing Data: Report		
Valid Range:		
Usual Range:		
Description: Discharge Medication Prescribed		
Definition: Indicate if discharge medication was prescrib	ed, not prescribed, contraindicated or blinded.	

Note: "Blinded" should be specified if the patient was in a research study and the prescribing of this specific medication is unknown.

Selections:	Coding/Sort	Selection(Choose one)	Explanation
	0	No	
	1	Yes	
	2	Contraindicated	
	3	Blinded	

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	Adv	erse Events
Field Name:	Cardiac Arrest	Adverse Event Seq No: ae001
Adverse Event ID:	1	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a cardiac arrest as documented by sudden cessation of cardiac activity so that the patient became unresponsive, with no normal breathing and no signs of circulation.	
Field Name:	Drug Reaction	Adverse Event Seq No: ae002
Adverse Event ID:	2	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a drug reaction as documented by anaphylaxis, rash, etc	
Field Name:	Cardiac Perforation	Adverse Event Seq No: ae003
Adverse Event ID:	3	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a cardiac perforation as documented by migration of pacing or defibrillator lead to epicardial surface, resulting in pain, pericardial effusion, failur to capture, capture of diaphragm, phrenic nerve, or intercostals muscle of sufficient magnitude to require repositioning.	
Field Name:	Cardiac Valve Injury	Adverse Event Seq No: ae004
Adverse Event ID:	4	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a cardiac valve injury as documented by manipulation of pacing or defibrillating leads that may tear a valve leaflet or chordae rendinae (usually manifests as a new regurgitant mumur appearing after the procedure).	
Field Name:	Conduction Block	Adverse Event Seq No: ae005
Adverse Event ID:	5	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a conduction block as documented by manipulation of pacing or defibrillating leads that may injure parts of the specialized cardiac conducting system. (Usually manifest as a new RBBB or new noset of complete heart block in a perso with preexisting LBBB).	
Field Name:	Coronary Venous Dissection	Adverse Event Seq No: ae006
Adverse Event ID:	6	

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Definition:	Indicate if the patient experienced a coronary venous dissection as documented by manipulation of pacing or defibrillating leads in the coronary sinus (CS) may result in a tear of the CS endothelium, with dissection into the CS wall. This may occasionally result in perforation of the CS.	
Field Name:	Hematoma	Adverse Event Seq No: ae007
Adverse Event ID:	7	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experience	d hematoma resulting in reoperation or transfusion.
Field Name:	Lead Dislodgement	Adverse Event Seq No: ae008
Adverse Event ID:	8	
Effective Date:	01/01/2004	Expiration Date:
Definition:	: Indicate if the patient experienced a lead dislodgement as documented by movement of lead sufficient to require repositioning.	
Field Name:	Hemothorax	Adverse Event Seq No: ae009
Adverse Event ID:	9	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a hemothorax as documented by accumulation of blood in thorax.	
Field Name:	Pneumothorax	Adverse Event Seq No: ae010
Adverse Event ID:	10	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a pneumothorax as documented by air in thorax sufficient to require chest tube.	
Field Name:	Peripheral Nerve Injury	Adverse Event Seq No: ae011
Adverse Event ID:	11	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced peripheral nerve injury as documented by sensory or motor loss of peripheral nerve function. This may result from external nerve compression as a result of positioning during an implantationprocedure, internal compression (e.g. secondary to hematoma formation) or direct nerve.	
Field Name:	Peripheral Embolus	Adverse Event Seq No: ae012
Adverse Event ID:	12	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a peripheral embolus as documented by acute occlusion of an artery resulting from embolization of a cardiac or proximal arterial thrombus.	
Field Name:	Phlebitis - Superficial	Adverse Event Seq No: ae013

