

**ADULT CORONARY ARTERY  
BYPASS GRAFT SURGERY IN THE  
COMMONWEALTH OF MASSACHUSETTS  
January 1 – December 31, 2004**

Mass-DAC  
Department of Health Care Policy  
Harvard Medical School  
October 2006

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## MASSACHUSETTS DATA ANALYSIS CENTER (MASS-DAC)

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## MASSACHUSETTS CARDIAC SURGERY CENTERS 2004

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Beth Israel Deaconess Medical Center (BIDMC) 330 Brookline Avenue Boston, MA 02115	Mount Auburn Hospital 330 Mount Auburn Street Cambridge, MA 02138
Boston Medical Center (BMC) 88 East Newton Street Boston, MA 02118	North Shore Medical Center - Salem Hospital 81 Highland Avenue Salem, MA 01970
Brigham & Women's Hospital (B&W) 75 Francis Street Boston, MA 02115	Southcoast Hospital Group - Charlton Memorial Hospital 363 Highland Avenue Fall River, MA 02720
Cape Cod Hospital 27 Park Street Hyannis, MA 02601	Saint Vincent Hospital at Worcester Medical Center 123 Summer Street Worcester, MA 01608
Caritas Saint Elizabeth's Medical Center 736 Cambridge Street Boston, MA 02135	Tufts-New England Medical Center (NEMC) 750 Washington Street Boston, MA 02111
Lahey Clinic 41 Mall Road Burlington, MA 01805	UMass Memorial Medical Center 55 Lake Avenue North Worcester, MA 01655

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## 1 - KEY FINDINGS

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- In 2004, there were **7289** hospital admissions in which at least one cardiac surgery was performed in Massachusetts. More than half (**54.7%**) of the admissions were those in which an isolated coronary artery bypass graft (CABG) surgery was undertaken.
- There were **407 fewer** isolated CABG surgery admissions performed in Massachusetts in 2004 as compared to 2003.
- Since 2002 (when data collection began), there has been a decrease of **617** isolated CABG surgeries in Massachusetts.
- **Fourteen** hospitals performed at least one CABG operation in Massachusetts in 2004.
- In the fourteen hospitals that performed cardiac surgery in 2004, the number of CABG surgery admissions ranged from **101 to 537**.
- The observed mortality rate (defined as the number of patients dying within 30 days of surgery divided by the number of patients undergoing CABG surgery) in Massachusetts during 2004 was **2.01%**.
- **Caritas Saint Elizabeth's Medical Center** was identified as **having higher than expected** 30-day mortality during 2004.

## 2 - INTRODUCTION

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### 2.1 - What is in this Report?

This report describes procedures for calculating hospital-specific risk standardized 30-day mortality rates following isolated coronary artery bypass graft (**CABG**) surgery performed in Massachusetts hospitals in 2004. Surgeries performed in United States Government Hospitals (e.g., VA Boston Healthcare System – Jamaica Plain Campus) are not included in this report. Information pertains to patients who were 18 years of age or older at the time of their surgery.

Not all hospitals in Massachusetts are permitted to perform cardiac surgery. Hospitals wishing to establish a new cardiac surgery program must submit an application to the Determination of Need Program in the Massachusetts Department of Public Health. In 2004, there were fourteen cardiac surgery programs in Massachusetts: eleven established and three relatively new programs. Of the three newer programs, two programs (Southcoast Hospital Group – Charlton Memorial Hospital and Cape Cod Hospital) began performing cardiac surgery in 2002. North Shore Medical Center - Salem Hospital began performing cardiac surgery in 2003. All Massachusetts hospitals with cardiac surgery programs submitted data to Mass-DAC.

This document is the 3rd annual report describing hospital-specific standardized mortality rates following isolated CABG surgery in Massachusetts (all reports are available at [www.massdac.org](http://www.massdac.org)). It describes standardized mortality rates for the fourteen cardiac surgery programs in Massachusetts that performed at least one isolated CABG surgery between January 1, 2004 and December 31, 2004.

## 2.2 - What is Coronary Artery Bypass Graft Surgery?

For a heart to function properly, it needs an oxygen-rich blood supply. Coronary arteries send oxygen-rich blood to the heart. When the coronary arteries are healthy, blood flows easily so that the heart muscle gets the oxygen it needs. Coronary artery disease begins when blood flow to the heart is reduced due to a build-up of plaque. Plaque may build up because of high cholesterol, high blood pressure, smoking, diabetes, genetic predisposition, or other factors. If the plaque build-up increases, the coronary arteries narrow and blood flow to the heart is reduced, often leading to angina (chest pain, arm pain, or jaw tightness that occurs with exertion, or in more serious cases, at rest). If blood flow is completely blocked by a sudden development of a clot within a coronary artery, this usually results in a heart attack or myocardial infarction, referred to as MI, which may irreversibly damage the heart muscle.

Coronary artery disease is usually treated by one of three methods (medication, coronary intervention, or cardiac surgery). The choice of treatment depends on the degree of blockage, patient symptoms, and the number of coronary arteries involved. Coronary artery bypass graft (CABG) surgery is a type of cardiac surgery that creates a new route or bypass around the blocked part of the artery, allowing the blood flow to reach the heart muscle again. During CABG surgery, the blocked coronary arteries are bypassed using some of the patient's own blood vessels. The internal mammary arteries are commonly used for the bypass; however, the saphenous vein in the leg or the radial artery in the arm can also be used. Surgical procedures in which CABG is the only major heart surgery performed are referred to as *isolated CABG* surgery.

### 2.3 - Why Report on CABG Surgery?

CABG surgery accounts for the majority of cardiac surgery performed nationally and is costly. In 2004, isolated CABG surgery accounted for 54.7% of the 7289 cardiac surgery hospital admissions (**Table 2.1**) in Massachusetts. Only patients undergoing isolated CABG surgery were used to determine hospital mortality rates in this report.

<b>Table 2.1: Surgical Procedure Type Classification of Adult Cardiac Surgeries During January 1, 2004 – December 31, 2004, Commonwealth of Massachusetts.</b>		
<b>Surgical Procedure Type</b>	<b>Number of Cardiac Surgery Admissions</b>	<b>% of All Cardiac Surgery Admissions</b>
<b>Isolated CABG</b>	<b>3986</b>	<b>54.7</b>
Mitral Valve Replacement (MVR)	131	1.8
Aortic Valve Replacement (AVR)	599	8.2
MVR + CABG	65	0.9
AVR + CABG	566	7.8
AVR + MVR	36	0.5
Other Cardiac Surgery	1837	25.2
Non-Cardiac (Thoracic) Procedures	69	1.0
<b>All Cardiac Surgery Admissions</b>	<b>7289</b>	<b>100</b>

### 2.4 - Who Receives CABG Surgery in Massachusetts?

Of patients undergoing isolated CABG surgery, 75% were males, more than half (60%) were aged 65 years or older, and the majority (91%) were white (**Table 2.2**). Patients who lived outside of Massachusetts at the time of their surgery comprised 9% of the 3986 isolated CABG surgery admissions.



<b>Table 2.2: Age-Sex-Race distribution for all adult Isolated CABG Surgeries (N = 3986) in MA hospitals during January 1, 2004 - December 31, 2004. Entries represent numbers of patients.</b>										
Age Group	Females					Males				
	White	African American	Hispanic	Other <sup>§</sup>	Total	White	African American	Hispanic	Other	Total
18-44	20	2	1	2	25	51	2	2	6	61
45-54	60	4	2	2	68	345	14	19	12	390
55-64	154	14	15	5	188	785	16	31	32	864
65-74	278	8	21	15	322	880	16	29	37	962
≥ 75	388	9	4	12	413	665	6	10	12	693
<b>Total</b>	<b>900</b>	<b>37</b>	<b>43</b>	<b>36</b>	<b>1016</b>	<b>2726</b>	<b>54</b>	<b>91</b>	<b>99</b>	<b>2970</b>

## 2.5 - Why Report Hospital-Specific Mortality Rates?

Data collected on quality can be used to provide useful information to both patients and health care providers, stimulate additional research, and foster improvements in quality of care. This report uses 30-day mortality as a measure of hospital quality, defined as death occurring within 30 days of the date of the operation, regardless of cause. While it is not the only important endpoint, mortality was selected as the primary measure of hospital quality because it is serious and unambiguous.

## 2.6 - What is Mass-DAC?

Mass-DAC is a data coordinating center responsible to the Massachusetts Department of Public Health for the collection, storage, and analysis of the cardiac data submitted by Massachusetts hospitals. Mass-DAC is located in the Department of Health Care Policy, Harvard Medical School in Boston ([www.massdac.org](http://www.massdac.org)). Mass-DAC is advised

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<sup>§</sup> Includes some missing

by several committees on an ongoing basis: the Massachusetts Cardiac Care Quality Advisory Commission, the Cardiac Advisory Board, the MA STS Outcomes Quality Improvement Committee, and the Cardiac Surgery Data Adjudication Committee. In addition, both the National Society of Thoracic Surgeons and the Massachusetts chapter of the Society of Thoracic Surgeons serve as resources.

## **2.7 - What Data are used in the Report?**

All Massachusetts hospitals that perform cardiac surgery are required by law to submit specific information to Mass-DAC. This report includes data submitted by hospitals in Massachusetts that performed isolated CABG surgery between January 1, 2004 and December 31, 2004. Key variables were rigorously verified by Mass-DAC. This process involved: (1) continuous data quality reports to data managers located at each hospital; (2) discussions with chiefs of every cardiac surgery department Massachusetts; (3) audits of selected chart information by an independent Cardiac Surgery Data Adjudication Committee; (4) a review by an external Cardiac Advisory Board; and (5) linkages and cross-checks with state administrative databases. The same procedures were used to collect data for CABG surgery performed during calendar years 2002 and 2003. All physicians who audited records completed the Harvard Medical School Human Subjects Training and were approved by the Internal Review Board (IRB).