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Adverse Event ID:	13	
Effective Date:	01/01/2004	Expiration Date:
Definition:	: Indicate if the patient experienced superficial phlebitis as documented by signs of superficial venous inflammatio, such as local erythema, tenderness or swelling.	
Field Name:	Phlebitis - Deep	Adverse Event Seq No: ae014
Adverse Event ID:	14	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced deep phlebitis as documented by occlusion of deep vein resulting in extremity swelling, plus or minus signs of inflammation.	
Field Name:	TIA	Adverse Event Seq No: ae015
Adverse Event ID:	15	
Effective Date:	01/01/2004	Expiration Date:
Definition:	: Indicate if the patient experienced a TIA as documented by loss of neurological function tha was abrupt in onset but with complete return of function within 24 hours.	
Field Name:	CVA/Stroke	Adverse Event Seq No: ae016
Adverse Event ID:	16	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced a cerebrovascular accident (CVA) as documented by a central neurological deficit persisting for > 72 hours.	
Field Name:	MI	Adverse Event Seq No: ae017
Adverse Event ID:	17	
Effective Date:	01/01/2004	Expiration Date:

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Implantable Cardioverter Defibrillators Registry v1.08

Definition:	Indicate if the patient experienced an MI during the EP lab visit or after lab visit until discharge (or before any subsequent lab visits) as documented by: Definitions: NON ST ELEVATION MYOCARDIAL INFARCTION (NSTEMI) The patient was hospitalized for a myocardial infarction documented in the medical record. AT LEAST ONE OF THE FOLLOWING BIOCHEMICAL INDICATORS for detecting myocardial necrosis must be present (see below for a definition of Reference Control Limits): 1) Troponin T or I: a) Maximal concentration of troponin T or I > the MI decision limit on at least one occasion
	 during the first 24 hours after the index clinical event. 2) CK-MB: a) Maximal value of CK-MB > 2 x the upper limit of normal on one occasion during the first hours after the index clinical event. OR b) Maximal value of CK-MB, preferable CK-MB mass, > upper limit of normal on two successive samples. 3) Total CK:
	a) In the absence of availability of a troponin or CK-MB assay, total CK > 2 x the upper limit of normal, or the B fraction of CK may be employed, but these last two biomarkers are considerably less satisfactory than CK-MB.
	 AND ONE OF THE FOLLOWING: 1) Either ST segment depression or T wave abnormalities; or 2) Ischemic symptoms in the presence or absence of chest discomfort. Ischemic symptoms may include: a) unexplained nausea and vomiting; or b) persistent shortness of breath secondary to left ventricular failure; or c) unexplained weakness, dizziness, lightheadedness, or syncope.
	ST ELEVATION MYOCARDIAL INFARCTION (STEMI) Indicate whether the patient was hospitalized for an ST Elevation Myocardial Infarction (STEMI) documented in the medical record. AT LEAST ONE OF THE FOLLOWING BIOCHEMICAL INDICATORS for detecting myocardial necrosis must be present (see below for a definition of Reference Control Limits): 1) Troponin T or I: a) Maximal concentration of troponin T or I > the MI decision limit on at least one occasion during the first 24 hours after the index clinical event. 2) CK-MB: a) Maximal value of CK-MB > 2 x the upper limit of normal on one occasion during the first hours after the index clinical event; OR b) Maximal value of CK-MB, preferable CK-MB mass, > upper limit of normal on two successive samples. 3) Total CK
	a) In the absence of availability of a troponin or CK-MB assay, total CK > 2 x the upper limit of normal, or the B fraction of CK may be employed, but these last two biomarkers are considerably less satisfactory than CK-MB.
	 AND ONE OF THE FOLLOWING ECG CHANGES: 1) ST-segment elevation: New or presumed new ST segment elevation at the J point in two or more continguous leads with the cut-off points >=0.2 mV in leads V1, V2, or V3, or >=0.1 mV in other leads; OR 2) Development of any Q wave in leads V1 through V3, or the development of a Q-wave > or = to 30 ms (0.03s) in leads I, II, aVL, aVF, V4, V5, or V6. (Q wave changes must be present in any two continguous leads, and be > or = to 1mm in depth.)