## **2.8 - Risk Adjustment for Assessing Hospital Mortality**

Specific "risk" factors are known to contribute to heart disease. These risk factors include high cholesterol, smoking, high blood pressure, family history of heart disease, diabetes, age, gender, and general health status prior to the CABG operation. Such factors also have an impact on the risk of mortality following surgery. Sicker patients or patients with more health-related risks may be more likely to die following a CABG operation than healthier patients. Moreover, patients who are sicker may be more likely to be treated at particular hospitals while patients who are healthier may be more likely to be treated at other hospitals. To compare hospitals fairly and not to penalize hospitals

that treat sicker patients, it is important to consider differences in patient health prior to surgery.

The statistical process of accounting for differences in patient sickness prior to their surgery is called *risk adjustment*. This statistical process aims to “level the playing field” by considering health risks that patients have prior to surgery. The hospital specific 30-day mortality rates in this report have been adjusted in order to account for differences in patient health prior to surgery.

## 2.9 - How are Hospital Differences in Patient Outcomes Measured?

If there are differences in hospital quality, due to staff, experience, or other factors, then the risks of hospital mortality for two patients having exactly the same risk factors prior to cardiac surgery but who are treated in different hospitals may not be the same. The statistical model used to calculate mortality rates in this report - a *hierarchical logistic regression* model - models the difference between the risks of mortality for patients with the same risk factors who have surgeries in different hospitals. This is accomplished through the inclusion of a hospital-specific (random) effect. If no key risk factor is missing in the statistical model, then the hospital-specific random effect represents quality for each hospital. If there are no differences in the hospital-specific effects across the hospitals, then there is no evidence of quality differences.

### 3 - IDENTIFYING OUTLYING CARDIAC SURGERY PROGRAMS

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The purpose of this report is to identify hospitals that have *unusually* high or *unusually* low mortality rates. Such hospitals are classified as “outlying” – however, the designation of outlying depends on how large the differences are. Two methods were used to identify outlying hospitals. The first method calculates a 95% interval estimate for each hospital’s risk-standardized mortality rate. If the interval estimate does not contain the state unadjusted 30-day hospital mortality rate, the hospital is designated as outlying.

However, because any one hospital could influence the estimates of the risk-standardized mortality rate for other hospitals in the state, Mass-DAC also calculates the predicted number of mortalities at each hospital using the experience of all **other** hospitals in Massachusetts. If there is a low probability that the actual number of mortalities and the predicted number of mortalities is the same, then the hospital is classified as “outlying.”

If the 95% interval estimate for a particular hospital excludes the state unadjusted 30-day hospital mortality rate **or** if the probability of the observed mortality based on mortality predicted from all other hospitals for a particular hospital is small, then Mass-DAC classified the hospital as outlying. It is important to note that the classification in this report is relative to all hospitals in Massachusetts performing CABG surgery.

#### 3.1 - Standardized Mortality Incidence Rates (SMIR)

Mass-DAC calculated a standardized mortality incidence rate (SMIR) and a corresponding 95% “posterior” interval for each hospital. The SMIR is interpreted as the projected mortality rate at the hospital **today** if hospital quality remained the same as in 2004. The SMIR consists of an estimate of the hospital’s underlying (true) risk-adjusted rate divided by an estimate of the mortality rate expected at the hospital given its case-mix. Each hospital’s SMIR should only be interpreted in the context of its posterior interval. If the 95% interval includes the unadjusted state rate, then the hospital mortality is **not different than expected**. If the interval excludes the state unadjusted rate, then the hospital’s SMIR is different from what was expected. In this case, if the upper limit of the interval is lower than the unadjusted state rate, then fewer patients than expected died. Such a hospital

would be categorized as having **lower than expected mortality**. If the lower limit of the interval is higher than the state unadjusted rate, then more patients than expected died. Such a hospital would be categorized as having **higher than expected mortality**.

### 3.2 - Cross-Validated P-Values

Because data from all hospitals are used to estimate the expected number of deaths in any hospital, there is a risk that outlying hospitals may influence the variation between hospitals in risk stratifying mortality rates. One method to identify hospitals as outlying is through “cross-validation.” This process involves systematically dropping each hospital from the data set and re-estimating the risk-adjusted model. Using the new model, the predicted number of deaths at the dropped hospital is calculated. This predicted number may be interpreted as the number of mortalities expected at the dropped hospital if the dropped hospital had the same level of quality as the remaining hospitals in the Commonwealth.

Mass-DAC compared the predicted number to the observed number of deaths at the dropped hospital and calculated a “probability.” This probability, loosely called a “p-value,” quantifies how **likely** the observed number of deaths would be if the dropped hospital had the same level of quality as all remaining cardiac surgery hospitals. Small p-values (those  $\leq 0.01$ ) indicate that the dropped hospital is outlying. When the p-value is small and the actual number of deaths is larger than that predicted by the remaining hospitals, the dropped hospital is classified as having **higher than predicted mortality**; when the p-value is small and the actual number of deaths is smaller than the number of deaths predicted by its peers, then the hospital is classified as having **lower than predicted mortality**. Mass-DAC repeated this procedure, eliminating each cardiac surgery hospital, and calculating a p-value for each hospital.

### 3.3 – Sensitivity Analyses

Several sensitivity analyses were undertaken to determine whether conclusions would change when making reasonable changes to some of the underlying assumptions.

These assumptions related to the amount of between-hospital variation in underlying quality of care. Mass-DAC varied assumptions regarding the amount and type of between-hospital variation in quality of care, and re-estimated the SMIRs as well as the cross-validated p-values.

## 4 - HOSPITAL QUALITY FOLLOWING ISOLATED CABG SURGERY: 2004

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Of the 3986 isolated CABG surgery admissions in 2004 in Massachusetts, 80 patients (2.01%) died within thirty days of the operation. **Table 4.1** lists the prevalence (%) of important risk factors and the relationship of each risk factor (controlling for all other risk factors) with 30-day mortality following surgery. For example, 75% of the 3986 CABG surgery admissions were male patients. Because age is measured in years, the table reports the average number of years over age 65 for the cohort.

Odds ratios greater than 1 correspond to increased risk of mortality while those less than 1 correspond to decreased risk of mortality. The odds ratio of 0.64 for males indicates that males are 0.64 times as likely as females to die within 30 days of a CABG operation. In contrast, patients having cardiogenic shock prior to a CABG operation are 5.53 times more likely to die within 30 days than patients not having cardiogenic shock.

**Figure 4.1** displays the SMIRs and corresponding 95% interval estimates. The solid black vertical line in the figure is the unadjusted state 30-day mortality rate of 2.01% (**Table 4.2** lists the values). Listed on the left-hand side of the figure are the total number of isolated CABG admissions and the expected 30-day mortality rates for each hospital. The expected mortality rate provides an overall assessment of case-mix severity at each hospital. The higher the expected mortality rate at a hospital is, the sicker the hospital's patients. Listed on the right-hand side are the estimated SMIRs. All 95% probability intervals include the unadjusted state rate of 2.01% with the exception of one (Saint Elizabeth's Medical Center).

**Figure 4.2** presents the cross-validated p-values under a number of different assumptions regarding between-hospital variability in quality. The p-value for Saint Elizabeth's Medical Center is 0.0054. The cross-validation analysis indicated that the actual number of mortalities was statistically higher than the predicted number of mortalities at Saint Elizabeth's Medical Center. This evidence suggests that, relative to all other cardiac surgery programs in Massachusetts, mortality following isolated CABG surgery at Saint Elizabeth's Medical Center is higher than expected.

In summary, the 2004 isolated CABG mortality data indicate one hospital is a statistical outlier, having higher than expected mortality. The 95% SMIR interval for Saint

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Elizabeth's Medical Center lies above the state rate of 2.01% (95% Interval: 2.03%, 7.47%). The cross-validated p-value indicates that Saint Elizabeth's Medical Center had statistically higher than predicted mortality relative to all other Massachusetts cardiac surgery hospitals. These results remained unchanged relative to different modeling assumptions.

Technical details can be found in **Technical Details on Adult Coronary Artery Bypass Graft Surgery in the Commonwealth of Massachusetts: January 1 - December 31, 2004**



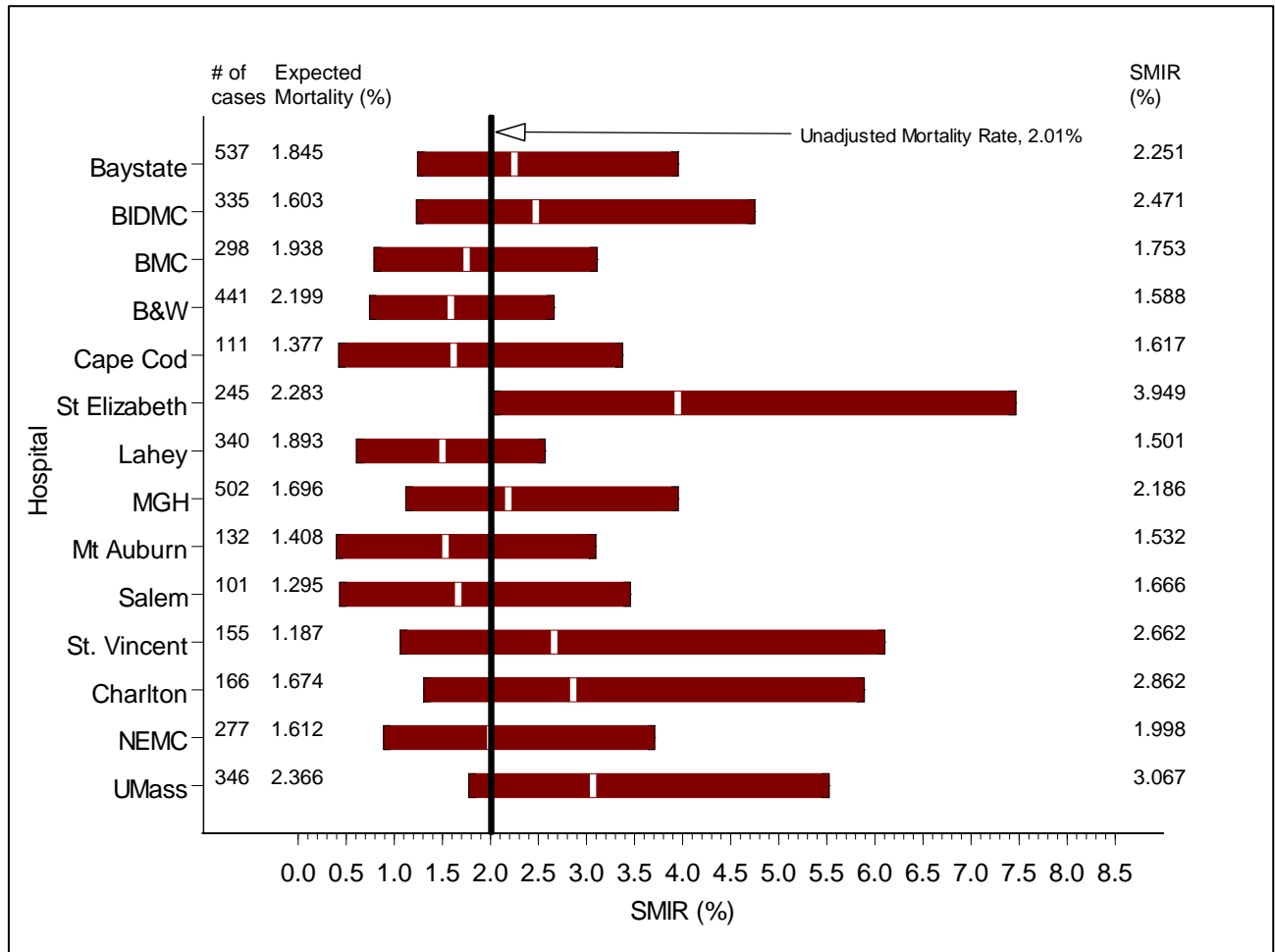
**Table 4.1: Prevalences and Adjusted Odds Ratios of 30-Day Mortality Following Isolated CABG Surgery in Adults, Massachusetts: 2004.** Based on 3986 surgeries with 80 deaths (2.01%). \* Average age of patients undergoing isolated CABG surgery is 65 +1.9 = 66.9 years of age.

Risk Factor	Prevalence (%)	Adjusted Odds Ratio	95% Interval for Adjusted Odds Ratio
Years over 65*	Mean = 1.9 yrs	1.05	(1.02, 1.07)
Male	74.5	0.64	(0.40, 1.11)
Renal Failure	5.8	3.83	(1.92, 6.65)
Diabetes	37.0	1.72	(0.99, 2.74)
Hypertension	82.7	0.67	(0.35, 1.26)
Peripheral Vascular Disease	17.7	3.55	(2.07, 5.54)
Prior CABG surgery	2.6	2.43	(0.67, 5.69)
Prior Percutaneous Coronary Intervention (PCI)	19.7	1.05	(0.54, 1.77)
Cardiogenic Shock	1.1	5.53	(1.36, 15.02)
Ejection Fraction (Ref = $\geq 40\%$ )			
< 30% or missing	11.8	3.97	(2.15, 6.65)
30 - 39	11.0	1.85	(0.82, 3.47)
Myocardial Infarction(MI)(Ref = None)			
< 6 hours	0.9	1.64	(0.11, 6.38)
7 - 24 hours	3.0	1.91	(0.50, 4.85)
1 - 7 days	22.1	0.95	(0.44, 1.76)
8 - 21 days	5.3	0.82	(0.22, 1.93)
> 21 days	21.8	1.71	(0.85, 3.05)
Status of CABG (Ref = Elective)			
Urgent	66.6	1.86	(0.87, 3.57)
Emergent/Salvage	3.1	2.15	(0.43, 6.13)
Pre-Op Intra-Aortic Balloon Pump	11.6	1.66	(0.73, 3.14)
<b>Between-Hospital Parameters</b>			
Between-Hospital Average logit, $\mu$		-5.94	-6.92, -5.00
Between-Hospital Variance in logits, $\tau^2$		0.3494	0.0136, 1.252

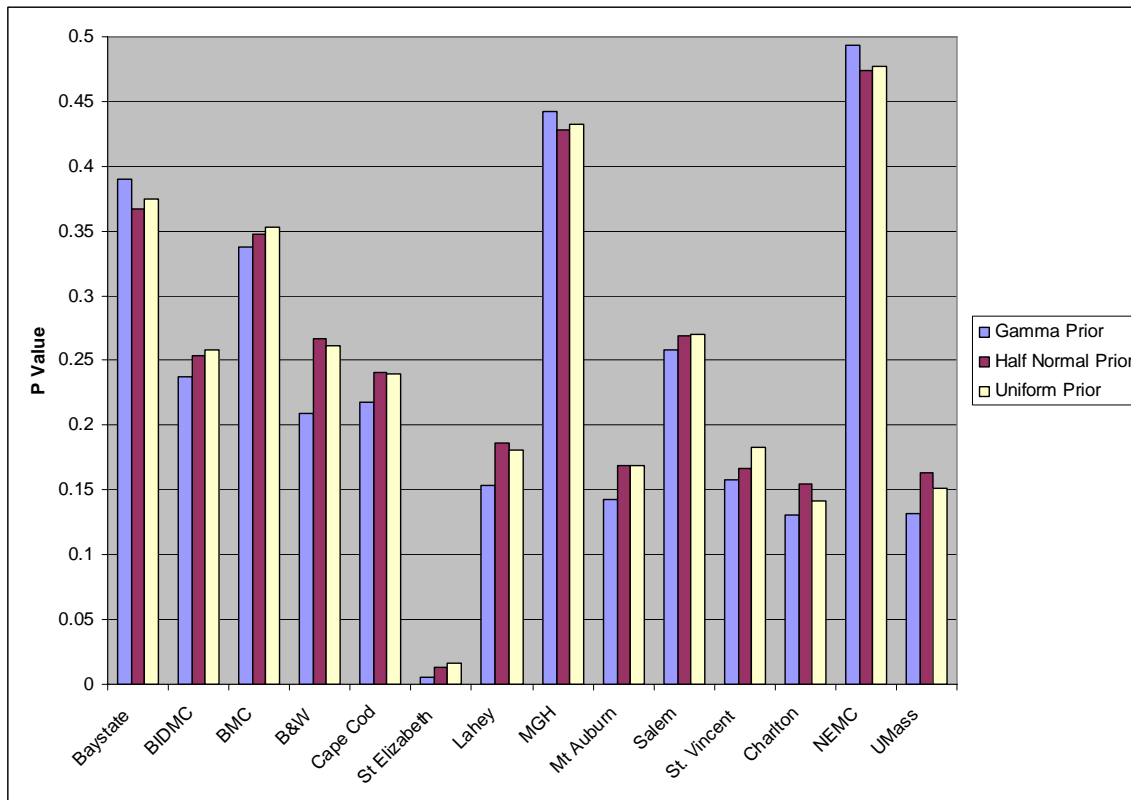
**Table 4.2: Expected 30-Day Mortality Rates, and Standardized 30-Day Mortality Incidence Rates (SMIR) point and interval estimates.** # of CABG cases refers to the number of Isolated CABG surgery admissions; expected mortality is the percentage of cases expected to die given the case-mix of the patients in the hospital. The SMIR is the hospital-specific standardized rate. Unadjusted state rate = 2.01%. \*\*Higher than expected mortality.

Hospital	# CABG Cases	30-Day Mortality (%)			
		Expected	Lower Limit of 95% SMIR Interval	SMIR	Upper Limit of 95% SMIR Interval
Baystate Medical Center	537	1.845	1.242	2.251	3.955
Beth Israel Deaconess Medical Center (BIDMC)	335	1.603	1.232	2.471	4.75
Boston Medical Center (BMC)	298	1.938	0.7889	1.753	3.109
Brigham & Women's Hospital (B&W)	441	2.199	0.7406	1.588	2.664
Cape Cod Hospital	111	1.377	0.421	1.617	3.375
Caritas Saint Elizabeth's Medical Center (St. Elizabeth)**	245	2.283	2.031	3.949	7.466
Lahey Clinic	340	1.893	0.6059	1.501	2.57
Massachusetts General Hospital (MGH)	502	1.696	1.12	2.186	3.955
Mount Auburn Hospital	132	1.408	0.3961	1.532	3.096
North Shore Medical Center-Salem Hospital (Salem)	101	1.295	0.4317	1.666	3.457
Saint Vincent Hospital at Worcester Medical Center (St. Vincent)	155	1.187	1.062	2.662	6.104
Southcoast Hospital Group-Charlton Memorial Hospital (Charlton)	166	1.674	1.307	2.862	5.889
Tufts- New England Medical Center (NEMC)	277	1.612	0.8899	1.998	3.709
UMass Memorial Medical Center (UMass)	346	2.366	1.776	3.067	5.524
<b>ALL</b>	<b>3986</b>			<b>2.01</b>	

**Figure 4.1. Ninety-Five Percent Posterior Intervals for Standardized Mortality Incidence Rates (SMIRs) Following Isolated CABG Surgery in Adults, Massachusetts, 2004.** # of cases refers to the number of isolated CABG surgery admissions; expected mortality rate is the percentage of cases expected to die given the case-mix of the patients in the hospital. The white vertical line in each box is the hospital's SMIR while the black vertical line denotes the unadjusted state 30-day mortality rate of 2.01%.



**Figure 4.2: Cross-Validated P-Values: Surgeries Performed in 2004.** P-values for each of the 14 cardiac surgery programs are listed on the y-axis; the x-axis identifies each hospital. Results are presented under a variety of assumptions for fitting the hierarchical regression model. For each hospital, p-values are presented assuming (from left to right) a gamma, half normal and uniform prior distribution, respectively.



## 5. IMPORTANT DEFINITIONS

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**Aortic Valve Repair:** Surgical repair of the aortic valve of the heart. The aortic valve is responsible for facilitating the flow of blood into the aorta.

**Aortic Valve Replacement:** A surgical procedure involving replacement of the aortic valve of the heart.

**Cardiac Catheterization:** A procedure that determines the extent and the location of the coronary artery obstruction or blockage.

**Cardiac Surgery** (as defined by the Massachusetts legislature for the Massachusetts Cardiac Study): Surgery on the heart and the thoracic great vessels. Examples of cardiac surgery include coronary artery bypass grafts, heart valve repair or replacement, heart transplantation, surgery of the thoracic aorta, repair of congenital heart defects, and minimally invasive heart surgery.

**Cardiovascular Disease:** Includes diseases of the heart or vessels that supply the body and the heart muscle with blood and oxygen.

**Coronary Artery Disease:** A disease affecting the coronary arteries in which the flow of oxygen containing blood to the heart muscle is partially or completely blocked, resulting in angina or a heart attack.

**Coronary Artery Bypass Graft [CABG] Surgery:** An operation in which the blocked coronary vessels are bypassed with the patient's own vessels to improve flow to the heart muscle. Coronary vessels are those vessels that supply the heart muscle with blood and oxygen.

**Cross-Validation:** Model validation is performed to ascertain whether predicted values from a statistical model are likely to accurately predict responses on future subjects or on

subjects not used to develop the analytical model. Cross-validation involves dropping a set of observations from the analytical process and the outcomes for the dropped set are predicted. This process is repeated many times in order to characterize the accuracy of the predictions.

**Mitral Valve Repair:** Surgical repair of the mitral valve of the heart. The mitral valve is responsible for facilitating the flow of blood from the left atrium into the left ventricle.

**Mitral Valve Replacement:** A surgical procedure which involves the replacement of the mitral valve of the heart.

**Percutaneous Coronary Intervention:** A non-surgical procedure designed to open and maintain the patency of obstructed coronary vessels. This treatment is an invasive procedure performed in the cardiac catheterization lab (e.g., outside of an operating room) by an interventional cardiologist in which a balloon, stent, or other device is delivered to the affected vessel to open and maintain its patency.

**Risk Factors:** Factors that contribute to an individual's risk of coronary artery disease or of death. These factors are classified as those that can be modified or changed by an individual, and those that cannot be changed. Examples of risk factors that cannot be modified include age, gender, family history of coronary artery disease, and ethnicity. Risk factors that can be controlled include diet, cholesterol levels, obesity, smoking, hypertension, inactive lifestyle, stress, and diabetes.

**Standardized Mortality Incidence Rate (SMIR):** The ratio of projected deaths (the number of deaths adjusted for the number of cases treated at the hospital and the hospital case mix) to expected deaths (the expected number of deaths calculated on the basis of the mortality experience of all cardiac surgery programs) multiplied by the state unadjusted rate. SMIRs are interpreted in terms of their corresponding probability intervals. If the probability interval includes the state rate, then the SMIR is no different from what was expected. If the interval excludes the state rate, then the SMIR is "significantly different" from what was expected. In this case, if the upper limit of the interval is lower than the

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state rate, then fewer patients than expected died; if the lower limit of the 95% interval is higher than the state rate, then more patients than expected died.

## **6 - ADVISORY COMMITTEES**

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Mass-DAC gratefully acknowledges the support from members of several Advisory Committees and physician volunteers (for data adjudication) who have donated their time to improve the quality of cardiac care in the Commonwealth of Massachusetts.



**The Massachusetts Cardiac Care Quality Advisory Commission** develops standards and criteria to be used by the Department of Public Health and Mass-DAC for the purpose of collecting, monitoring, and validating patient-specific outcome data from all hospitals in the Commonwealth of Massachusetts performing cardiac surgery or PCIs.

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David Torchiana, M.D.  
 Chairman and Chief Executive Officer  
 Massachusetts General Physicians  
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**The Mass-DAC Cardiac Advisory Board** advises Mass-DAC on data quality; on identification of risk factors affecting patient outcomes; and on appropriateness, interpretation, and limitations of analytic results.

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**The Mass-DAC Cardiac Surgery Data Adjudication Committees** review patient-specific data elements and corresponding data documentation submitted by hospitals to Mass-DAC in order to determine validity.

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The following Cardiac Surgeons generously volunteered their time to join the Cardiac Surgery Adjudication Committee in adjudicating the 2004 data:

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**The Massachusetts STS Quality and Outcomes Committee** advises Mass-DAC on surgeon and operator specific mortality.

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Massachusetts General Physicians Organization  
Boston, MA

**APPENDIX 1:**  
**STS DATA COLLECTION TOOL - VERSION 2.41**

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# The Society of Thoracic Surgeons Adult Cardiac Surgery Database Data Collection Form

Version 2.41

## A. Administrative

Participant ID: | | | | | | | | Cost Link Field: | | | | | | | | Optional STS Trial Link Number: | | | | | | | | Optional

## B. Demographics

Patient Medical Record Number: \_\_\_\_\_ not harvested

Last Name: \_\_\_\_\_

First: \_\_\_\_\_ MI: \_\_\_\_\_ not harvested

Date of Birth: \_\_\_\_/\_\_\_\_/\_\_\_\_ optional harvest

Age: \_\_\_\_\_ system calculation

Gender: (Male) (Female)

Race: (Caucasian) (Black) (Hispanic) (Asian) (Native American) (Other)

Social Security (or National ID ) Number: \_\_\_\_\_ not harvested

ZIP or Postal Code: \_\_\_\_\_ optional harvest

Referring Cardiologist's Name: \_\_\_\_\_ not harvested

Referring Physician's Name: \_\_\_\_\_ not harvested

## C. Hospitalization

Hospital Name: \_\_\_\_\_ controlled list Primary Payor: \_\_\_\_\_ not harvested

Date of Admission: \_\_\_\_/\_\_\_\_/\_\_\_\_

Date of Surgery: \_\_\_\_/\_\_\_\_/\_\_\_\_

Date of Discharge: \_\_\_\_/\_\_\_\_/\_\_\_\_

Same Day Elective Admission: No Yes

Initial ICU Hours: \_\_\_\_\_ Readmn to ICU: No Yes → if yes, Additional ICU Hours \_\_\_\_\_ Total Hours in ICU: \_\_\_\_\_ calculated

## D. Pre-Operative Risk Factors

Weight: \_\_\_\_\_ (kg) Height: \_\_\_\_\_ (cm)

Smoker: No Yes → if yes, Current Smoker: No Yes

Family History of CAD: No Yes

Diabetes: No Yes → if yes, select one: Diabetes Control: (None) (Diet) (Oral) (Insulin)

Hypercholesterolemia: No Yes

Last Creatinine Preop: \_\_\_\_\_

Renal Failure: No Yes → if yes, Dialysis: No Yes

Hypertension: No Yes

Cerebrovascular Accident: No Yes → if yes, When: (Recent <= 2 weeks) (Remote > 2 weeks)

Infectious Endocarditis: No Yes → if yes, Infectious Endocarditis Type: (Treated) (Active)

Chronic Lung Disease: (No) (Mild) (Moderate) (Severe)

Immunosuppressive Trtment: No Yes

Peripheral Vascular Disease: No Yes

Cerebrovascular Disease: No Yes → if yes, CVD Type: (Coma) (CVA) (RIND) (TIA) (Non Invasive > 75%) ( Previous Carotid Surgery)

## E. Previous Interventions

Previous CV Interventions: No Yes ↓ if yes, complete this section

# of Prior Cardiac Operations Requiring Cardiopulmonary Bypass: \_\_\_\_\_ # of Prior Cardiac Operations Without Cardiopulmonary Bypass: \_\_\_\_\_

Previous Surgery:

Coronary Artery Bypass: No Yes

Valve: No Yes

Previous Other Cardiac: No Yes

Prior PTCA including Balloon and/or Atherectomy: No Yes → if yes, Interval: <= 6 hours > 6 hours

Previous non-surgical Stent Placement: No Yes → if yes, Interval: <= 6 hours > 6 hours

Thrombolysis: No Yes → if yes, Interval: <= 6 hours > 6 hours

Previous non-surgical Balloon Valvuloplasty: No Yes

**F. Pre Operative Cardiac Status**

Myocardial Infarction: No Yes → if yes, When: (<= 6 hours) (> 6 hours but <24 hours) (1 - 7 days) (8 - 21 days) (> 21 days)

Congestive Heart Failure: No Yes

Angina: No Yes → if yes, Type: Stable Unstable ↓ if unstable

Unstable Type: (Rest Angina) (New Class 3) (Recent Accel) (Variant Angina) (Non-Q MI) (Post- Infarct Angina)

Cardiogenic Shock: No Yes → if yes Type: (Refractory Shock) (Hemodynamic Instability)

Resuscitation: No Yes

Arrhythmia: No Yes → if yes, Type: (Sust VT/VF) (Heart Block) (AFib/Flutter)

Classification: CCS: 0 I II III IV NYHA: I II III IV

**G. Pre Operative Medications**

Digitalis: No Yes	Beta Blockers: No Yes	Nitrates – I.V.: No Yes	Anticoagulants: No Yes	Diuretics: No Yes
Inotropic Agents: No Yes	Steroids: No Yes	Aspirin: No Yes	Ace Inhibitors: No Yes	Oth Anti-Platelets: No Yes

**H. Pre Operative Hemodynamics and Cath**

Number of Diseased Coronary Vessels: (None) (One) (Two) (Three)

Left Main Disease > 50%: No Yes

Ejection Fraction Done? No Yes → if yes, Ejection Fraction: \_\_\_\_\_ → Method: (LV gram) (Radionucleotide) (Estimate) (ECHO)

Pulmonary Artery Mean Pressure Done? No Yes → if yes, Pulmonary Artery Mean Pressure: \_\_\_\_\_

Aortic Stenosis: No Yes → if yes, Gradient: _____	Aortic Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe
Mitral Stenosis: No Yes	Mitral Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe
Tricuspid Stenosis: No Yes	Tricuspid Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe
Pulmonic Stenosis: No Yes	Pulmonic Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe

**J. Operative**

Surgeon's Name: \_\_\_\_\_ controlled list Surgeon Group: \_\_\_\_\_ controlled list

Status of the procedure:  
Emergent Salvage

Emergent → Reason: (Shock Circ Supp) (Shock No Circ Supp) (Pulm Edema) (AEMI) (Ongoing Ischemia) (Valve Dysfnctn) (Aortic Dissection)

Urgent → Reason: (AMI) (IABP) (Worsening CP) (CHF) (Anatomy) (USA) (Rest Angina) (Valve Dysfunction) (Aortic Dissection)

Elective

Coronary Artery Bypass: No Yes (if yes, complete Section K)

<u>Aortic:</u>	<u>Mitral:</u>	<u>Tricuspid:</u>	<u>Pulmonic:</u>
No	No	No	No
Replacement	Annuloplasty only	Annuloplasty Only	Replacement
Repair/Reconstruction	Replacement	Replacement	Reconstruction
Root Reconstruction Valve Conduit	Reconstruction w/ Annuloplasty	Reconstruction w/ Annuloplasty	
Reconstruction w/ Valve Sparing	Reconstruction w/out Annuloplasty	Reconstruction w/out Annuloplasty	
Resuspension Aortic Valve		Valvectomy	
Resection Sub-Aortic Stenosis			

Other Cardiac Procedure: No Yes ↓ (if yes, complete Section N)

Other Non-Cardiac Procedure: No Yes ↓ (if yes, complete Section O)

**K. Coronary Surgery**

Unplanned CABG: No Yes

Number of Distal Anastomoses with Arterial Conduits: \_\_\_\_\_

Number of Distal Anastomoses with Vein Grafts: \_\_\_\_\_

IMAs Used as Grafts: (Left IMA) (Right IMA) (Both IMAs) (No IMA)

Number of IMA Distal Anastomoses: \_\_\_\_\_

Radial Artery(ies) Used as Grafts: (No Radial) (Left Radial) (Right Radial) (Both Radials)

Number of Radial Artery Distal Anastomoses: \_\_\_\_\_

Number of Gastro-Epiploic Artery Distal Anastomoses: \_\_\_\_\_

L. Valve Surgery		↓ Key M = Mechanical, B = Bioprosthesis, H = Homograft, A = Autograft, R = Ring									
Aortic Prosthesis -	Implant Type:	None	M	B	H	A	R	Implant:	_____	Size:	____(mm)
	Explant Type:	None	M	B	H	A	R	Explant:	_____	Size:	____(mm)
Mitral Prosthesis -	Implant Type:	None	M	B	H	A	R	Implant:	_____	Size:	____(mm)
	Explant Type:	None	M	B	H	A	R	Explant:	_____	Size:	____(mm)
Tricuspid Prosthesis -	Implant Type:	None	M	B	H	A	R	Implant:	_____	Size:	____(mm)
	Explant Type:	None	M	B	H	A	R	Explant:	_____	Size:	____(mm)
Pulmonic Prosthesis -	Implant Type:	None	M	B	H	A	R	Implant:	_____	Size:	____(mm)
	Explant Type:	None	M	B	H	A	R	Explant:	_____	Size:	____(mm)

Valve Key	
<u>Mechanical</u> M1= ATS Mechanical Prosthesis M2= Björk-Shiley Convex-Concave Mechanical Prosthesis M3= Björk-Shiley Monostrut Mechanical Prosthesis M4= CarboMedics Mechanical Prosthesis M5= Edwards Tekna Mechanical Prosthesis M6= Lillehei-Kaster Mechanical Prosthesis M7= Medtronic-Hall Mechanical Prosthesis M8= OmniCarbon Mechanical Prosthesis M9= OmniScience Mechanical Prosthesis M10= On-X Mechanical Prosthesis M11= Sorin Bicarbon (Baxter Mira) Mechanical Prosthesis M12= Sorin Monoleaflet Allcarbon Mechanical Prosthesis M13= St. Jude Medical Mechanical Prosthesis M14= Starr-Edwards Caged-Ball Prosthesis M15= Ultracor Mechanical Prosthesis	B12= Hancock Modified Orifice Porcine Bioprosthesis B13= Ionescu-Shiley Pericardial Bioprosthesis B14= Labcor Stented Porcine Bioprosthesis B15= Labcor Stentless Porcine Bioprosthesis B16= Medtronic Freestyle Stentless Porcine Bioprosthesis B17= Medtronic Intact Porcine Bioprosthesis B18= Medtronic Mosaic Porcine Bioprosthesis B19= Mitroflow Pericardial Bioprosthesis B20= Sorin Pericarbon Stentless Pericardial Bioprosthesis B21= St. Jude Medical - Toronto SPV Stentless Porcine Bioprosthesis B22= St. Jude Medical-Bioimplant Porcine Bioprosthesis
<u>Bioprosthetic</u> B1= Baxter Prima Plus Stentless Porcine Bioprosthesis B2= Baxter Prima Stentless Porcine Bioprosthesis B3= Biocor Porcine Bioprosthesis B4= Biocor Stentless Porcine Bioprosthesis B5= CarboMedics PhotoFix Pericardial Bioprosthesis B6= Carpentier-Edwards Pericardial Bioprosthesis B7= Carpentier-Edwards Standard Porcine Bioprosthesis B8= Carpentier-Edwards Supra-Annular Porcine Bioprosthesis B9= Cryolife O'Brien Stentless Porcine Bioprosthesis B10= Hancock Standard Porcine Bioprosthesis B11= Hancock II Porcine Bioprosthesis	<u>Homograft</u> H1= Homograft Aortic – Subcoronary H2= Homograft Aortic Root/Cylinder H3= Homograft Mitral H4= Homograft Pulmonic Root H5= Cryolife Homograft
	<u>Autograft</u> A1= Autograft Pulmonic Root
	<u>Ring</u> R1= Carpentier-Edwards Classic Ring R2= Carpentier-Edwards Physio Ring R3= Cosgrove-Edwards Ring R4= Medtronic Sculptor Ring R5= Medtronic-Duran Ring R6= Sorin-Puig-Messana Ring R7= St. Jude Medical Sequin Ring
	777= Other

M. Operative Techniques	
Cardiopulmonary Bypass Used:	No Yes → if yes, Conversion to CPB: No Yes
Primary Indication for minimally Invasive approach:	(Surg/Pat Choice) (ContraindicatedStd Approach) (Comb Cath Intervention)
Primary Incision:	Full Sternotomy    Partial Sternotomy    Transverse Sternotomy    Right Vertical Parasternal    Left Vertical Parasternal Right Anterior Thoracotomy    Left Anterior Thoracotomy    Posterolateral Thoracotomy    Xiphoid    Epigastric    Subcostal
Total # of Incisions: _____	Conversion to Stnd Incision: No Yes → if yes, Indication: (Exposure) (Bleeding) (Rhythm) (Hypotension) (Conduit)
Cannulation Meth:	(Aorta and Fem/Jug Vein) (Fem Art and Fem/Jug Vein) (Aorta and Atrial/Caval) (Fem Art and Atrial/Caval) (Other)
Aortic Occlusion Method:	(None) (Cross-clamp) (Balloon Occlusion)
Intracoronary Shunt used during distal anastomoses:	No Yes
Suture Technique:	(Running) (Interrupted) (Stapler) (Combination)
Vessel Stabilization Technique:	(None) (Suture Snare) (Suction Device) (Compression) (Other)
IMA Harvest Technique:	(None) (Direct Vision) (Thoracoscopy) (Combination)
Acute Flow Patency Assess of Grafts (Periop):	(None) (IntaOp Doppler) (IntraOp Angio) (Postop Angio) (Postop Doppler)

N. Other Cardiac Procedures		
No Yes Left Ventricular Aneurysm Repair	No Yes Vent Septal Defect Repair	No Yes Atrial Septal Defect Repair
No Yes Batista	No Yes SVR	No Yes Congenital Defect Repair
No Yes Transmyocard Laser Revasc	No Yes Cardiac Trauma	No Yes Cardiac Transplant
No Yes Permanent Pacemaker	No Yes AICD	No Yes Other



O. Other Non Cardiac Procedures												
No	Yes	Aortic Aneurysm	No	Yes	Carotid Endarterectomy	No	Yes	Other Vascular	No	Yes	Other Thoracic	
P. CPB and Support												
Skin Incision Start Time: _____ 24 hour clock				Skin Incision Stop Time: _____ 24 hour clock								
Cross Clamp Time (min): _____				Perfusion Time (min): _____				Cardioplegia: No Yes				
IABP	No	Yes	→ if yes, When Inserted: (Preop) (Intraop) (Postop)									
	If yes, → Indication:		(Hemodynamic Instab)	(PTCA Support)	(Unst. Angina)	(CPB Wean)	(Prophylatic)					
Ventricular Assist Device:		No	Yes									
Q. Post Operative												
Blood Products Used:		No	Yes									
Initial # of Hrs Ventilated Postop: _____				Re-intubated During Hosp Stay: No Yes → if yes, Addl Hours Ventilated Postop: _____								
Total Hours Ventilated Postop:		_____										
R. Complications In hospital Complications: No Yes ↓ if yes, at least one complication below must be selected												
Operative	No	Yes	ReOp for Bleeding/Tamponade			Infection			No	Yes	Sternum – Deep	
	No	Yes	ReOp for Valvular Dysfunction						No	Yes	Thoracotomy	
	No	Yes	ReOp for Graft Occlusion						No	Yes	Leg	
	No	Yes	ReOp for Other Cardiac Problem						No	Yes	Septicemia	
	No	Yes	ReOp for Other Non Cardiac Problem						No	Yes	Urinary Tract Infection	
	No	Yes	Perioperative Myocardial Infarction									
Neurologic	No	Yes	Stroke			Pulmonary			No	Yes	Prolonged Ventilation	
	No	Yes	Transient						No	Yes	Pulmonary Embolism	
	No	Yes	Continuous Coma >=24Hrs						No	Yes	Pneumonia	
Renal	No	Yes	Renal Failure			Vascular			No	Yes	Vascular - Aortic Dissection	
	No	Yes	Dialysis						No	Yes	Iliac/Femoral Dissection	
										No	Yes	Acute Limb Ischemia
Other	No	Yes	Heart Block									
	No	Yes	Cardiac Arrest						No	Yes	Gastro-Intestinal Complication	
	No	Yes	Anticoagulant Complication						No	Yes	Multi-System Failure	
	No	Yes	Tamponade						No	Yes	Atrial Fibrillation	
S. Discharge (Note: this section is blank if patient dies during initial hospital stay)												
Aspirin: No Yes		Ace-Inhibitors: No Yes		Beta Blockers: No Yes		Lipid Lowering: No Yes		Other Anti-Platelets: No Yes				
Discharge Location: (Home)		(Extended Care/TCU)		(Other Hospital)		(Nursing Home)		(Other)				
T. Mortality												
Mortality - Mortality: No Yes		Discharge Status: Alive Dead			Status at 30 days after surgery: Alive Dead							
Mortality - Operative Death: No Yes		Mortality - Date ___/___/____ (mm/dd/yyyy)										
Location of Death: (OR) (Hospital) (Home) (Other Facility)												
Primary Cause of Death (select only one): (Cardiac) (Neurological) (Renal) (Vascular) (Infection) (Pulmonary) (Valvular) (Other)												
U. Readmission (Note: this section is blank if patient dies during initial hospital stay)												
Readmit <=30 Days from Date of Procedure: No Yes		↓ if yes, select the most predominate reason										
Readmission Reason:												
(Anticoagulant Complications)				(Arrhythmias/Heart Block/Pacemaker Insertion/AICD)				(CHF)				
(MI/Recurrent Angina)				(Pericardial Effusion/Tamponade)				(Pneumonia/ Respiratory Complication)				
(Valve Dysfunction)				(Infection Deep Sternum)				(Infection Leg)				
Cardiac Cath)				(PTCA Stent)				(Renal Failure)				
TIA)				(Reop for Graft Occlusion)				(Reop for Bleeding)				
(Permanent CVA)				(Acute Vascular Complication)				(Other)				

**APPENDIX 2:**  
**STS DATA COLLECTION TOOL - VERSION 2.52.1**

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(Variables not harvested by STS are harvested by Mass-DAC)



The Society of Thoracic Surgeons  
Adult Cardiac Surgery Database  
Data Collection Form  
Version 2.52.1

**A. Administrative**

Participant ID: |\_|\_|\_|\_|\_|\_|\_|\_| Record ID \_\_\_\_\_  
Cost Link Field: \_\_\_\_\_ STS Trial Link Number: |\_|\_|\_|\_|\_|\_|\_|\_| Patient ID \_\_\_\_\_

**B. Demographics**

Last Name: \_\_\_\_\_ First Name: \_\_\_\_\_ Patient M.I.: \_\_\_\_\_ **Name Fields Not Harvested**  
Date of Birth (mm/dd/yyyy): \_\_\_/\_\_\_/\_\_\_\_ Patient Age: \_\_\_\_\_ **System Calculation**  
Gender: Male Female  
Social Security (or National Patient ID) Number: \_\_\_\_\_ **Not Harvested** Medical Record Number: \_\_\_\_\_ **Not Harvested**  
Patient ZIP or Postal Code: \_\_\_\_\_ Race: Caucasian Black Hispanic Asian Native American Other  
Referring Cardiologist's Name: \_\_\_\_\_ **Not Harvested** Referring Physician's Name: \_\_\_\_\_ **Not Harvested**

**C. Hospitalization**

Hospital Name: \_\_\_\_\_ Hospital ZIP Code |\_\_\_\_\_| Hospital State |\_|\_|\_|  
Payor: \_\_\_\_\_ **Not Harvested**  
Date of Admission: \_\_\_/\_\_\_/\_\_\_\_ Date of Surgery: \_\_\_/\_\_\_/\_\_\_\_ Date of Discharge: \_\_\_/\_\_\_/\_\_\_\_  
ICU Visit: Yes No **If Yes, →** Initial ICU Hours: \_\_\_\_\_  
Readmn to ICU: Yes No **If Yes, →** Additional ICU Hours \_\_\_\_\_  
Total Hours in ICU: \_\_\_\_\_

**D. Risk Factors**

Weight (kg): \_\_\_\_\_ Height (cm): \_\_\_\_\_  
Smoker: Yes No **If Yes, →** Current Smoker: Yes No  
Family History of Coronary Artery Disease: Yes No  
Diabetes: Yes No **If Yes, select one: →** Diabetes Control: None Diet Oral Insulin  
Dyslipidemia: Yes No  
Last Creatinine Level Preop: \_\_\_\_\_  
Renal Failure: Yes No **If Yes, →** Dialysis: Yes No  
Hypertension: Yes No  
Cerebrovascular Accident: Yes No **If Yes, →** When: Recent <= 2 weeks Remote > 2 weeks  
Infectious Endocarditis: Yes No **If Yes, →** Infectious Endocarditis Type: Treated Active  
Chronic Lung Disease: No Mild Moderate Severe  
Immunosuppressive Therapy: Yes No  
Peripheral Vascular Disease: Yes No  
Cerebrovascular Disease: Yes No **If Yes, →** CVD Type: Coma CVA RIND TIA Non Invasive > 75% Prior Carotid Surgery

**E. Previous CV Interventions**

Incidence: First CV Surgery First Re-op CV Surgery Second Re-op CV Surgery Third Re-op CV Surgery Fourth or More Re-op Surgery  
Previous CV Interventions: Yes No **If Yes, complete the rest of this section ↓**  
Previous Coronary Artery Bypass: Yes No  
Previous Valve: Yes No  
Previous Other Cardiac – Intrapericardial or Great Vessel: Yes No  
Previous Other Cardiac – AICD: Yes No  
Previous Other Cardiac – Pacemaker: Yes No **If Yes, →** Previous Other Cardiac – Pacemaker Type: Biventricular Univentricular  
Previous Other Cardiac – PCI: Yes No **If Yes, →** Previous Other Cardiac – PCI Interval: <= 6 Hours > 6 Hours

**F. Preoperative Cardiac Status**

Myocardial Infarction: Yes No **If Yes, →** When: <= 6 hours > 6 hours but <24 hours 1 - 7 days 8 - 21 days > 21 days

Congestive Heart Failure: Yes No

Angina: Yes No **If Yes, →** Angina Type: Stable Unstable

Cardiogenic Shock: Yes No **If Yes, →** Cardiogenic Shock Type: Refractory Shock Hemodynamic Instability

Resuscitation: Yes No

Arrhythmia: Yes No **If Yes, →** Arrhythmia Type: Sust VT/VF Heart Block AFib/Flutter None

Classification - NYHA: I II III IV

**G. Preoperative Medications**

Beta Blockers: Yes No

ACE Inhibitors: Yes No

Nitrates I.V.: Yes No

Anticoagulants: Yes No **If Yes, →** Anticoagulants Medication Name: Heparin (Unfractionated) Heparin (Low Molecular) Thrombin Inhibitors

Coumadin: Yes No

Inotropes: Yes No

Steroids: Yes No

Aspirin: Yes No

Lipid-Lowering: Yes No **If Yes, →** Lipid Lowering Medication Name: Statin Non statin

ADP Inhibitors: Yes No

Glycoprotein IIb/IIIa Inhibitor: Yes No **If Yes, →** Glycoprotein IIb/IIIa Inhibitor Medication Name: Abciximab (ReoPro)  
Eptifibatid (Integrilin)  
Tirofiban (Aggrastat)

**H. Hemodynamics and Cath**

Number of Diseased Coronary Vessels: None One Two Three

Left Main Disease >= 50%: Yes No

Ejection Fraction Done? Yes No **If Yes, →** Ejection Fraction: \_\_\_\_\_

Method: LV gram Radionucleotide Estimate ECHO

Pulmonary Artery Mean Pressure Done? Yes No **If Yes, →** Pulmonary Artery Mean Pressure: \_\_\_\_\_

Aortic Stenosis: Yes No **If Yes, →** Gradient: \_\_\_\_\_

Mitral Stenosis: Yes No

Tricuspid Stenosis: Yes No

Pulmonic Stenosis: Yes No

Aortic Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe

Mitral Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe

Tricuspid Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe

Pulmonic Insufficiency: 0=None 1=Trivial 2=Mild 3= Moderate 4= Severe

**I. Operative**

Surgeon's Name: \_\_\_\_\_ Surgeon ID: \_\_\_\_\_

Status of the procedure: ↓

Elective

Urgent → Reason: AMI IABP Worsening CP CHF Anatomy USA Rest Angina

Valve Dysfunction Aortic Dissection Angiographic Accident

Emergent → Reason: Shock Circ Support Shock No Circ Support Pulmonary Edema AEMI

Ongoing Ischemia Valve Dysfunction Aortic Dissection Angiographic Accident

Emergent Salvage

Robotic Technology Assisted: Yes No

Coronary Artery Bypass: Yes No → If Yes, also complete Section J

Valve Surgery: Yes No → If Yes, also complete Section K

Ventricular Assist Device: Yes No → If Yes, also complete Section L

Other Cardiac Procedure: Yes No → If Yes, also complete Section M

Other Non-Cardiac Procedure: Yes No → If Yes, also complete Section N

Skin Incision Start Time: \_\_\_\_\_ 24 hour clock Skin Incision Stop Time: \_\_\_\_\_ 24 hour clock

CPB Utilization: None Combination Full ↓

If Combination, → Combination Plan: Planned Unplanned → If Unplanned, Unplanned Combination Reason: Exposure/visualization

- Bleeding
- Inadequate size and/or diffuse disease of distal vessel
- Hemodynamic Instability
- Conduit quality and/or trauma
- Other

If Combination or Full, → Perfusion Time (min): \_\_\_\_\_

- Cannulation Method: → Aorta and Fem/Jug Vein
- Fem Art and Fem/Jug Vein
- Aorta and Atrial/Caval
- Fem Art and Atrial/Caval
- Other

Aortic Occlusion: → None

Aortic Crossclamp → If Aortic Crossclamp or Balloon Occlusion, → Cross Clamp Time (min): \_\_\_\_\_

- Balloon Occlusion
- Partial Crossclamp

Cardioplegia: Yes No

IABP: Yes No → If Yes, When Inserted: → Preoperatively Intraoperatively Postoperatively

Indication: → Hemodynamic Instab PTCA Support Unstable Angina CPB Wean Prophylactic

Intraop Blood Products: Yes No → If Yes, Red Blood Cell Units \_\_\_\_\_

Fresh Frozen Plasma Units \_\_\_\_\_

Cryoprecipitate Units \_\_\_\_\_

Platelet Units \_\_\_\_\_

## J. Coronary Bypass

Number of Distal Anastomoses with Arterial Conduits: \_\_\_\_\_

Number of Distal Anastomoses with Venous Conduits: \_\_\_\_\_

Anastomotic Device Used: Yes No If Yes, → Anastomotic Device: Glue Magnets Clips Staples Other

IMAs Used as Grafts: Left IMA Right IMA Both IMAs No IMA If Left, Right, or Both ↓

IMA Harvest Technique: Direct Vision Thoracoscopy Combination Robotic Assisted

Number of IMA Distal Anastomoses: \_\_\_\_\_

Radial Artery Used: No Radial Left Radial Right Radial Both Radials If Left, Right, or Both ↓

Number of Radial Artery Distal Anastomoses: \_\_\_\_\_

Number of Gastro-Epiploic Artery Distal Anastomoses: \_\_\_\_\_

Number of Other Arterial Distal Anastomoses: \_\_\_\_\_

## K. Valve Surgery

<u>Aortic:</u>	<u>Mitral:</u>	<u>Tricuspid:</u>	<u>Pulmonic:</u>
No	No	No	No
Replacement	Annuloplasty Only	Annuloplasty Only	Replacement
Repair/Reconstruction	Replacement	Replacement	Reconstruction
Root Reconstruction w/ Valve Conduit	Reconstruction w/ Annuloplasty	Reconstruction w/ Annuloplasty	
Replacement + Aortic Graft Conduit	Reconstruction w/out Annuloplasty	Reconstruction w/out Annuloplasty	
Root Reconstruction w/ Valve Sparing		Valvectomy	
Resuspension Aortic Valve with replacement ascending Aorta			
Resuspension Aortic Valve without replacement ascending Aorta			
Resection Sub-Aortic Stenosis			

Annular Enlargement: Yes No

↓ Key M = Mechanical B = Bioprosthesis H = Homograft A = Autograft (Ross) R = Ring/Annuloplasty BA = Band/Annuloplasty

Aortic Prosthesis -	Implant Type:	None M B H A R BA	Implant: _____	Size: _____
Mitral Prosthesis -	Implant Type:	None M B H A R BA	Implant: _____	Size: _____
Tricuspid Prosthesis -	Implant Type:	None M B H A R BA	Implant: _____	Size: _____
Pulmonic Prosthesis -	Implant Type:	None M B H A R BA	Implant: _____	Size: _____

### Valve Key

#### Mechanical

ATS Mechanical Prosthesis = M1  
 Björk-Shiley Convex-Concave Mechanical Prosthesis = M2  
 Björk-Shiley Monostrut Mechanical Prosthesis = M3  
 CarboMedics Mechanical Prosthesis = M4  
 CarboMedics Carbo-Seal Ascending Aortic Valved Conduit Prosthesis = M16  
 CarboMedics Carbo-Seal Valsalva Ascending Aortic Valved Conduit Prosthesis = M17  
 CarboMedics Reduced Cuff Aortic Valve = M18  
 CarboMedics Standard Aortic Valve = M19  
 CarboMedics Top-Hat Supra-annular Aortic Valve = M20  
 CarboMedics OptiForm Mitral Valve = M21  
 CarboMedics Standard Mitral Valve = M22  
 CarboMedics Orbis Universal Valve = M23  
 CarboMedics Small Adult Aortic and Mitral Valves = M24  
 Edwards Tekna Mechanical Prosthesis = M5  
 Lillehei-Kaster Mechanical Prosthesis = M6  
 MCRI On-X Mechanical Prosthesis = M10  
 Medtronic-Hall/Hall Easy-Fit Mechanical Prosthesis = M7  
 Medtronic ADVANTAGE Mechanical Prosthesis = M25  
 OmniCarbon Mechanical Prosthesis = M8  
 OmniScience Mechanical Prosthesis = M9  
 Sorin Bicarbon (Baxter Mira) Mechanical Prosthesis = M11  
 Sorin Monoleaflet Allcarbon Mechanical Prosthesis = M12  
 St. Jude Medical Mechanical Prosthesis or St. Jude Medical® Mechanical Heart Valve = M13  
 SJM® Masters Series Mechanical Heart Valve = M26  
 SJM® Masters Series Aortic Valve Graft Prosthesis = M27  
 St. Jude Medical® Mechanical Heart Valve Hemodynamic Plus (HP) Series = M28  
 SJM® Masters Series Hemodynamic Plus Valve with FlexCuff™ Sewing Ring = M29  
 SJM Regent™ Valve = M30  
 Starr-Edwards Caged-Ball Prosthesis = M14  
 Ultracor Mechanical Prosthesis = M15

#### Bioprosthetic

Baxter Prima Stentless Porcine Bioprosthesis – Subcoronary = B24  
 Baxter Prima Stentless Porcine Bioprosthesis – Root = B25  
 Biocor Porcine Bioprosthesis = B3  
 Biocor Stentless Porcine Bioprosthesis – Subcoronary = B26  
 Biocor Stentless Porcine Bioprosthesis – Root = B27  
 CarboMedics PhotoFix Pericardial Bioprosthesis = B5  
 Carpentier-Edwards Duraflex Porcine Bioprosthesis = B28  
 Carpentier-Edwards Prima Plus Stentless Porcine Bioprosthesis – Subcoronary = B29  
 Carpentier-Edwards Prima Plus Stentless Porcine Bioprosthesis – Root = B30  
 Carpentier-Edwards PERIMOUNT Pericardial Bioprosthesis = B6  
 Carpentier-Edwards Standard Porcine Bioprosthesis = B7  
 Carpentier-Edwards Supra-Annular Aortic Porcine Bioprosthesis = B8  
 Cryolife O'Brien Stentless Porcine Bioprosthesis – Subcoronary = B31  
 Cryolife O'Brien Stentless Porcine Bioprosthesis – Root = B32  
 Hancock Standard Porcine Bioprosthesis = B10  
 Hancock II Porcine Bioprosthesis = B11

Hancock Modified Orifice Porcine Bioprosthesis = B12  
 Ionescu-Shiley Pericardial Bioprosthesis = B13  
 Labcor Stented Porcine Bioprosthesis = B14  
 Labcor Stentless Porcine Bioprosthesis – Subcoronary = B33  
 Labcor Stentless Porcine Bioprosthesis – Root = B34  
 Medtronic Freestyle Stentless Porcine Bioprosthesis – Subcoronary = B35  
 Medtronic Freestyle Stentless Porcine Bioprosthesis – Root = B36  
 Medtronic Intact Porcine Bioprosthesis = B17  
 Medtronic Mosaic Porcine Bioprosthesis = B18  
 Medtronic Contegra Bovine Jugular Bioprosthesis = B37  
 Mitroflow Pericardial Bioprosthesis = B19  
 St. Jude Medical - Toronto SPV Stentless Porcine Bioprosthesis or SJM Toronto SPV® Valve = B21  
 St. Jude Medical-Bioimplant Porcine Bioprosthesis = B22  
 SJM Biocor™ Valve = B38  
 SJM Epic™ Valve = B39  
 SJM Toronto Root™ Bioprosthesis = B40  
 Sorin Pericarbon Stentless Pericardial Bioprosthesis = B20

#### Homograft

CryoLife Aortic Homograft = H6  
 CryoLife Pulmonary Homograft = H7  
 CryoLife CryoValve SG(Decellularized)Aortic Homograft = H8  
 CryoLife CryoValve SG Pulmonary Homograft = H9  
 Homograft Aortic – Subcoronary = H1  
 Homograft Aortic Root = H2  
 Homograft Mitral = H3  
 Homograft Pulmonic Root = H4  
 LifeNet CV Allografts = H10

#### Autograft

Pulmonary Autograft to aortic root (Ross Procedure) = A1

#### Ring - Annuloplasty

CarboMedics AnnuloFlo Ring = R8  
 CarboMedics AnnuloFlex Ring = R9  
 CarboMedics CardioFix Bovine Pericardium with PhotoFix Technology = R10  
 Carpentier-Edwards Classic Annuloplasty Ring = R1  
 Carpentier-Edwards Physio Annuloplasty System Ring = R2  
 Cosgrove-Edwards Annuloplasty System Ring = R3  
 Edwards MC<sup>3</sup> Tricuspid Annuloplasty System G Future Band = R11  
 Genesee Sculptor Annuloplasty Ring = R12  
 Medtronic Sculptor Ring = R4  
 Medtronic-Duran AnCore Ring = R5  
 Sorin-Puig-Messana Ring = R6  
 St. Jude Medical Sequin Ring or SJM® Séguin Annuloplasty Ring = R7  
 SJM Tailor™ Annuloplasty Ring = R13

#### Band – Annuloplasty

Medtronic Colvin Galloway Future Band = Ba1  
 Medtronic Duran Band = Ba2  
 Medtronic Duran – Ancore Band = Ba3

Other = 777

L. **VAD**

Previous VAD: Yes No

Please note that future references to "initial VAD" refer to the initial VAD for this hospitalization, not a VAD placed during a previous hospitalization.

**Current Circulatory Support: For Initial VAD only**

Indication for VAD: (Bridge to Transplant) (Bridge to Recovery) (Destination) (Separation from CPB) (Device Malfunction)

Intubated Pre VAD: Yes No

Hemodynamics Pre VAD: May be obtained Prior to induction in the OR, or in an ICU immediately prior to OR

PCWP: \_\_\_mm/Hg CVP: \_\_\_mm/Hg PVR: \_\_\_woods units CI: \_\_\_L/ (min x m2)

RV Function: (Normal) (Mildly Impaired) (Moderately Impaired) (Severely Impaired)

RV Function method: \_\_\_ (Pre-op ECHO) (Intra-op pre VAD TEE)

VO2 Measured: Yes No

Peak VO2: \_\_\_ml/kg/min

VAD Device Data:

Implant Type: Fill in below: (RVAD) (LVAD) (BiVAD)

Product Type: Fill in below: 1. HeartQuest VAD 2. Lion Heart 3. Novacor LVAS 4. Heartsaver VAD 5. Jarvik 2000 6. DeBakey VAD 7. TandemHeart pVAD 8. AB-180 iVAD 9. CardioWest TAH 10. Thoratec iVAD 11. HeartMate VE 12. HeartMate IP LVAS 13. HeartMate SNAP-VE 14. HeartMate XVE 15. HeartMate II 16. HeartMate III 17. BVS5000i 18. AbioCor 19. InCor 20. Excor 21. Other

Explant Reason: Fill in below: 1. Cardiac Transplant 2. Recovery 3. Device Transfer 4. Device Related Infection 5. Device Malfunction

**Initial Implant Data**

<u>Implant Type</u>	<u>Product Type</u>	<u>Implant Date</u>	<u>Explant</u>	<u>Explant Date</u>	<u>Explant Reason</u>	<u>Cardiac Tx</u>	<u>Tx Date</u>
_____	_____	___/___/___	Y N	___/___/___	_____	Y N	___/___/___

Initial VAD Cannulation/Attachment Sites:

LVAD Inflow: (LA) (LV)

RVAD Inflow: (RA) (RV)

**Additional Implant(s) Data**

<u>Implant(s) Type</u>	<u>Product Type</u>	<u>Implant Date</u>	<u>Explant</u>	<u>Explant Date</u>	<u>Explant Reason</u>	<u>Cardiac Tx</u>	<u>Tx Date</u>
_____	_____	___/___/___	Y N	___/___/___	_____	Y N	___/___/___
_____	_____	___/___/___	Y N	___/___/___	_____	Y N	___/___/___

**Primary VAD Complications Data:**

Intracranial Bleed:	Yes	No
Embolic Stroke:	Yes	No
Driveline/Cannula Infection:	Yes	No
Pump Pocket Infection:	Yes	No
VAD Endocarditis:	Yes	No
Device Malfunction:	Yes	No

Additional Complications (not specific to initial VAD as above) to be collected in section "P", Complications.

**VAD Status:** Discharged from hospital: (with VAD) (without VAD)

**M. Other Cardiac Procedures**

Yes	No	Left Ventricular Aneurysm Repair	Yes	No	Ventricular Septal Defect Repair	Yes	No	Atrial Septal Defect Repair
Yes	No	Batista	Yes	No	Surgical Ventricular Restoration	Yes	No	Congenital Defect Repair
Yes	No	Transmyocard Laser Revasc	Yes	No	Cardiac Trauma	Yes	No	Cardiac Transplant

Arrhythmia Correction Surgery → None

- Permanent Pacemaker
- Permanent Pacemaker with Cardiac Resynchronization Therapy (CRT)
- Implanted Cardioverter Defibrillator (ICD)
- ICD with CRT

If "Permanent Pacemaker with CRT" or "ICD with CRT", then answer ↓

Arrhythmia Correction Surgery – Lead Placement → Epicardial Endocardial

Atrial Fibrillation Correction Surgery → None

- Standard Surgical Maze Procedure
- Other Surgical Ablative Procedure
- Combination of Standard and Other [If Other or Combo, then answer ↓](#)

Atrial Fibrillation Surgery – Energy Source →

- Unipolar Radiofrequency
- Bipolar Radiofrequency
- Microwave
- Cryothermia
- Other
- Combination of above

Yes	No	Aortic Aneurysm	<a href="#">If Yes, →</a>	Yes	No	Ascending Aorta
				Yes	No	Aortic Arch
				Yes	No	Descending Aorta
				Yes	No	Thoracoabdominal Aorta

Yes No Other

**N. Other Non Cardiac Procedures**

Yes	No	Carotid Endarterectomy	Yes	No	Other Vascular	Yes	No	Other Thoracic	Yes	No	Other
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**O. Post Operative**

Blood Products Used Postoperatively: Yes No → [If Yes,](#)

- Red Blood Cell Units \_\_\_\_\_
- Fresh Frozen Plasma Units \_\_\_\_\_
- Cryoprecipitate Units \_\_\_\_\_
- Platelet Units \_\_\_\_\_

Extubated in OR: Yes No [If No, →](#) Initial # Hrs Ventilated Postop: \_\_\_\_\_

Re-intubated During Hosp Stay: Yes No [If Yes, →](#) Addl Hours Ventilated Postop: \_\_\_\_\_

Total Hours Ventilated Postop: \_\_\_\_\_

**P. Complications** In Hospital Complications: Yes No

Operative:

- Yes No ReOp for Bleeding Tamponade
- Yes No ReOp for Valvular Dysfunction
- Yes No ReOp for Graft Occlusion
- Yes No ReOp for Other Cardiac Problem
- Yes No ReOp for Other Non Cardiac Problem
- Yes No Perioperative MI

Infection:

- Yes No Sternum – Deep
- Yes No Thoracotomy
- Yes No Leg
- Yes No Septicemia



Neurologic:  
Yes No Postoperative Stroke for >72 hours  
Yes No Transient Neurologic Deficit  
Yes No Continuous Coma >=24Hrs

Pulmonary:  
Yes No Prolonged Ventilation  
Yes No Pulmonary Embolism  
Yes No Pneumonia

Renal:  
Yes No Renal Failure [If Yes, ↓](#)  
Yes No Dialysis (Newly Required)

Vascular:  
Yes No Illiac/Femoral Dissection  
Yes No Acute Limb Ischemia

Other:  
Yes No Heart Block  
Yes No Cardiac Arrest  
Yes No Anticoagulant Complication  
Yes No Tamponade  
Yes No Gastro-Intestinal Complication

Yes No Multi-System Failure  
Yes No Atrial Fibrillation  
Yes No Aortic Dissection  
Yes No Other

#### Q. Mortality

Mortality: Yes No Discharge Status: Alive Dead Status at 30 days after surgery: Alive Dead Unknown  
Operative Death: Yes No [Only answered if Mortality = Yes](#)  
Mortality - Date \_\_\_/\_\_\_/\_\_\_ (mm/dd/yyyy) [Only answered if Mortality = Yes](#)  
Location of Death: OR during initial surgery Hospital Home Other Care Facility OR during reoperation [Only answered if Mortality = Yes](#)  
Primary Cause of Death (select only one): [Only answered if Mortality = Yes](#)  
Cardiac Neurologic Renal Vascular Infection Pulmonary Valvular Unknown Other

#### R. Discharge (Note: This section is only answered if Discharge Status is "Alive")

ADP Inhibitors: Yes No

Antiarrhythmics: Yes No [If Yes, ↓](#)

Antiarrhythmics – Discharge – Medication Name: Amiodarone Other

Aspirin: Yes No

Ace-Inhibitors: Yes No

Beta Blockers: Yes No

Lipid Lowering: Yes No [If Yes, ↓](#)

Lipid Lowering – Discharge – Medication Type: Statin Non statin

Coumadin: Yes No

Discharge Location: Home Extended Care/TCU Other Hospital Nursing Home Other

Cardiac Rehabilitation Referral: Yes No Not Applicable

Smoking Cessation Counseling: Yes No Not Applicable

S. **Readmission** (Note: This section is only answered if Discharge Status is "Alive")

Readmit <=30 Days from Date of Procedure: Yes No ↓ If Yes, select the primary reason and procedure

Readmit Reason:

- Anticoagulation Complication – Valvular
- Anticoagulation Complication - Pharmacological
- Arrhythmias/Heart Block
- Congestive Heart Failure
- Myocardial Infarction and/or Recurrent Angina
- Pericardial Effusion and/or Tamponade
- Pneumonia or other Respiratory Complication
- Coronary Artery Dysfunction
- Valve Dysfunction
- Infection - Deep Sternum
- Infection – Conduit Harvest Site
- Renal Failure
- TIA
- Permanent CVA
- Acute Vascular Complication
- Subacute Endocarditis
- VAD Complication
- Other – Related Readmission
- Other – Nonrelated Readmission

Readmit Reason – Primary Procedure:

- OR for Bleeding
- Pacemaker Insertion/AICD
- PCI
- Pericardiotomy/Pericardiocentesis
- OR for Coronary Arteries
- OR for Valve
- OR for Sternal Debridement/Muscle Flap
- Dialysis
- OR for Vascular
- No Procedure Performed
- Other Procedure
- Unknown