Defining Reference Control Values (MI Diagnostic Limit and Upper Limit of Normal):

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Definition:	: Reference values must be determined in each laboratory by studies using specific assays with appropriate quality control, as reported in peer-reviewed journals. Acceptable imprecision (coefficient of variation) at the 99th percentile for each assay should be defined as < or = to 10%. Each individual laboratory should confirm the range of reference values in their specific setting.	
Field Name:	Pericardial Tamponade	Adverse Event Seq No: ae018
Adverse Event ID:	18	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced fluid in the pericardial space compromising cardiac filling, and requiring intervention as documented by either: 1) Echo showing pericardial fluid and signs of tamponade such as right heart compromise, or 2) Systemic hypotension due to pericardial fluid compromising cardiac function.	
Field Name:	AV Fistula	Adverse Event Seq No: ae019
Adverse Event ID:	19	
Effective Date:	01/01/2004	Expiration Date:
Definition:	Indicate if the patient experienced an AV fistula as documented by a connection between the access artery and the accompanying vein that is demonstrated by arteriography or ultrasound and most often characterized by a continuous bruit.	
Field Name:	Infection Related to Device	Adverse Event Seq No: ae020
Adverse Event ID:	20	
Effective Date:	01/01/2004	Expiration Date:
Definition:	efinition: Indicate if the patient experienced an infection related to the device.	

Appendix B - Medications

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Implantable Cardioverter Defibrillators Registry v1.08

	Medications		
Field Name:	ACE-Inhibitor (any)	Medication Seq No: m001	
Category:	Ace Inhibitor	Medication ID: 1	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Amiodarone	Medication Seq No: m002	
Category:	Antiarrhythmic Agent	Medication ID: 2	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Disopyramide	Medication Seq No: m003	
Category:	Antiarrhythmic Agent	Medication ID: 3	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Dofetilide	Medication Seq No: m004	
Category:	Antiarrhythmic Agent	Medication ID: 4	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Flecainide	Medication Seq No: m005	
Category:	Antiarrhythmic Agent	Medication ID: 5	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Mexiletine	Medication Seq No: m006	
Category:	Antiarrhythmic Agent	Medication ID: 6	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Procainamide	Medication Seq No: m007	
Category:	Antiarrhythmic Agent	Medication ID: 7	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Propafenone	Medication Seq No: m008	
Category:	Antiarrhythmic Agent	Medication ID: 8	
Effective Date:	01/01/2004	Expiration Date:	
Field Name:	Quinidine	Medication Seq No: m009	
Category:	Antiarrhythmic Agent	Medication ID: 9	
Effective Date:	01/01/2004	Expiration Date:	

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Field Name:	Sotalol	Medication Seq No: m010
Category:	Antiarrhythmic Agent	Medication ID: 10
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Other Anti. Arrhy.	Medication Seq No: m011
Category:	Antiarrhythmic Agent	Medication ID: 11
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Hydralazine	Medication Seq No: m012
Category:	Antihypertensive	Medication ID: 12
Effective Date:	01/01/2004	Expiration Date:
Field Name:	ARB (any)	Medication Seq No: m013
Category:	ARB	Medication ID: 13
Effective Date:	01/01/2004	Expiration Date:
Field Name:	ASA	Medication Seq No: m014
Category:	ASA	Medication ID: 14
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Beta-Blocker (any)	Medication Seq No: m015
Category:	Beta Blocker	Medication ID: 15
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Diltiazem	Medication Seq No: m016
Category:	Calcium Channel Blocker	Medication ID: 16
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Verapamil	Medication Seq No: m017
Category:	Calcium Channel Blocker	Medication ID: 17
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Other CCB	Medication Seq No: m018
Category:	Calcium Channel Blocker	Medication ID: 18
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Coumadin	Medication Seq No: m019
Category:	Coumadin	Medication ID: 19

Appendix B - Medications

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Effective Date:	01/01/2004	Expiration Date:
Field Name:	Digoxin	Medication Seq No: m020
Category:	Digoxin	Medication ID: 20
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Diuretic (any)	Medication Seq No: m021
Category:	Diuretic	Medication ID: 21
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Nitroglycerin SL, PRN	Medication Seq No: m022
Category:	Nitrate	Medication ID: 22
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Nitroglycerin Long Acting	Medication Seq No: m023
Category:	Nitrate	Medication ID: 23
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Clopidogrel	Medication Seq No: m024
Category:	Platelet Aggregation Inhibitor	Medication ID: 24
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Ticlopidine	Medication Seq No: m025
Category:	Platelet Aggregation Inhibitor	Medication ID: 25
Effective Date:	01/01/2004	Expiration Date:
Field Name:	Statin (any)	Medication Seq No: m026
Category:	Statin	Medication ID: 26
Effective Date:	01/01/2004	Expiration Date: