de cardiologie

Communauté. Connaissances. Leadership.

THE CANADIAN CARDIOVASCULAR SOCIETY **DATA DICTIONARY**

A CCS Consensus Document

HEART FAILURE DATA ELEMENTS AND DEFINITIONS

FINAL VERSION v1.1

Last Updated: July 16, 2013

Copyright © 2013 The Canadian Cardiovascular Society This publication may not be reproduced or modified without the permission of The Canadian Cardiovascular Society.

For authorised reproduction, please obtain permission from:

The Canadian Cardiovascular Society 222 Queen Street, Suite 1403 Ottawa. Ontario Canada K1P 5V9

Email: healthpolicy@ccs.ca

Background

The Canadian Cardiovascular Society Data Dictionary is comprised of multiple "chapter" data elements and definitions that reflect national input and consensus on definitions within several spheres of cardiovascular disease, treatment and subspecialty expertise.

THIS CHAPTER'S DATA ELEMENTS AND DEFINITIONS ARE SPECIFIC TO **HEART FAILURE** AND SHOULD BE CONSIDERED AS A SUPPLEMENT TO THE CORE ELEMENTS CHAPTER.

This Dictionary also contains the supporting data elements and definitions for the Canadian Cardiovascular Society Quality Indicators E-Catalogues for Heart Failure. The data elements in this dictionary have been identified into classification levels as follows:

ESSENTIAL: Are deemed as a minimum recommended data element to be used as a standard to enable reporting of key quality indicators and to allow cross-comparison with other centres using these common data elements.

SPECIALIZED: Are deemed as an expanded set of recommended data element to be used for more in-depth data collection and analysis.

Visit www.ccs.ca/ for the latest version of the CCS Quality Indicators E-Catalogues for HF and other areas.

Definitions

The following are the definitions of terminology and key outcome indicators used throughout this data dictionary, as well as terminology and key outcome indicators used by Heart Failure Quality Indicators E-Catalogue.

| Term | Definition | | |
|--|--|--|--|
| TERMINOLOGY | | | |
| Heart Failure (HF) | Clinical syndrome characterized by abnormal cardiac systolic and/or diastolic function and resulting symptoms of low cardiac output of venous congestion.or congestion due to abnormal myocardial function. | | |
| Acute Heart Failure | Heart failure with new or worsening signs and symptoms which develop over a period of less than 30 days. | | |
| Chronic Heart | Same as 'Congestive Heart Failure (CHF)' | | |
| Failure | Same as stable chronic heart failure | | |
| Newly Diagnosed HF | Heart failure diagnosed for the first time. | | |
| | Date of Newly Diagnosed HF (YYYYMMDD) | | |
| Electrocardiographic documentation (ECG) | 12-lead ECG, rhythm strip, Holter monitor, intracardiac electrograms, event recorder, or electrical cardiac activity measured by any implanted device that does not have intracardiac electrode. | | |
| Rheumatic mitral valve disease | As rheumatic mitral regurgitation cannot be reliably diagnosed without a pathological specimen, "rheumatic mitral valve disease" is defined as mitral stenosis (usually an echocardiographic diagnosis) that is moderate or greater in severity (valve area ≤1.5 cm² [Source: BSE Echocardiography: Guidelines for Valve Quantification]). | | |
| Qualifying HF | Episode of HF that resulted in first entry into the database, regardless of whether or not it is newly diagnosed. Date of Qualifying HF (YYYYMMDD) | | |
| HF Management | Method of heart failure treatment which includes ongoing follow up, provision of patient and family education, provision of self-care skill teaching, | | |
| Documented History | The patient has been told by a physician that they clearly have this diagnosis or there is a medicarecord of this diagnosis/event. | | |

| | KEY OUTCOME INDICATORS | | | |
|------------|--|--|--|--|
| IV Therapy | Any therapy provided via the intravenous route with the intention to effect control improvement of heart failure such as | | | |
| | Sodium (Na), Potassium (K), BUN and Creatinine. | | | |
| Stroke | Stroke is an acute onset of a focal neurologic deficit of presumed vascular origin lasting for ≥24 hours or resulting in death. Stroke [is] [can be] categorized as ischemic or hemorrhagic or cause unknown (based on computed tomographic or magnetic resonance scanning or autopsy) [but in this instance all strokes are included]. Fatal stroke is defined as death from any cause within 30 days of stroke. [Modified from Source: Am Heart J 2009;157:810.e1] | | | |
| | Stroke must be confirmed by imaging of the brain (computed tomographic or magnetic resonance scanning) or by autopsy. | | | |
| | Date of Stroke (YYYYMMDD): date of onset of symptoms of stroke | | | |
| | CHA ₂ DS ₂ VASc score at time of stroke = CHA ₂ DS ₂ -VASc Score | | | |
| | 1. Score = 0 | | | |
| | 2. Score = 1 | | | |
| | 3. Score = 2 | | | |
| | 4. Score = 3 | | | |
| | 5. Score = 4 or greater | | | |
| | 6. Score Unknown/Uncertain | | | |
| | Antithrombotic therapy at time of stroke = Antithrombotic Therapy | | | |
| | No antithrombotic therapy | | | |
| | 2. Anticoagulation only | | | |
| | i. Warfarin or other vitamin K antagonist | | | |
| | ii. Dabigatran | | | |
| | iii. Rivaroxaban | | | |
| | iv. Apixaban | | | |
| | 3. Antiplatelet only | | | |
| | Anticoagulation and antiplatelet | | | |
| | i. Warfarin or other vitamin K antagonist | | | |
| | ii. Dabigatran | | | |
| | iii. Rivaroxaban | | | |
| | iv. Apixaban | | | |
| | 5. Unknown/Uncertain | | | |

Contraindication to Anticoagulation

List of examples from the ROCKET AF Study:

- Active internal bleeding
- History of, or condition associated with, increased bleeding risk, including:
 - Major surgical procedure or trauma within 30 days before randomization
 - Clinically significant gastrointestinal bleeding within 6 months before randomization
 - History of intracranial, intraocular, spinal, or atraumatic intra-articular bleeding
 - Chronic hemorrhagic disorder
 - o Known intracranial neoplasm, arteriovenous malformation, or aneurysm
 - Planned invasive procedure with potential for uncontrolled bleeding, including major surgery

[Source: Am Heart J 2010;159:340-7.e1]

Date when Contraindication was First Noted (YYYYMMDD)

Systemic Embolus

Systemic embolism is an acute vascular occlusion of the extremities or any organ (kidneys, mesenteric arteries, spleen, retina or grafts) and must be documented by angiography, surgery, scintigraphy, or autopsy. [Modified from Source: Am Heart J 2009;157:810.e1]

Date of Systemic Embolus (YYYYMMDD): date of the onset of symptoms of systemic embolus

CHA₂DS₂VASc Score at time of Systemic Embolus = CHA₂DS₂-VASc Score

- 1. Score = 0
- 2. Score = 1
- 3. Score = 2
- 4. Score = 3
- 5. Score = 4 or greater
- 6. Score Unknown/Uncertain

Antithrombotic Therapy at Time of Systemic Embolus = Antithrombotic Therapy

- 1. No antithrombotic therapy
- 2. Anticoagulation only
 - i. Warfarin or other vitamin K antagonist
 - ii. Dabigatran
 - iii. Rivaroxaban
 - iv. Apixaban
- Antiplatelet only
- 4. Anticoagulation and antiplatelet
 - i. Warfarin or other vitamin K antagonist
 - ii. Dabigatran
 - iii. Rivaroxaban
 - iv. Apixaban
- 5. Unknown/Uncertain

TIA

Same as stroke but symptoms resolve within <24h and no imaging evidence of cerebral infarct or hemorrhage.

Date of TIA (YYYYMMDD): date of onset of symptoms of TIA

CHA₂DS₂VASc at Time of TIA = CHA₂DS₂-VASc Score

- 1. Score = 0
- 2. Score = 1
- 3. Score = 2
- 4. Score = 3
- 5. Score = 4 or greater
- 6. Score Unknown/Uncertain

Antithrombotic Therapy at Time of TIA = Antithrombotic Therapy

- 1. No antithrombotic therapy
- 2. Anticoagulation only
 - i. Warfarin or other vitamin K antagonist
 - ii. Dabigatran
 - iii. Rivaroxaban
 - iv. Apixaban
- 3. Antiplatelet only
- 4. Anticoagulation and antiplatelet
 - i. Warfarin or other vitamin K antagonist
 - ii. Dabigatran
 - iii. Rivaroxaban
 - iv. Apixaban
- 5. Unknown/Uncertain

Major Hemorrhage

Major hemorrhage is defined by ≥1 of the following criteria:

- Overt bleeding associated with reduction in haemoglobin level of at least 2.0 g/L;
- Overt bleeding leading to transfusion of at least 2 U of blood or packed cells; or
- Symptomatic bleeding in a critical area or organ such as intraocular, intracranial, intraspinal or intramuscular with compartment syndrome, retroperitoneal bleeding, intraarticular bleeding, or pericardial bleeding.

[Modified from Source: Am Heart J 2009;157:810.e2]

In the AF Quality Indicators e-Catalogue the Cross-sectional Analysis is based on hospitalization for major hemorrhage as defined above.

Date of Major Bleeding (YYYYMMDD) = date of onset of symptoms of bleeding or detection of overt bleeding when asymptomatic

CHA2DS2VASc at Time of TIA = CHA2DS2-VASc Score

- 1. Score = 0
- 2. Score = 1
- 3. Score = 2
- 4. Score = 3
- 5. Score = 4 or greater
- 6. Score Unknown/Uncertain

Antithrombotic Therapy at Time of Major Hemorrhage = Antithrombotic Therapy

- 1. No antithrombotic therapy
- 2. Anticoagulation only
 - i. Warfarin or other vitamin K antagonist
 - ii. Dabigatran
 - iii. Rivaroxaban
 - iv. Apixaban
- 3. Antiplatelet only
- 4. Anticoagulation and antiplatelet
 - Warfarin or other vitamin K antagonist
 - ii. Dabigatran
 - iii. Rivaroxaban
 - iv. Apixaban
- 5. Unknown/Uncertain

CV Hospitalization

Primary reason for hospitalization was cardiovascular categorized by reason(s) for hospitalization (Check all that apply):

- 1. Rhythm management of AF/AFL
- 2. Bleeding
- 3. Acute HF
- 4. MI
- 5. Other Acute Coronary Syndrome
- 6. Rhythm management for other SVT
- 7. Bradycardia Management
- 8. Rhythm management for VT/VF/SCD
- 9. Other, specify

Date of CV Hospitalization (YYYYMMDD)

| Non-CV Hospitalization Only | Primary reason for hospitalization was non-cardiovascular and no secondary CV problem during hospitalization | | | | |
|---|---|--|--|--|--|
| | Date of Non-CV Hospitalization (YYYYMMDD) | | | | |
| Non-CV Hospitalization with Secondary CV Problem | Primary reason for hospitalization was non-cardiovascular but a secondary cardiovascular problem developed during hospitalization categorized by CV problem(s) (Check all that apply): 1. Rhythm management for AF/AFL 2. Bleeding 3. Acute HF 4. MI 5. Other Acute Coronary Syndrome 6. Rhythm management for other SVT 7. Bradycardia Management 8. Rhythm management for VT/AF/SCD 9. Other, specify Date of Non-CV Hospitalization with Secondary CV Problem (YYYYMMDD) | | | | |
| CV Emergency Department Visit (whether or not followed by hospital admission) | Primary reason for Emergency Department Visit was cardiovascular categorized by reason(s) for ER Visit (Check all that apply): 1. Rhythm management of AF/AFL 2. Bleeding 3. Acute HF 4. MI 5. Other Acute Coronary Syndrome 6. Rhythm management for other SVT 7. Bradycardia management 8. Rhythm management for VT/VF/SCD 9. Other, specify Date of CV Emergency Department Visit (YYYYMMDD) | | | | |
| Lost to Follow-up | Patient is permanently lost to any further follow-up due to moving or any other administrative or other reason they are no longer included in the database. | | | | |
| | Date of Last Contact (YYYYMMDD) | | | | |

TABLE OF CONTENTS

| PART 1 – DEMOGRAPHICS | 10 |
|--|----------------------|
| PART 2 – HISTORY & RISK FACTORS | 11 14 |
| PART 3 – SYMPTOMS AND FUNCTIONAL ASSESSMENT | 16 |
| PART 4 – PHYSICAL EXAM AND VITAL SIGNS FOR THIS EPISODE OF CARE | 18 |
| PART 5 – MEDICATIONS | 19 23 |
| PART 6 – DEVICESA. IMPLANTABLE CARDIAC DEVICES (PACEMAKERS, DEFIBRILLATORS AND MONITORING DEVICES) B. MECHANICAL CARDIAC ASSISTS DEVICES | 33 |
| PART 7 – LABORATORY RESULTS | 37 |
| PART 8 – DIAGNOSTIC TESTS | 40 |
| PART 9 – COUNSELLING | 44 |
| PART 10 – CARE EPISODE | 45 46 46 48 |
| PART 11 – OUTCOMES | 51 |
| ACKNOWLEDGEMENT | 52 |
| DISCLAIMER | 52 |
| COPYRIGHT | 52 |

PART 1 – DEMOGRAPHICS

| Note: All Demographic data elements and definitions are defined in the Core Elements Data Dictionary Chapter. |
|---|
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |
| |

PART 2 – HISTORY & RISK FACTORS

- INDICATE DATE DATA COLLECTED (YYYYMMDD)

NOTE: The following data elements and definitions are specific to this Chapter and should be considered as a supplement to History & Risk Factors section defined in the Core Elements Data Dictionary Chapter.

A. MEDICAL HISTORY AND COMORBIDITIES

NOTE: Includes any occurrence between birth and this episode of care, unless otherwise indicated.

| FIELD NAME | CLASSIFICATION | DEFINITION |
|--------------------------|----------------|---|
| History of Dementia | SPECIALIZED | Patient has a documented history of dementia, Alzheimer's disease, chronic confusion (at least one month in duration), or senility. 1. Yes 2. No |
| | 00501411750 | 3. Unknown |
| History of Depression | SPECIALIZED | Patient has a documented history of treated depression, or is currently taking antidepressant medication. |
| | | 1. Yes 2. No |
| | | 3. Unknown |
| History of Sleep | SPECIALIZED | Patient has a documented history of sleep apnea. |
| Apnea | 01 2011 (21222 | 1. Yes |
| | | If yes, specify: |
| | | a. Central |
| | | b. Obstructive |
| | | c. Mixed |
| | | d. Unknown |
| | | If yes, are they on CPAP/BiPAP |
| | | a. Yes |
| | | b. No 2. No |
| | | 3. Unknown |
| History of Liver | SPECIALIZED | Patient has a documented history of chronic hepatitis or cirrhosis. |
| Disease | OI LOW LIZED | 1. Yes |
| 2.000.00 | | 2. No |
| | | 3. Unknown |
| History of Anemia | SPECIALIZED | Patient has a documented history of anemia. |
| | | 1. Yes |
| | | 2. No |
| | 00501411750 | 3. Unknown |
| History of Asthma | SPECIALIZED | Patient has a documented history of asthma/Reactive Airways disease. |
| | | 1. Yes 2. No |
| | | 3. Unknown |
| History of Thyroid | SPECIALIZED | Patient has a documented history of thyroid disorder. |
| Disorder | OI LOW LIZED | 1. Yes |
| | | If yes, specify |
| | | a. Hypothyroid |
| | | b. Hyperthyroid |
| | | c. Unknown |
| | | 2. No |
| | | 3. Unknown |

| History of | SPECIALIZED | Patient has a documented history of exposure to cardiotoxic chemotherapy. |
|---------------------------------|----------------|---|
| exposure to | 01 2011 (21222 | 1. Yes, |
| Cardiotoxic | | if yes, select Class: |
| Chemotherapy | | a. Anthracyclines: Adriamycin, Daunorubicin, Doxorubicin, |
| | | Epirubicin, Idarubicin, etc. |
| | | b. Mitoxantrone |
| | | c. Cyclophosphamide |
| | | d. Mitomycin C |
| | | e. Trastuzamab (Herceptin) |
| | | f. Tyrosine kinase inhibitor |
| | | g. Other, specify |
| | | h. Unknown |
| | | 2. No 3. Unknown |
| History of | SPECIALIZED | Unknown Patient has a documented history of thoracic radiation therapy. |
| Thoracic | SPECIALIZED | 1. Yes |
| Radiation | | If Yes, then specify: |
| Radiation | | a. Radiation therapy was received |
| | | i. Before 20 years of age |
| | | ii. After 20 years of age |
| | | b. Location |
| | | i. Mediastinal |
| | | ii. Chest |
| | | iii. Breast |
| | | iv. Other, specify |
| | | c. Total radiation dose (mCu) |
| | | 2. No |
| Illiata and Alaska la | ODECIALIZED | 3. Unknown |
| History of Alcohol Consumption/ | SPECIALIZED | Patient has a documented history of alcohol consumption/dependency. 1. Yes |
| Dependency | | If yes, categorize alcohol consumption history |
| Dependency | | a. Prior |
| | | b. Current |
| | | i. < 14 units* per week |
| | | ii. ≥14 units* per week |
| | | 2. No |
| | | 3. Unknown |
| | | |
| | | *Unit = 1 oz hard = 1 beer = 1 glass wine |
| History of | SPECIALIZED | Patient has a documented history of substance abuse. |
| Substance Abuse | | 1. Yes |
| | | If Yes, specify |
| | | a. Cocaine/crack b. Amphetamine |
| | | c. Opiates/heroin |
| | | d. Other, specify |
| | | 2. No |
| | | 3. Unknown |
| HIV Status | SPECIALIZED | Patient has a documented history of HIV seropositivity. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Unknown |

| History of | SPECIALIZED | Patient has a documented history of cancer, excluding non-melanoma basal |
|------------------------------|--------------|--|
| Malignancy | | cell skin cancers. |
| | | 1. Yes |
| | | If yes, specify cancer site. Select all that apply and indicate date of |
| | | first diagnosis, if available. |
| | | a. Breast |
| | | b. Cervical |
| | | c. Lung |
| | | d. Ovarian e. Prostate |
| | | f. Colon |
| | | g. Liver |
| | | h. Bone |
| | | i. Amyloidosis |
| | | j. Lymphoma/Leukemia |
| | | k. Other, specify |
| | | If yes, specify cancer therapies. Select all that apply. |
| | | a. Chemotherapy |
| | | b. Radiation Therapy |
| | | c. Surgery d. Transplantation |
| | | e. Other, specify |
| | | 2. No |
| | | 3. Unknown |
| History of Primary | SPECIALIZED | Patient has a documented history of primary muscular disease. |
| Muscular | | 1. Yes |
| Disease | | If yes, specify: |
| | | a. Muscular Dystrophy |
| | | b. Myasthenia Gravis |
| | | c. Dermatomyositis |
| | | d. Other, specify |
| | | 2. No 3. Unknown |
| History of | SPECIALIZED | Patient has a documented history of arthritis and/or collagen vascular |
| Arthritis or | OI LOW LIZED | disease. |
| Collagen | | 1. Yes |
| Vascular Disease | | If yes, specify all that apply: |
| | | a. Lupus Erythematosis |
| | | b. Scleroderma |
| | | c. Osteoarthritis |
| | | d. Gout |
| | | e. Rheumatoid Arthritis f. Seronegative arthropathy |
| | | g. Other, Specify |
| | | 2. No |
| | | 3. Unknown |
| History of | SPECIALIZED | Patient has a documented history of influenza immunization. |
| Influenza | | 1. Yes |
| Immunization | | If yes, Month and Year of most recent immunization should be noted. |
| | | 2. No |
| 111 4 | 00501115 | 3. Unknown |
| History of | SPECIALIZED | Patient has a documented history of pneumococcal immunization. |
| Pneumococcal Immunization | | 1. Yes |
| iniiiuiiization | | If yes, Month and Year of most recent immunization should be noted. 2. No |
| | | 3. Unknown |
| | | C. Cindomi |

B. CARDIOVASCULAR HISTORY

| FIELD NAME | CLASSIFICATION | DEFINITION |
|--|----------------|--|
| History of Arrhythmogenic Disease, Syndrome, or Substrate | SPECIALIZED | Patient has a documented history of <u>any</u> of the following arrhythmogenic conditions. Select all that apply. 1. None 2. Ventricular Tachycardia/fibrillation 3. Arrhythmogenic Right Ventricular Cardiomyopathy (ARVC) 4. Brugada syndrome 5. Wolf-Parkinson-White syndrome 6. Long QT syndrome 7. Hypertrophic cardiomyopathy (HCM) 8. Unknown |
| Family History of Sudden Cardiac Death | SPECIALIZED | Patient has a family history (parent, sibling or children) of sudden cardiac death, defined as natural death due to cardiac causes, within one hour of any new or changing cardiovascular symptom. The time and mode of death are unexpected even though pre-existing heart disease may have been known to be present. Sudden death without obvious cause is considered sudden cardiac death. Unwitnessed death without any evidence of new CV symptoms are considered SCD. 1. Yes 2. No 3. Unknown Age at time of sudden cardiac death may be specified. |
| History of Valvular Disease | SPECIALIZED | Patient has a documented history of primary valvular disease or history of valvular disease of other etiology. Select all that apply. 1. Yes If yes, may include: a. History of acute rheumatic fever/carditis (usually determined through correspondence with major and minor criteria) i. Yes ii. No iii. Unknown b. History of valve disease with echocardiographic findings suggestive of or diagnostic of rheumatic valvular disease. i. Yes ii. No iii. Unknown c. Congenital (present at birth or occurring association with congenital heart disease syndrome) d. Degenerative (acquired during adulthood, usually after age 50) e. Infectious (acquired as a result of infectious endocarditis) f. Toxic (for example, as a result of exposure to fenfluramine phentermine dexfenfluramine) g. Myxomatous h. Other (specify) i. Unknown 2. No 3. Unknown Valve affected and Year of the first episode may be helpful. |

| History of | ESSENTIAL | Patient has a documented history of congenital cardiac lesions. Also |
|--------------------|-----------|--|
| Congenital Cardiac | | indicate if surgery was performed on lesion(s). Select all that apply. |
| Lesions | | 1. Yes |
| | | If yes, specify each type and if surgery was performed: |
| | | a. Cyanotic |
| | | i. Congenitally Corrected Transposition |
| | | ii. Ebstein's Anomaly |
| | | iii. Hypoplastic Left Heart |
| | | iv. Pulmonary Atresia |
| | | v. Single Ventricle |
| | | vi. Tetralogy of Fallot |
| | | vii. Total Anomalous Pulmonary Venous Return |
| | | viii. Transposition of Great Vessels |
| | | ix. Tricuspid Atresia |
| | | x. Truncus Arteriosus |
| | | xi. Other, specify |
| | | b. Non-cyanotic |
| | | i. Aortic Stenosis |
| | | ii. Atrial Septic Defect (ASD) |
| | | iii. Atrioventricular Canal (endocardial cushion defect) |
| | | iv. Coarctaction of the Aorta |
| | | v. Patent Ductus Arteriosus (PDA) |
| | | vi. Pulmonic Stenosis |
| | | vii. Ventricular Septal Defect (VSD) |
| | | viii. Other, specify |
| | | 2. No |
| | | 3. Unknown |

C. HEART FAILURE ETIOLOGY

| FIELD NAME | CLASSIFICATION | DEFINITION |
|-------------------|----------------|--|
| Etiology of Heart | SPECIALIZED | Etiology of heart failure. Select all that apply and identify the primary and/or |
| Failure | | secondary etiology: |
| | | 1. Ischemic/CAD |
| | | 2. Valvular |
| | | 3. Infiltrative - Amyloid |
| | | 4. Infiltrative – Sarcoid |
| | | 5. Iron overload |
| | | 6. Myocarditis |
| | | 7. Hypertrophic |
| | | 8. Hypertensive |
| | | 9. Idiopathic |
| | | 10. Tachyarrhythmia Induced |
| | | 11. Familial |
| | | 12. ARVC |
| | | 13. Substance Abuse |
| | | 14. Alcoholic |
| | | 15. Pregnancy |
| | | 16. Chemotherapy |
| | | 17. Other(s), specify |

PART 3 – SYMPTOMS AND FUNCTIONAL ASSESSMENT

INDICATE ATE ATA COLLECTED (YYYYMMDD)

NOTE: The following data elements and definitions are specific to this Chapter and should be considered as a supplement to Symptoms section defined in the Core Elements Data Dictionary Chapter.

| FIELD NAME | CLASSIFICATION | DEFINITION |
|---------------------------------|----------------|--|
| Dyspnea at Rest | SPECIALIZED | Indicate if the patient describes frequent uncomfortable awareness of breathing while resting in a sitting position. 1. Yes 2. No Date of onset and duration may be helpful. |
| Dyspnea on Exertion | SPECIALIZED | Indicate if the patient describes uncomfortable awareness of breathing while exerting him/herself. 1. Yes If yes, Indicate degree of activity required to elicit dyspnea symptom a. Running or other sport (specify sport) b. Walking up an incline (specify distance) c. Walking on a flat surface (specify distance) d. Stopping to rest while dressing e. Standing (specify length of time) f. Other activity (i.e. shopping or housework), specify 2. No Date of onset and duration may be helpful. |
| Orthopnea | SPECIALIZED | Indicate if the patient describes at least one of the following: Uncomfortable awareness of breathing while in a supine position Positioning with 3 or more pillows or in a chair or recliner to maintain comfortable breathing during sleep Recurrent supine cough without other known cause may be an orthopnea equivalent None of the above Date of onset and duration may be helpful. |
| Paroxysmal Nocturnal Dyspnea | SPECIALIZED | Indicate if the patient describes awakening suddenly from sleep with uncomfortable awareness of breathing, or with general distress relieved by the upright position. Any report of this symptom lasting greater than 5 minutes is considered positive. 1. Yes 2. No Date of onset and duration may be helpful. |
| Swelling | SPECIALIZED | Indicate if the patient reports swelling or puffiness in extremities, bloating in abdomen, and/or other areas. 1. Yes 2. No Date of onset and duration may be helpful. |
| Fatigue | SPECIALIZED | Indicate if the patient describes unusual tiredness and inability to perform usual activities. 1. Yes 2. No Date of onset and duration may be helpful. |

| Syncope | SPECIALIZED | Indicate if the patient describes sudden loss of consciousness not related to anesthesia, with spontaneous recovery as reported by patient or observer. Patients losing consciousness prior to an implantable cardiac defibrillator (ICD) discharge will be considered to have syncope. 1. Yes 2. No Date of most recent episode may be helpful. |
|--------------------------------|-----------------------|--|
| FUNCTIONAL ASSESSME | NT AND STAGING OF DIS | EASE |
| NYHA Functional Capacity^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| ACC/AHA Heart Failure Stage | SPECIALIZED | Indicate the ACC/AHA Heart Failure Stage: A. Patient at high risk for developing heart failure but who has no structural disorder of the heart. B. Patient with a structural disorder of the heart but who has never developed symptoms of heart failure. C. Patient with past or current symptoms of heart failure associated with structural heart disease. D. Patient with end-stage disease who requires SPECIALIZED treatment strategies such as mechanical circulatory support, continuous inotropic infusions, cardiac transplantation or hospice care. |

PART 4 – PHYSICAL EXAM AND VITAL SIGNS FOR THIS EPISODE OF CARE

| FIELD NAME | CLASSIFICATION | DEFINITION |
|------------------------------|----------------|---|
| Heart Rate* | SPECIALIZED | Indicate the patients first recorded Heart Rate (in beats per minute) reading. Note: if zero, take the first obtainable reading. |
| | | *This data element and definition originates from the Acute Coronary Syndrome (ACS) Data Dictionary and is duplicated here for consistency. |
| Systolic Blood Pressure* | ESSENTIAL | Indicate the patients first recorded Systolic Blood Pressure (mm Hg) reading. Note: If zero, take the first obtainable reading. Note which arm is used |
| | | *This data element and definition originates from the Acute Coronary Syndrome (ACS) Data Dictionary and is duplicated here for consistency. |
| Diastolic Blood Pressure* | SPECIALIZED | Indicate the patients first recorded Diastolic Blood Pressure (mm Hg) reading. Note: If zero, take the first obtainable reading. Note which arm is used |
| | | *This data element and definition originates from the Acute Coronary Syndrome (ACS) Data Dictionary and is duplicated here for consistency. |
| Jugular Venous Pressure | SPECIALIZED | Record the height above sternal angle (cm). |
| Hepatojugular Reflux | SPECIALIZED | Indicate the presence or absence of Hepatojugular Reflux. 1. Positive 2. Negative |
| Respiratory Rate | SPECIALIZED | Indicate the patient's first recorded Respiratory Rate in respiratory cycles per minute. |
| Third Heart Sound | SPECIALIZED | Indicate the presence or absence of Third Heart Sound. 1. Present 2. Absent |
| Fourth Heart Sound | SPECIALIZED | Indicate the presence or absence of Fourth Heart Sound. 1. Present 2. Absent |
| Lung Examination | SPECIALIZED | Indicate the findings in the Lung Examination. Select all that apply. 1. Normal 2. Rales 3. Wheezing 4. Decreased breath sounds, or dullness to percussion |
| Peripheral Edema | SPECIALIZED | Indicate the presence or absence of Peripheral Edema. 1. Present 2. Absent |
| Ascites | SPECIALIZED | Indicate the presence or absence of Ascites. 1. Present 2. Absent |
| Hepatomegaly | SPECIALIZED | Indicate the presence or absence of Hepatomegaly. 1. Present 2. Absent |

PART 5 - MEDICATIONS

NOTE: The following data elements and definitions are specific to this Chapter and should be considered as a supplement to the Medications section defined in the Core Elements Data Dictionary Chapter.

A. PRE-ENCOUNTER

| FIELD NAME | CLASSIFICATION | DEFINITION |
|--------------------------|----------------|--|
| Aspirin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Clopidogrel [^] | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Prasugrel [^] | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Ticagrelor^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Digoxin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Anti-arrhythmics^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Carvedilol | SPECIALIZED | Indicate if the patient has been taking Carvedilol routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Bisoprolol | SPECIALIZED | Indicate if the patient has been taking Bisoprolol routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | 00501411750 | 5. Not Tolerated |
| Metoprolol | SPECIALIZED | Indicate if the patient has been taking Metroprolol routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Other Beta-blocker | SPECIALIZED | Indicate if the patient has been taking any other Beta-blocker(s) routinely |
| Other Beta-blocker | SPECIALIZED | prior to this encounter. |
| | | 1. Yes, specify drug(s) |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| ACE Inhibitors | ESSENTIAL | Indicate if the patient has been taking ACE Inhibitors routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |

| Angiotensin II | ESSENTIAL | Indicate if the patient has been taking Angiotensin II Receptor Blockers |
|--------------------|---------------|--|
| Receptor Blockers | LOOLIVII/(L | routinely prior to this encounter. |
| Treespier Diseases | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Warfarin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Dabigatran | SPECIALIZED | Indicate if the patient has been taking Dabigatran routinely prior to this |
| | OI LOW LEIZED | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Rivaroxaban | SPECIALIZED | Indicate if the patient has been taking Rivaroxaban routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Apixaban | SPECIALIZED | Indicate if the patient has been taking Apixaban routinely prior to this |
| • | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Loop Diuretics | SPECIALIZED | Indicate if the patient has been taking Loop Diuretics routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Metolazone | SPECIALIZED | Indicate if the patient has been taking Metolazone routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| Non Motologo | ODEOLAL IZED | 5. Not Tolerated |
| Non-Metolazone | SPECIALIZED | Indicate if the patient has been taking Non-Metolazone Thiazide Diuretics |
| Thiazide Diuretics | | routinely prior to this encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |

| Mineralocorticoid Receptor Antagonists | SPECIALIZED | Indicate if the patient has been taking Mineralocorticoid Receptor Antagonists (Spironolactone or Eplerenone) routinely prior to this encounter. 1. Yes, specify a. Spironolactone b. Eplerenone c. Other, specify 2. No 3. Contraindicated 4. Blinded 5. Not tolerated |
|---|----------------------------|--|
| Other Diuretic Agents | SPECIALIZED | Indicate if the patient has been taking any other Diuretic Agents routinely prior to this encounter. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Direct Renin | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Inhibitors^ | CDECIALIZED | ADefer to Core Flamente chanter for definition |
| Statins^ Other Lipid Lowering | SPECIALIZED SPECIALIZED | ^Refer to Core Elements chapter for definition. ^Refer to Core Elements chapter for definition. |
| Agents^ | OF LUIALIZED | Refer to Oure Elements chapter for definition. |
| Insulin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Oral | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Antihyperglycemics [^] | | |
| Non-Insulin Injectables^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Dihydropyridine Calcium Channel Blockers^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Non-Dihydropyridine Calcium Channel Blockers^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Ivabradine | SPECIALIZED | Indicate if the patient has been taking Ivabradine routinely prior to this encounter. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Supplemental Potassium or Magnesium | SPECIALIZED | Indicate if the patient has been taking any supplemental Potassium or Magnesium routinely prior to this encounter. 1. Yes, specify a. Potassium b. Magnesium 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Intravenous (IV) Inotropic Agent | SPECIALIZED | Indicate if the patient has had intravenous Positive Inotrope administered prior to this encounter. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |

| Intravenous (IV) | SPECIALIZED | Indicate if the patient has had intravenous Nesiritide administered prior to |
|---------------------|-------------|--|
| Nesiritide | | this encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Intravenous (IV) | SPECIALIZED | Indicate if the patient has had intravenous Vasodilator administered prior |
| Vasodilator Agents | | to this encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Oral or Intravenous | SPECIALIZED | Indicate if the patient has had oral or intravenous Vasopressin Antagonist |
| (IV) Vasopressin | | administered prior to this encounter. |
| Antagonist | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Morphine Sulfate | SPECIALIZED | Indicate if patient has been taking Morphine Sulfate routinely (orally or |
| | | intravenously) prior to this encounter. May be administered for pain or |
| | | pulmonary edema. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded 5. Not Tolerated |
| Hydromorphone | SPECIALIZED | Indicate if patient has been taking Hydromorphone routinely prior to this |
| Tiyaromorphone | SFLOIALIZED | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Fentanyl | SPECIALIZED | Indicate if patient has been taking Fentanyl routinely prior to this |
| | | encounter. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Nitrate Therapy | SPECIALIZED | Indicate if patient has been using Nitrate Therapy routinely prior to this |
| | | encounter. Nitroglycerin may be in the form of topical, oral, or sublingual. |
| | | Nitroglycerin spray or pills used on an as-needed basis only should be |
| | | noted in this category as sublingual. |
| | | 1. Yes |
| | | a. Topical |
| | | b. Oral |
| | | c. Sublingual 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| | | J. NOT TOICIALED |

| Hydralazine SPECIALIZED Indicate if the patient has been taking Hydralazine routinely encounter. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated | |
|---|---------------------|
| 1. Yes 2. No 3. Contraindicated 4. Blinded | |
| 2. No 3. Contraindicated 4. Blinded | |
| 3. Contraindicated 4. Blinded | |
| 4. Blinded | |
| | |
| | |
| | hrania waa |
| Oxygen Therapy SPECIALIZED Indicate if the patient has been using Oxygen Therapy for contribution to this appropriate this appropriate. | inronic use |
| rountinely prior to this encounter. | |
| 1. Yes | |
| 2. No | |
| 3. Contraindicated | |
| 4. Blinded | |
| 5. Not Tolerated | |
| Female Hormone SPECIALIZED Indicate if the patient has been using Female Hormone Rep | olacement |
| Replacement Therapy routinely prior to this encounter. | |
| Therapy 1. Yes | |
| 2. No | |
| 3. Contraindicated | |
| 4. Blinded | |
| 5. Not Tolerated | |
| Male Hormone SPECIALIZED Indicate if the patient has been using Male Hormone Repla | cement |
| Replacement Therapy routinely prior to this encounter. | |
| Therapy 1. Yes | |
| 2. No | |
| 3. Contraindicated | |
| 4. Blinded | |
| | |
| 5. Not Tolerated | latar varitia ali i |
| Inhaled SPECIALIZED Indicate if the patient has been using an Inhaled Bronchodi | lator routinely |
| Bronchodilator prior to this encounter. | |
| 1. Yes | |
| 2. No | |
| 3. Contraindicated | |
| 4. Blinded | |
| 5. Not Tolerated | |
| Inhaled SPECIALIZED Indicate if the patient has been using an Inhaled Corticoste | roid routinely |
| Corticosteroid prior to this encounter. | |
| 1. Yes | |
| 2. No | |
| 3. Contraindicated | |
| 4. Blinded | |
| 5. Not Tolerated | |
| Non-Steroidal Anti- SPECIALIZED Indicate if the patient has been taking any Non-Steroidal Ar | nti- |
| Inflammatory Drug Inflammatory Drugs routinely prior to this encounter. As NS | |
| (NSAID) generally contraindicated in patients with heart failure, spec | |
| for their use should be noted. | |
| 1. Yes | |
| 2. No | |
| 3. Contraindicated | |
| 4. Blinded | |
| | |
| 5. Not Tolerated | nto |
| Vitamins, Food SPECIALIZED Indicate if the patient is taking any vitamins, food suppleme | riis, |
| Supplements, and homeopathic treatments routinely prior to this encounter. | |
| Other Non- | |
| Prescription If yes, please specify | |
| Treatments 2. No | |

B. DURING HEALTHCARE ENCOUNTER

| Aspirin^ SPECIALIZED Refer to Core Elements chapter for definition. | FIELD NAME | CLASSIFICATION | DEFINITION |
|--|--------------------|----------------|--|
| SPECIALIZED Refer to Core Elements chapter for definition. | Aspirin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Prasugre ^ SPECIALIZED Refer to Core Elements chapter for definition. | | SPECIALIZED | |
| PECIALIZED Refer to Core Elements chapter for definition. | | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Digoxin* SPECIALIZED Nefer to Core Elements chapter for definition. | | SPECIALIZED | |
| Anti-arrhythmics | | SPECIALIZED | |
| SPECIALIZED Indicate if Carvedilol was administered at any point in time during this episode of health care. | | | |
| 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 6. Not Tolerated 7. Not Tolerated 7. Yes 7. Ye | | SPECIALIZED | Indicate if Carvedilol was administered at any point in time during this |
| 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 6. Not Tolerated 6. Not Tolerated 7. Not | | | episode of health care. |
| Bisoprolol SPECIALIZED Indicate if Bisoprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 6. Not Tolerated 7. No | | | |
| Bisoprolol SPECIALIZED SPECIALIZED Indicate if Bisoprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Metoprolol SPECIALIZED Indicate if Metroprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Other Beta-blocker SPECIALIZED Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Applications in II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 5. Not Tolerated 6. Not Tolerated 7. Not Tolerated 7. Yes 7. No 7. Contraindicated 7. Not Tolerated 8. Blinded 8. Blinded 9. Not Tolerated 9. Not Toler | | | |
| SPECIALIZED Indicate if Bisoprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Metroprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Metroprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated | | | |
| SPECIALIZED Indicate if Bisoprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated SPECIALIZED Indicate if Metroprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated SPECIALIZED Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated SPECIALIZED Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Applications were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Applications in II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Applications in II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Applications in II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Applications in II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Applications in II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 1. Yes 2. No 3. Contraindicated | | | |
| episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Metroprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 1. Yes 1. Yes 1. Not Tolerated Warfarin^ SPECIALIZED Afer to Core Elements chapter for definition. Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 4. Blinded 5. Not Tolerated 5. Not Tolerated 6. SPECIALIZED Afer to Core Elements chapter for definition. Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| Netoprolol SPECIALIZED Indicate if Metroprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 6. Not Tolerated 7. N | Bisoprolol | SPECIALIZED | |
| ACE Inhibitors SSENTIAL SSENTIAL Contraindicated | | | |
| Metoprolol SPECIALIZED Indicate if Metroprolol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Other Beta-blocker Other Beta-blocker SPECIALIZED Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED Afeer to Core Elements chapter for definition. Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated | | | |
| Metoproiol SPECIALIZED Indicate if Metroproiol was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 5. Not Tolerated 6. N | | | |
| SPECIALIZED Indicate if Metroproloi was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin* | | | |
| SPECIALIZED | | | |
| episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Other Beta-blocker SPECIALIZED Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated | Metoprolol | SPECIALIZED | |
| 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 1. Yes 2. No 3. Contraindicated 4. Blinded 4. Blinded 5. Not Tolerated 1. Yes 2. No 3. Contraindicated 4. Blinded 4. Blinded 5. Not Tolerated 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 4. Blinded 5. Not Tolerated 6. Not Tolerated 7. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 7. N | Metoproloi | OI LOIALIZED | |
| Contraindicated A Blinded Blin | | | |
| 3. Contraindicated 4. Blinded 5. Not Tolerated | | | |
| A. Blinded S. Not Tolerated | | | |
| Indicate if any other Beta-blocker(s) were administered at any point in time during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Ves | | | |
| during this episode of health care. 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED Arefer to Core Elements chapter for definition. Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated | | | 5. Not Tolerated |
| 1. Yes, specify drug(s) 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin* SPECIALIZED ARefer to Core Elements chapter for definition. Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 5. Not Tolerated 6. Not Tolerated 7. Not Tolera | Other Beta-blocker | SPECIALIZED | Indicate if any other Beta-blocker(s) were administered at any point in time |
| ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | during this episode of health care. |
| ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated The provided in the point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated Arefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Arefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| ACE Inhibitors ESSENTIAL Indicate if ACE Inhibitors were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin* SPECIALIZED *Refer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated AREfer to Core Elements chapter for definition. SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | 1 |
| this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED Arefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | ACE Inhibitore | FOOTNIAL | |
| 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Uarfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 6. Not Tolerated 7. Not Tolerated 7. Not Tolerated 8. Not Tolerated 9. Not Tolerate | ACE inhibitors | ESSENTIAL | |
| 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | 1 |
| 3. Contraindicated 4. Blinded 5. Not Tolerated Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| Angiotensin II Receptor Blockers ESSENTIAL Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| Angiotensin II Receptor Blockers Indicate if Angiotensin II Receptor Blockers were administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated 6. Not Tolera | | | |
| Point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ^Refer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | Angiotensin II | ESSENTIAL | |
| 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ^Refer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| 3. Contraindicated 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ^Refer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| 4. Blinded 5. Not Tolerated Warfarin^ SPECIALIZED ^Refer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | 2. No |
| Warfarin^ SPECIALIZED ARefer to Core Elements chapter for definition. Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| Warfarin^SPECIALIZED^Refer to Core Elements chapter for definition.DabigatranSPECIALIZEDIndicate if Dabigatran was administered at any point in time during this episode of health care.1. Yes2. No3. Contraindicated | | | |
| Dabigatran SPECIALIZED Indicate if Dabigatran was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated | 10/ C : A | 00501411355 | |
| episode of health care. 1. Yes 2. No 3. Contraindicated | | | |
| 1. Yes 2. No 3. Contraindicated | Dabigatran | SPECIALIZED | |
| 2. No 3. Contraindicated | | | |
| 3. Contraindicated | | | |
| | | | |
| | | | 4. Blinded |
| 5. Not Tolerated | | | |

| Rivaroxaban | SPECIALIZED | Indicate if Rivaroxaban was administered at any point in time during this |
|----------------------|---|---|
| | | episode of health care. 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Apixaban | SPECIALIZED | Indicate if Apixaban was administered at any point in time during this |
| • | | episode of health care. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| Loop Diuretics | SPECIALIZED | 5. Not Tolerated Indicate if Loop Diuretics were administered at any point in time during |
| Loop Didietics | SPECIALIZED | this episode of health care. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Metolazone | SPECIALIZED | Indicate if Metolazone was administered at any point in time during this |
| | | episode of health care. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated 4. Blinded |
| | | 5. Not Tolerated |
| Non-Metolazone | SPECIALIZED | Indicate if Non-Metolazone Thiazide Diuretics were administered at any |
| Thiazide Diuretics | | point in time during this episode of health care. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| Mineralocorticoid | SPECIALIZED | Not Tolerated Indicate if Mineralocorticoid Receptor Antagonists (Spironolactone or |
| Receptor | SPECIALIZED | Eplerenone) were administered at any point in time during this episode of |
| Antagonists | | health care. |
| , intagonioto | | 1. Yes, specify |
| | | a. Spironolactone |
| | | b. Eplerenone |
| | | c. Other, specify |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded 5. Not tolorated |
| Other Diuretic | SPECIALIZED | 5. Not tolerated Indicate if any other Diuretic Agents were administered at any point in time |
| Agents | OI LOIALIZED | during this episode of health care. |
| J | | 1. Yes, specify drug(s) |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | 000000000000000000000000000000000000000 | 5. Not Tolerated |
| Direct Renin | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Inhibitors^ Statins^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Other Lipid Lowering | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Agents [^] | OI LOIALIZED | Note: to object terrients onapter for definition. |
| Insulin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| - * | | 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 |

| Oral Antihyperglycemics^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
|--|-------------|--|
| Non-Insulin Injectables^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Dihydropyridine Calcium Channel Blockers^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Non-Dihydropyridine Calcium Channel Blockers^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Ivabridine | SPECIALIZED | Indicate if Ivabridine was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Supplemental Potassium or Magnesium | SPECIALIZED | Indicate if Supplemental Potassium or Magnesium were administered at any point in time during this episode of health care. 1. Yes, specify a. Potassium b. Magnesium 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Intravenous (IV) Inotropic Agent | SPECIALIZED | Indicate if Positive Inotrope was intravenously administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Intravenous (IV) Nesiritide | SPECIALIZED | Indicate if Nesiritide was intravenously administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Intravenous (IV) Vasodilator Agents | SPECIALIZED | Indicate if Vasodilator Agents was intravenously administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Oral or Intraveneous (IV) Vasopressin Antagonist | SPECIALIZED | Indicate if Vasopressin Antagonist was orally or intravenously administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |

| Morphine Sulfate | SPECIALIZED | Indicate if Morphine Sulfate was administered at any point in time during this episode of health care. May be administered for pain or pulmonary edema. 1. Yes 2. No 3. Contraindicated 4. Blinded |
|--|-------------|--|
| Hydromorphone | SPECIALIZED | 5. Not Tolerated Indicate if Hydromorphone was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Fentanyl | SPECIALIZED | Indicate if Fentanyl was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Nitrate Therapy | SPECIALIZED | Indicate if Nitrate Therapy was administered at any point in time during this episode of health care. Nitroglycerin may be in the form of topical, oral, or sublingual. Nitroglycerin spray or pills used on an as-needed basis only should be noted in this category as sublingual. 1. Yes a. Topical b. Oral c. Sublingual 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Hydralazine | SPECIALIZED | Indicate if Hydralazine was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Oxygen Therapy | SPECIALIZED | Indicate if Oxygen Therapy for chronic use was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Female Hormone Replacement Therapy | SPECIALIZED | Indicate if Female Hormone Replacement Therapy was administered at any point in time during this episode of health care. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |

| Male Hormone | SPECIALIZED | Indicate if Male Hormone Replacement Therapy was administered at any |
|---------------------|-------------|--|
| Replacement | | point in time during this episode of health care. |
| Therapy | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Inhaled | SPECIALIZED | Indicate if an Inhaled Bronchodilator was administered at any point in time |
| Bronchodilator | | during this episode of health care. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Inhaled | SPECIALIZED | Indicate if an Inhaled Corticosteroid was administered at any point in time |
| Corticosteroid | | during this episode of health care. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Non-Steroidal Anti- | SPECIALIZED | Indicate if any Non-Steroidal Anti-Inflammatory Drugs were administered |
| Inflammatory Drug | | at any point in time during this episode of health care. As NSAIDs are |
| (NSAID) | | generally contraindicated in patients with heart failure, specific indications |
| | | for their use should be noted. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Vitamins, Food | SPECIALIZED | Indicate if any Vitamins, Food Supplements, Homeopathic Treatments |
| Supplements, and | | were administered at any point in time during this episode of health care. |
| Other Non- | | 1. Yes |
| Prescription | | If yes, please specify |
| Treatments | | 2. No |

C. AT DISCHARGE

| FIELD NAME | CLASSIFICATION | DEFINITION |
|--------------------------|----------------|---|
| Aspirin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Clopidogrel [^] | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Prasugrel [^] | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Ticagrelor^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Digoxin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Anti-arrhythmics^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Carvedilol | SPECIALIZED | Indicate if Carvedilol was continued or prescribed at discharge |
| | | Yes No Contraindicated Blinded Not Tolerated |
| Bisoprolol | SPECIALIZED | Indicate if Bisoprolol was continued or prescribed at discharge. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |

| Metoprolol | SPECIALIZED | Indicate if Metroprolol was continued or prescribed at discharge. |
|--------------------|--------------|---|
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Other Beta-blocker | SPECIALIZED | Indicate if any other Beta-blocker(s) were continued or prescribed at |
| | 0. 20 | discharge. |
| | | 1. Yes, specify drug(s) |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| ACE Inhibitors | ESSENTIAL | |
| ACE inhibitors | ESSENTIAL | Indicate if ACE Inhibitors was continued or prescribed at discharge. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | =005::=::: | 5. Not Tolerated |
| Angiotensin II | ESSENTIAL | Indicate if Angiotensin II Receptor Blockers were continued or prescribed |
| Receptor Blockers | | at discharge. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Warfarin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Dabigatran | SPECIALIZED | Indicate if Dabigatran was continued or prescribed at discharge |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Rivaroxaban | SPECIALIZED | Indicate if Rivaroxaban was continued or prescribed at discharge |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Apixaban | SPECIALIZED | Indicate if Apixaban was continued or prescribed at discharge. |
| , tpixubuii | OI LOIALIZED | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Loop Diurctica | SPECIALIZED | |
| Loop Diuretics | SPECIALIZED | Indicate if Loop Diuretics was continued or prescribed at discharge. |
| | | 1. Yes 2. No |
| | | |
| | | 3. Contraindicated |
| | | 4. Blinded |
| Matalaa | 00501411755 | 5. Not Tolerated |
| Metolazone | SPECIALIZED | Indicate if Metolazone was continued or prescribed at discharge. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |

| Non Motolozono | SPECIALIZED | Indicate if Non-Metolazone Thiazide Diuretics was continued or |
|-------------------------------------|-------------|---|
| Non-Metolazone | SPECIALIZED | |
| Thiazide Diuretics | | prescribed at discharge. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Mineralocorticoid | SPECIALIZED | Indicate Mineralocorticoid Receptor Antagonists (Spironolactone or |
| Receptor | | Eplerenone) were continued or prescribed at discharge. |
| Antagonists | | 1. Yes, specify |
| | | a. Spironolactone |
| | | b. Eplerenone |
| | | c. Other, specify |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not tolerated |
| Other Diuretic | SPECIALIZED | Indicate if any other Diuretic Agents were continued or prescribed at |
| Agents | • | discharge |
| | | 1. Yes, specify drug(s) |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Direct Renin | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Inhibitors^ | SFLOIALIZED | Refer to Gore Elements chapter for definition. |
| Statins^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Other Lipid Lowering | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Agents^ | SPECIALIZED | Refer to Core Elements chapter for definition. |
| Insulin^ | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Oral | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Antihyperglycemics^ | SFLOIALIZED | Refer to Core Liements chapter for definition. |
| Non-Insulin | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Injectables^ | SPECIALIZED | Refer to Core Elements chapter for definition. |
| Dihydropyridine | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Calcium Channel | SPECIALIZED | Refer to Core Elements chapter for delimition. |
| Blockers [^] | | |
| | SPECIALIZED | ^Refer to Core Elements chapter for definition. |
| Non-Dihydropyridine Calcium Channel | SPECIALIZED | Refer to Gore Elements chapter for definition. |
| Blockers [^] | | |
| Ivabradine | SPECIALIZED | Indicate if lyabrading was continued or properited at discharge |
| ivapradine | SPECIALIZED | Indicate if Ivabradine was continued or prescribed at discharge. |
| | | 1. Yes 2. No |
| | | |
| | | 3. Contraindicated |
| | | 4. Blinded |
| Cumplement 4-1 | ODECIALIZED | 5. Not Tolerated |
| Supplemental | SPECIALIZED | Indicate if supplemental Potassium or Magnesium was continued or |
| Potassium or | | prescribed at discharge. |
| Magnesium | | 1. Yes, specify |
| | | a. Potassium |
| | | b. Magnesium |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |

| Manushina Culfata | CDECIALIZED | Indicate if Marchine Culfete was continued as prescribed at discharge |
|--|-------------|---|
| Morphine Sulfate | SPECIALIZED | Indicate if Morphine Sulfate was continued or prescribed at discharge. May be administered for pain or pulmonary edema. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Hydromorphone | SPECIALIZED | Indicate if Hydromorphone was continued or prescribed at discharge. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Fentanyl | SPECIALIZED | Indicate if Fentanyl was continued or prescribed at discharge. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Nitrate Therapy | SPECIALIZED | Indicate if Nitrate Therapy was continued or prescribed at discharge. Nitroglycerin may be in the form of topical, oral, or sublingual. Nitroglycerin used on an as-needed basis only should be noted in this category. 1. Yes a. Topical b. Oral c. Sublingual 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Hydralazine | SPECIALIZED | Indicate if Hydralazine was continued or prescribed at discharge. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Oxygen Therapy | SPECIALIZED | Indicate if Oxygen Therapy for chronic use was continued or prescribed at discharge. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Female Hormone Replacement Therapy | SPECIALIZED | Indicate if Female Hormone Replacement Therapy was continued or prescribed at discharge. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |
| Male Hormone Replacement Therapy | SPECIALIZED | Indicate if Male Hormone Replacement Therapy was continued or prescribed at discharge. 1. Yes 2. No 3. Contraindicated 4. Blinded 5. Not Tolerated |

| Inhaled | SPECIALIZED | Indicate if an Inhaled Bronchodilator was continued or prescribed at |
|---------------------------------------|--------------|---|
| Bronchodilator | OI EOIAEIZED | discharge. |
| Bronchodilator | | 1. Yes |
| | | '' '' |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Inhaled | SPECIALIZED | Indicate if an Inhaled Corticosteroid was continued or prescribed at |
| Corticosteroid | | discharge. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | 4. Blinded |
| | | 5. Not Tolerated |
| Non-Steroidal Anti- | SPECIALIZED | Indicate if a Non-Steroidal Anti-Inflammatory Drug was continued or |
| Inflammatory Drug | | prescribed at discharge. As NSAIDs are generally contraindicated in |
| (NSAID) | | patients with heart failure, specific indications for their use should be |
| (1137112) | | noted. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Contraindicated |
| | | o. Contramidated |
| | | 4. Blinded |
| \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ | 00501411750 | 5. Not Tolerated |
| Vitamins, Food | SPECIALIZED | Indicate if any Vitamins, Food Supplements, Homeopathic Treatments |
| Supplements, and | | were continued or prescribed at discharge. |
| Other Non- | | 1. Yes |
| Prescription | | If yes, please specify |
| Treatments | | 2. No |

PART 6 - DEVICES

- INDICATE DATE DATA COLLECTED (YYYYMMDD)

A. IMPLANTABLE CARDIAC DEVICES (PACEMAKERS, DEFIBRILLATORS AND MONITORING DEVICES)

| FIELD NAME | CLASSIFICATION | DEFINITION |
|----------------|----------------|--|
| Device implant | SPECIALIZED | Indicate if the patient was implanted with an implantable cardiac device during this episode of health care. 1. Yes If yes, indicate device type and date of device implant (YYYYMMDD): a. S-ICD (single chamber Implantable defibrillator only), Date of implant b. D-ICD (dual chamber Implantable cardioverter defibrillator), Date of implant c. CRT (Biventricular pacemaker without defibrillator), Date of implant d. CRT-D (biventricular pacemaker with defibrillator, Date of implant e. Dual chamber Pacer (Dual chamber pacemaker), Date of implant f. Single chamber pacer (Single chamber pacemaker), Date of implant 2. No |
| Referral By | OPTIONAL | Specialty of Referral MD for this device implant. 1. Generalist 2. Internist 3. Cardiologist 4. Other, specify |
| Indication | SPECIALIZED | Primary Prevention Prophylaxis of sudden cardiac death in patient without previous event. If multiple indications, use lower numbered reason for primary and higher numbered reason for secondary indication. Secondary Prevention Prophylaxis of sudden cardiac death in patients with previously aborted sudden cardiac death due to VT or VF Syncope: History of or for the prevention of syncope Heart Failure Other, specify |

B. MECHANICAL CARDIAC ASSISTS DEVICES

| FIELD NAME | CLASSIFICATION | DEFINITION |
|------------------------|----------------|--|
| Device Implant | SPECIALIZED | Indicate if the patient was implanted with a device during this episode of health care. Select all that apply. 1. Yes If yes, indicated the device implanted and date of device implant (YYYYMMDD) a. LVAD, Date implanted b. RVAD, Date implanted c. BiVAD, Date implanted d. TAH, Date implanted e. ECMO, please specify: i. V-A ECMO, Date implanted ii. V-V ECMO, Date implanted f. other, specify and Date implanted 2. No |
| Device Strategy | SPECIALIZED | Indicate device strategy for this implantation. 1. Bridge-to-Transplant (patient listed) 2. Bridge-to-Recovery 3. Bridge-to-Decision, specify: |
| Device Brand (LVAD) | SPECIALIZED | If the patient was implanted with an LVAD, indicate the device brand. 1. Not implanted with an LVAD 2. HeartMate II 3. Heartware 4. Thoratec PVAD 5. Jarvik 2000 6. Berlin Heart, please specify: a. EXCOR b. INCOR 7. Arrow Lionheart 8. Micromed Debakey Heart 9. Biomedicus 10. Tandem Heart 11. Levitronix Centrimag 12. Abiomed Impella a. LD 2.5 b. LD 5.0 c. 8F 4.0 L percutaneous 13. Other, specify |

¹ This is the standard European format for date and also follows the CIHI Discharge Abstract Database (DAD) for dates. All data elements across all chapters should consistently use the YYYYMMDD format.

| Device Brand | SPECIALIZED | If the patient was implanted with an RVAD, Indicate the device brand: |
|----------------------|-------------|--|
| (RVAD) | SFLCIALIZED | Not implanted with an RVAD |
| (KVAD) | | 2. Thoratec PVAD |
| | | 3. Biomedicus |
| | | |
| | | 4. Levitronix Centrimag |
| | | 5. Abiomed Impella, please specify: |
| | | a. RD 2.5 |
| | | b. RD 5.0 |
| | | 6. Other, specify |
| Device Brand | SPECIALIZED | If the patient was implanted with an BiVAD, indicate the device brand: |
| (BiVAD) | | Not implanted with a BIVAD |
| | | 2. Thoratec, please specify: |
| | | a. IVAD |
| | | b. PVAD |
| | | 3. Other, specify |
| Device Brand | SPECIALIZED | If the patient was implanted with a TAH, indicate the device: |
| (TAH) | | SynCardia Cardiowest |
| | | 2. Other, specify |
| Is this VAD an | SPECIALIZED | If the patient was implanted with a VAD, indicate if it was an investigational |
| investigational | | device. |
| device? | | 1. Yes |
| | | 2. No |
| | | 3. Unknown |
| Is patient part of a | SPECIALIZED | Indicate if this patient is part of a device clinical trial. |
| device clinical | | 1. Yes |
| trial? | | 2. No |
| | | 3. Unknown |
| INTERMACS | SPECIALIZED | Indicate the corresponding INTERMACS patient profile at the time of this |
| Patient Profile | | device implant. |
| | | 1. INTERMACS 1 |
| | | 2. INTERMACS 2 |
| | | 3. INTERMACS 3 |
| | | 4. INTERMACS 4 |
| | | 5. INTERMACS 5 |
| | | 6. INTERMACS 6 |
| Final Insul | 00501411750 | 7. INTERMACS 7 |
| First Implant | SPECIALIZED | Indicate if the patient was implanted with a device prior to this episode of |
| | | health care. |
| | | 1. Yes |
| | | If yes, then indicate former implant(s): |
| | | a. LVAD |
| | | b. RVAD |
| | | c. BiVAD |
| | | d. TAH |
| | | e. ECMO |
| | | f. Other, specify |
| | | 2. No |
| | | 3. Unknown |

| Complications of | SPECIALIZED | Indicate if the patient suffered a complication as a result of the device implant |
|------------------|--------------|---|
| Device Implant | 0. 20 | (excluding LVAD therapy) during or requiring hospitalization. Indicate all that |
| (excluding LVAD | | apply. |
| Therapy) | | 1. Pneumothorax |
| Therapy) | | 2. Hemothorax |
| | | |
| | | 3. Infection |
| | | 4. Death |
| | | 5. Stroke |
| | | 6. Pericardial Tamponade |
| | | 7. Myocardial Infarction |
| | | 8. Cardiac Arrest |
| | | 9. Respiratory Failure |
| | | 10. Cardiogenic Shock |
| | | 11. Ventricular Arrhythmia (Requiring treatment) |
| | | 12. Coronary Sinus Dissection |
| | | 13. Atrial Pacer Lead Dislodgement |
| | | 14. RV Pacer Lead Dislodgement |
| | | 15. Defibrillator Lead Dislodgement (ventricle) |
| | | 16. Coronary Sinus Lead Dislodgement |
| | | 17. Diaphragmatic Pacing |
| | | 18. Other, specify |
| Complications of | SPECIALIZED | Indicate if the patient suffered a complication as a result of LVAD therapy |
| LVAD Therapy | OI LOW LIZED | during or requiring hospitalization. Indicate all that apply. |
| LVAD Illerapy | | 1. Death |
| | | 2. Tamponade |
| | | |
| | | 3. Low output syndrome |
| | | 4. RV failure |
| | | 5. MOSF |
| | | 6. Pneumonia |
| | | 7. Sepsis |
| | | 8. UTI |
| | | 9. PE |
| | | 10. Other infection |
| | | 11. Device thrombosis |
| | | 12. Vasogenic hypotension (post pump vasoplegia) |
| | | 13. Embolism |
| | | 14. Stroke |
| | | 15. TIA |
| | | 16. Driveline infection |
| | | 17. Pump dysfunction |
| | | 18. Device infection |
| | | 19. ARF |
| | | 20. Bleeding requiring transfusion |
| | | 21. Hemolysis |
| | | 22. Other, specify |

PART 7 – LABORATORY RESULTS

Note: The laboratory testing elements listed below are those reflecting the laboratory tests most likely to be followed longitudinally by cardiologists taking care of heart failure patients for the purpose of metabolic and cardiovascular risk mitigation via direct interventional management of the laboratory abnormalities.

These laboratory result data elements are useful adjunct to clinical evaluation, assessing the appropriate use of cardiovascular procedures (including LVAD implantation risk factors), prognosis (Seattle HF model) and for quality-performance measurement, and for meaningful use reporting.

| FIELD NAME | CLASSIFICATION | DEFINITION |
|-------------------------------|----------------|--|
| BNP | SPECIALIZED | Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) Not done |
| Nt-proBNP | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) 2. Not done |
| Troponin | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units d. Date of test (YYYYMMDD) 2. Not done |
| Total cholesterol (mmol/L) | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) 2. Not done |
| LDL cholesterol (mmol/L) | SPECIALIZED | Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) Not done |
| Uric Acid (umol/L) | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) 2. Not done |
| Albumin (mg/L) | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) 2. Not done |

| TOLL | ODEOLALIZED | 4 V |
|------------------|--------------|---|
| TSH | SPECIALIZED | 1. Yes |
| | | Indicate the following: a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) |
| | | 2. Not done |
| Creatinine | ESSENTIAL | 1. Yes |
| (mmol/L) | LOOLIVIIAL | Indicate the following: |
| (IIIIIOI/L) | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) |
| | | 2. Not done |
| BUN (mmol/L) | ESSENTIAL | 1. Yes |
| , , | | Indicate the following: |
| | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) |
| | | 2. Not done |
| | | BUN is defined as blood urea nitrogen. |
| Sodium (mmol/L) | ESSENTIAL | 1. Yes |
| | | Indicate the following: |
| | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) 2. Not done |
| Potassium | ESSENTIAL | 1. Yes |
| (mmol/L) | ESSENTIAL | Indicate the following: |
| (IIIIIOI/L) | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) |
| | | 2. Not done |
| Hemoglobin A1c | SPECIALIZED | 1. Yes |
| | | Indicate the following: |
| | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) |
| | | 2. Not done |
| Hemoglobin (g/L) | SPECIALIZED | 1. Yes |
| | | Indicate the following: |
| | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) 2. Not done |
| Hematocrit | SPECIALIZED | 1. Yes |
| | OI LOIALIZED | Indicate the following: |
| | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) |
| | | 2. Not done |
| WBC count | SPECIALIZED | 1. Yes |
| | | Indicate the following: |
| | | a. Value |
| | | b. Units |
| | | c. Date of test (YYYYMMDD) |
| | | 2. Not done |

| Lymphocytes (%) | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) 2. Not done |
|-----------------|-------------|--|
| Platelet (µL) | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) 2. Not done |
| INR | SPECIALIZED | 1. Yes Indicate the following: a. Value b. Units c. Date of test (YYYYMMDD) 2. Not done |

PART 8 – DIAGNOSTIC TESTS

| FIELD NAME | CLASSIFICATION | DEFINITION |
|-------------------------------|----------------|--|
| (LV) Function^ | ESSENTIAL | ^Refer to Core Elements Chapter for definition. |
| Ejection Fraction Modality | ESSENTIAL | Indicate the modality used to determine the ejection fraction. When multiple measures are available, the most recent is preferred. 1. No documentation of ejection fraction 2. Magnetic resonance imaging (MRI) 3. Echocardiography 4. LV Angiogram 5. CT Angiogram 6. Other nuclear imaging technique including Radionuclide ventriculography 7. Other Indicate the Date of Ejection Fraction (YYYYMMDD) |
| Electrocardiography | SPECIALIZED | Indicate the results of first 12-lead electrocardiography results. When a quantitative range is given, provide the midpoint of the range. Documented findings may include: 1. Rhythm: a. Sinus rhythm b. Atrial fibrillation or flutter c. Indicate if paroxysmal or persistent? d. Paced or other rhythm 2. Heart rate (beats per minute) 3. Left bundle branch block (LBBB) 4. Right bundle branch block (RBBB) 5. Non-specific intraventricular conduction delay 6. Location of abnormal Q waves (0.03 second in width and 1 mm [0.1 mV] in depth in at least 2 contiguous leads) 7. QRS duration (in milliseconds): may be reported as the measured duration, or categorically as shorter than 120 milliseconds; 121 to 150 milliseconds; or longer than 150 milliseconds 8. Heart block, indicate degree a. None b. 1st degree c. 2nd type 1 (Wenckebach) d. 2nd type 2 e. 3rd degree |
| Chest Radiography | ESSENTIAL | Indicate the Date of Electrocardiography (YYYYMMDD) Indicate the results of the first Radiological examination of the chest. Documented findings from the chest X-ray pertinent to heart failure patients may include: 1. Presence of Pulmonary vascular redistribution, pulmonary congestion, or pulmonary edema 2. Presence of Cardiomegaly 3. Presence of Pleural effusion(s) Indicate the Date and Time of Chest Radiology (YYYYMMDD) |

| ISCHEMIA AND INFARCT E | VALUATION | |
|------------------------|-------------|---|
| Myocardial Imaging | ESSENTIAL | Indicate the following |
| | | 1. Type of imaging: |
| | | a. Radionuclide (nuclear) imaging |
| | | i. Planar |
| | | ii. SPECT |
| | | iii. PET |
| | | b. MRI |
| | | c. Echocardiogram |
| | | d. CT |
| | | e. Coronary angiogram |
| | | f. Unknown |
| | | Stress-induced Perfusion defect: |
| | | a. Yes |
| | | b. No |
| | | c. Unknown |
| | | Fixed Perfusion defect: |
| | | a. Yes |
| | | b. No |
| | | c. Unknown |
| | | Ladicate the Date of Localine (AAAAAAADD) |
| Cananamulasiana | SPECIALIZED | Indicate the Date of Imaging (YYYYMMDD) |
| Coronary Lesions | SPECIALIZED | Indicate the extent of coronary lesions. |
| | | Normal (< 20% stenosis in all epicardial vessels) |
| | | 2. Obstructive (> 50% in one or more vessels) |
| | | a. Yes, if yes, select one: |
| | | i. Single vessel. Indicate if LM or Prox LAD |
| | | involvement |
| | | 1. Neither |
| | | 2. If Yes, specify |
| | | a. >50% LM |
| | | b. >50% Prox LAD |
| | | ii. Two vessel. Indicate if LM or Prox LAD |
| | | involvement |
| | | 1. Neither |
| | | 2. If Yes, (Select all that apply) |
| | | a. >50% LM |
| | | b. >50% Prox LAD |
| | | iii. Three vessel. Indicate if LM or Prox LAD |
| | | involvement |
| | | 1. Neither |
| | | 2. Yes, (Select all that apply) |
| | | a. >50% LM |
| | | b. >50% Prox LAD |
| | | b. No |
| | | Note: This data definition is duplicated here from the Atrial Fibrillation Data |
| | | Dictionary Chapter and is consistent with the originating definition from the |
| | | |
| | | Coronary Angiography/Revascularization Data Dictionary Chapter. |

| Right Heart | SPECIALIZED | Indicate the results of a Right heart catheterization with or without |
|-----------------|--------------|---|
| Catheterization | OI LOIALIZED | |
| Catheterization | | pulmonary angiography. Documented findings may include: |
| | | No right heart catheterization documented |
| | | 2. RA pressure (mm Hg): mean right atrial pressure from pulmonary |
| | | artery catheter. Indicate number and reference range. |
| | | PA systolic pressure (mm Hg): systolic pulmonary pressure from |
| | | pulmonary artery catheter. Indicate number and reference range. |
| | | 4. PA diastolic pressure (mm Hg): diastolic pulmonary pressure from |
| | | pulmonary artery catheter. Indicate number and reference range. |
| | | 5. Mean pulmonary artery occlusion pressure from pulmonary artery |
| | | catheter (wedge pressure, mm Hg). May be recorded with or |
| | | without V-wave. Indicate number and reference range. |
| | | 6. Cardiac output/index (liters per minute). Indicate number and |
| | | reference range. |
| | | 7. Pulmonary vascular resistance (Wood's units). Indicate number |
| | | and reference range. |
| | | Systemic vascular resistance (dynes/second/cm2). Indicate |
| | | number and reference range. |
| | | |
| | | Indicate the Date of Catherization (YYYYMMDD) |
| Heart Biopsy | SPECIALIZED | Biopsy of the endomyocardium. |
| | | 1. Yes |
| | | 2. No |
| | | 3. Unknown |
| | | |
| | | Indicate the Date of Biopsy (YYYYMMDD) |

| Stress Testing | SPECIALIZED | Cardiovascular stress test including exercise (treadmill, bicycle) and |
|----------------|------------------|--|
| 23.000 .0009 | 0. 20.7 (2.12.25 | pharmacological stress. Documented findings may include: |
| | | Maximal (symptom limited) or submaximal test |
| | | Workload achieved. May be expressed as Watts, exercise stage |
| | | achieved (include exercise protocol) or metabolic equivalents |
| | | (METS). |
| | | · · · · · · |
| | | Reason for terminating exercise test: Cheet pain. |
| | | a. Chest pain |
| | | b. Dyspnea |
| | | c. Dizziness |
| | | d. Leg fatigue |
| | | e. Other discomfort (specify) |
| | | f. Tachyarrhythmias |
| | | g. Asymptomatic tachyarrhythmias |
| | | 4. Evidence for ischemia on stress test (positive or negative): |
| | | a. Positive: on an exercise tolerance test, the patient developed either |
| | | i. Greater than or equal to 1 mm of horizontal or |
| | | downsloping ST-segment depression or elevation |
| | | for at least 60 to 80 milliseconds (ms) after the |
| | | end of the QRS complex, in asymptomatic |
| | | patients, or ii. New ST-segment depression greater than or |
| | | equal to 2 mm (0.2 mV) (horizontal or |
| | | downsloping) believed to represent ischemia |
| | | even in the absence of ischemic discomfort. If the |
| | | patient had an equivalent type of exercise test |
| | | (e.g., exercise thallium or MIBI test, stress |
| | | echocardiography, or dipyridamole, thallium, or |
| | | adenosine radioisotope scan) that showed |
| | | definite evidence of ischemia (e.g., an area of clear reversible ischemia), this should be |
| | | considered a positive test. |
| | | b. Negative: no evidence of ischemia (i.e., no typical angina pain |
| | | and no ST depressions, no imaging evidence for ischemia) |
| | | 5. 6-minute walk test. Record the distanced walked during 6-minute |
| | | walk (on a flat surface), in meters. |
| | | Indicate the Date of Stress Test (YYYYMMDD) |
| CPET | SPECIALIZED | Record the following. |
| | | Exercise time Mayimum wantland ashiound |
| | | 2. Maximum workload achieved 3. VO2 (ml/kg/min: L/min) |
| | | 3. VO2 (ml/kg/min; L/min) 4. % predicted |
| | | 5. VE/VCO2 |
| | | 6. RER |
| | | Indicate Date of CPET (YYYYMMDD) |
| | l | maiodic bate of of ET (TTTTWINDD) |

PART 9 - COUNSELLING

- INDICATE DATE DATA COLLECTED (YYYYMMDD)

| FIELD NAME | CLASSIFICATION | DEFINITION | |
|--------------------|---|---|--|
| FIELD NAME | CLASSIFICATION | DEFINITION | |
| ADVANCE DIRECTIVES | E/END OF LIFE MANAGE | MENT/PROGNOSIS | |
| Do Not | SPECIALIZED | Indicate if there is any explicit documentation by physician and/or patient | |
| Resuscitate | | indicating that no resuscitative efforts are to be performed in the event of | |
| (DNR) | | circulatory or respiratory arrest. | |
| | | 1. Yes | |
| | | 2. No | |
| Advance Care | SPECIALIZED | Indicate if there is any documentation of discussion carried out with the | |
| Planning | | patient and/or family (by physician or nurse) about advance directive. | |
| | | 1. Yes | |
| | | 2. No | |
| EDUCATION ON HEAR | EDUCATION ON HEART FAILURE AND SELF-MANAGEMENT SKILLS - | | |
| PATIENT EDUCATION: | ASSESSMENT OF LEAR | NING READINESS | |
| Education Level^ | SPECIALIZED | ^Refer to Core Elements Chapter for definition. | |
| | | | |
| EDUCATION ON HEAR | T FAILURE AND SELF-MA | ANAGEMENT SKILLS - | |
| EDUCATION/COUNSEL | LING INTERVENTION (No | OTE: SHOULD BE THE SAME FOR INPATIENT AND OUTPATIENT COUNSELLING) | |
| | T | | |
| Referral to | SPECIALIZED | Indicate if there is any documentation in the medical record that the patient | |
| Dietician for Diet | | was referred to Dietician for weight management and/or advanced nutritional | |
| Counselling | | instruction. | |
| | | 1. Yes | |
| 5.4.4 | 00501411755 | 2. No | |
| Referral to | SPECIALIZED | Indicate if there is any documentation in the medical record that the patient | |
| Cardiac | | was referred to cardiac rehabilitation or other structured exercise program. | |
| Rehabilitation | | 1. Yes | |
| Program | | 2. No | |

PART 10 - CARE EPISODE

- INDICATE DATE DATA COLLECTED (YYYYMMDD)

A. DEMOGRAPHICS

| FIELD NAME | CLASSIFICATION | DEFINITION |
|---|----------------|--|
| Care Period | SPECIALIZED | For inpatient admission, note the date the patient was admitted to the hospital and the date the patient was discharged from the hospital. For outpatient, note the date (day, month, year) of the encounter (physician visit, nurse visit, consultation, procedures, and so on). For emergency department visit without hospital admission, note the date |
| Presentation to Health Care Facility | SPECIALIZED | (day, month, year) of the encounter (physician visit, nurse visit, consultation, procedures, and so on). Indicate the type of health care contact for this episode of care. 1. Emergency department visit only without hospital admission 2. Hospital admission for HF 3. other cardiovascular problem 4. non-cardiovascular problem (e.g. pneumonia) 5. Planned admission 6. Regularly scheduled outpatient visit 7. urgent outpatient visits 8. Remote monitoring 9. Telephone contact 10. Electronic communication (e.g. email, messaging) 11. Other, specify |
| Location of Health Care Encounter | SPECIALIZED | Indicate the type of location of health care encounter. 1. Acute-care hospital 2. Long-term care facility 3. Emergency department 4. Caregiver office 5. Heart failure clinic a. Cardiology practice b. Primary care physician office c. Other caregiver office 6. Other, specify |
| Initial Location Type | SPECIALIZED | Indicate the type of facility where the patient was first evaluated at. 1. Academic Teaching Hospital 2. Community Hospital |
| Date-Time of First Medical Contact* | SPECIALIZED | Indicate the date and time of first medical contact. Note: First medical contact is defined as arrival of paramedic to patient or arrival at emergency if patient self-transports to the emergency department. 1. Date and Time values 2. Not Available *Note: this data element originates in the ACS Chapter and is duplicated here for consistency in definitions. |
| Arrival Date-Time At First Facility* | SPECIALIZED | Indicate the date and time the patient arrived at first facility. 1. Date and Time values 2. Not available *Note: this data element originates in the ACS Chapter and is duplicated here for consistency in definitions. |

| Admission Date- Time* | SPECIALIZED | If admitted, indicate the date and time the patient was admitted as an inpatient to first facility for the current episode of care. 1. Date and Time values 2. Not available *Note: this data element originates in the ACS Chapter and is duplicated here for consistency in definitions. |
|--------------------------------|-------------|--|
| Initial Means of Transport* | SPECIALIZED | Indicate the means of transportation to the facility where the patient was evaluated first. Select code: 1. Ambulance 2. Air 3. Other 4. Not Available *Note: this data element originates in the ACS Chapter and is duplicated here for consistency in definitions. |

B. INTER-HOSPITAL TRANSFER

| FIELD NAME | CLASSIFICATION | DEFINITION |
|---|----------------|--|
| Inter-Hospital Transfer?* | SPECIALIZED | Was the patient transferred to another hospital after assessment at first facility (regardless of whether treatment was received at first hospital or not)? 1. Yes 2. No *Note: this data element originates in the ACS Chapter and is duplicated here for consistency in definitions. |
| Inter-Hospital Transfer - Means of Transport* | SPECIALIZED | Indicate the means of inter-hospital transfer. Select code: 1. Ambulance 2. Air 3. Other 4. Not Available *Note: this data element originates in the ACS Chapter and is duplicated here for consistency in definitions. |
| Date-Time of Patient Transfer between Facilities* | SPECIALIZED | Indicate the date and time the patient left the initial hospital. 1. Date and Time values 2. Not available *Note: this data element originates in the ACS Chapter and is duplicated here for consistency in definitions. |

C. Invasive Therapeutic Procedures

| FIELD NAME | CLASSIFICATION | DEFINITION |
|---|----------------|--|
| Coronary Artery Bypass Graft (CABG) Surgery | SPECIALIZED | Indicate the number and types of grafts and surgical approach may be further specified: 1. None 2. Number and placement of vein bypass grafts 3. Number and placement of arterial bypass grafts 4. Standard bypass surgery approach |
| | | Approach may be further specified: 1. Median Sternotomy Approach 2. Small Thoracotomy 3. With or without cardiopulmonary bypass |

| Valve Repair | SPECIALIZED | Indicate the type(s) of repair. Valve(s) and procedure(s) may be specified. 1. None 2. Mitral 3. Aortic, and/or 4. Tricuspid valve surgical repair. 5. Percutaneous Mitral repair (Mitraclip) |
|--|-------------|---|
| Valve Replacement | SPECIALIZED | Indicate the type of replacement. Valve(s) and procedure(s) may be specified. 1. None 2. Mitral 3. Aortic 4. Tricuspid |
| Valvuloplasty | SPECIALIZED | Indicate if the patient required valvuloplasty for stenotic valve lesions. Valve(s) and procedure(s) may be specified. 1. None 2. Mitral 3. Aortic 4. Tricuspid |
| Ventricular Remodeling Surgery | SPECIALIZED | Indicate if the patient required ventricular remodeling surgery. 1. None 2. Aneurysectomy 3. Anterior Ventricular Resection (Surgical Anterior Ventricular Restoration [SAVR], Dor procedure) |
| Intervention for Hypertrophic Cardiomyopathy | SPECIALIZED | Indicate if the patient required hypertrophic cardiomyopathy treatment. 1. Septal Myectomy 2. Septal Myectomy with mitral valve replacement or repair 3. Percutaneous Septal Alcohol embolization |
| Pericardiectomy | SPECIALIZED | Indicate if the patient required surgical removal of the pericardium, usually because of constrictive pericardial disease or infection. 1. Yes 2. No |
| Pericardiocentesis, Surgical | SPECIALIZED | Indicate if the patient required surgical drainage of fluid in the pericardim. 1. Yes 2. No |
| Closure of Patent Foramen Ovale (PFO) or Atrial Septal Defect | SPECIALIZED | Indicate if the patient required open surgical PFO closure or correction of atrial septal defect. 1. Yes, specify: a. Stroke b. Left to Right Shunt c. Right to Left Shunt d. Percutaneous e. Other, specify 2. No |
| Surgery for Congenital Heart Disease | SPECIALIZED | Indicate if the patient required surgery for congenital heart disease and specify what was required. 1. Yes, specify: a. Fontan Procedure b. Mustard Procedure c. Senning Procedure d. Other procedure, specify 2. No |
| Atrial Fibrillation Surgery | SPECIALIZED | Indicate if the patient required atrial fibrillation surgery. 1. Yes, specify a. Maze b. Modified Maze Procedure 2. No |

| Organ | SPECIALIZED | Indicate if the patient has required an organ transplant. |
|-----------------|-------------|--|
| Transplantation | | 1. Yes, select one |
| | | a. Heart |
| | | b. Heart/Lung |
| | | c. Lung, Single/Double |
| | | d. Kidney |
| | | e. Liver |
| | | f. Other, please specify (may include combination of organs) |
| | | 2. No |

D. COURSE IN-HOSPITAL

| FIELD NAME | CLASSIFICATION | DEFINITION |
|---------------|----------------|--|
| Renal Failure | SPECIALIZED | Documentation in the medical record of Renal Failure: 1. Yes – defined as a 50% increase in baseline/first Creatinine recorded in hospital 2. No 3. Unknown |
| Dialysis | SPECIALIZED | Documentation in the medical record of a new requirement for dialysis. 1. Yes, indicate duration a. Temporary b. Permanent 2. No |

E. CIRCULATORY/ VENTILATORY SUPPORT

| FIELD NAME | CLASSIFICATION | DEFINITION |
|----------------------------------|----------------|--|
| Mechanical Ventilator Support | SPECIALIZED | Indicate if patient required mechanical ventilatory support. Specify dates of initiation and termination of therapy. 1. Yes, specify: a. Mechanical ventilation/intubation b. CPAP c. BiPAP 2. No |

F. AT DISCHARGE FROM HOSPITAL

| FIELD NAME | CLASSIFICATION | DEFINITION |
|------------------|----------------|---|
| Discharge Status | ESSENTIAL | Indicate whether the patient was alive or deceased at discharge from this hospitalization. 1. Alive 2. Deceased |
| | | *This data element and definition originates from the Acute Coronary Syndrome (ACS) Data Dictionary and is duplicated here for consistency. |

| Death | ESSENTIAL | Specify date of death. |
|---|-----------|--|
| | | Indicate cause of death, if available: 1. Cardiovascular: a. Myocardial infarction b. Cardiogenic shock c. Heart failure d. Sudden cardiac death e. Cardiac arrest f. Arrhythmia (specify) g. Stroke h. Other, specify 2. Non-cardiovascular: a. Pulmonary embolism b. Cancer c. Trauma d. Sepsis e. Chronic obstructive lung disease f. Renal failure i. Other, specify 3. Indicate location of death: a. At home b. In hospice care c. In hospital j. Other, specify d. Unknown |
| Disposition after Health Care Encounter | ESSENTIAL | Indicate disposition after health care encounter. 1. Discharged to home or self care (routine discharge) 2. Discharged/transferred to another short-term general hospital for inpatient care 3. Discharged/transferred to skilled nursing facility (SNF) 4. Discharged/transferred to an intermediate care facility (ICF) 5. Discharged/transferred to another type of institution 6. Discharged/transferred to home under care of organized home health service organization 7. Left against medical advice or discontinued care 8. Discharged/transferred to home under care of a home IV drug therapy provider 9. Expired 10. Hospice-home 11. Hospice-medical facility 12. Discharged/transferred to an inpatient rehabilitation facility including rehabilitation distinct part units of a hospital Specify date. |
| Discharge Date | ESSENTIAL | Indicate the date the patient was discharged from hospital or left against medical advice or was transferred to another centre or died during this admission. |
| | | *This data element and definition originates from the Acute Coronary Syndrome (ACS) Data Dictionary and is duplicated here for consistency. |

| Patient Referral | ESSENTIAL | Patient referred to other care: 1. Heart Failure Specialty Clinic 2. Heart Failure Transitional Care by advanced practice nurses |
|------------------|-----------|--|
| | | Heart Failure Disease Management Program |
| | | Evaluation for Heart Transplant |
| | | Transitional Care (specify duration): |
| | | Home Health Care Heart Failure Nurse Case Manager |
| | | 3. Hospice or Palliative Care |
| | | Home Telemonitoring Ambulatory Cardiac Telemetric Monitoring (e.g. mobile cardiac outpatient telemetry) |
| | | Period of time enrolled in program and/or qualitative characterization of level of patient's success/participation in the program(s) may be specified. |

PART 11 – OUTCOMES

| FIELD NAME | CLASSIFICATION | DEFINITION |
|----------------------|----------------|--|
| Alive 30 Days post | SPECIALIZED | 1. Yes |
| service date end | | 2. No |
| Mortality (6 and 12 | SPECIALIZED | Specify date of death. |
| months) | | |
| | | Indicate cause of death, if available: |
| | | 1. Cardiovascular: |
| | | a. Myocardial infarction |
| | | b. Cardiogenic shock c. Heart failure |
| | | d. Sudden cardiac death |
| | | e. Cardiac arrest |
| | | f. Arrhythmia (specify) |
| | | g. Stroke |
| | | h. Other, specify |
| | | 2. Non-cardiovascular: |
| | | a. Pulmonary |
| | | b. Cancer |
| | | c. Trauma |
| | | d. Sepsis |
| | | e. Chronic obstructive lung disease |
| | | f. Renal failure |
| | | g. Other, specify 3. Indicate location of death, if known: |
| | | a. At home |
| | | b. In hospice care |
| | | c. In hospital |
| | | d. Other, specify |
| Follow up after last | SPECIALIZED | Date of first medical visit. |
| episode of care | | |
| | | GP/Specialist/Cardiologist/Heart Failure Clinics |
| | | 2. Patient referred to other care: |
| | | a. Heart Failure Specialty Clinic |
| | | b. Heart Failure Transitional Care by advanced practice nurses |
| | | c. Heart Failure Disease Management Programd. Evaluation for Heart Transplant |
| | | 3. Transitional Care (specify duration): |
| | | a. Home Health Care |
| | | b. Heart Failure Nurse Case Manager |
| | | c. Hospice or Palliative Care |
| | | d. Home Telemonitoring |
| | | e. Ambulatory Cardiac Telemetric Monitoring (e.g. mobile |
| | | cardiac outpatient telemetry) |
| | | Period of time enrolled in program and/or qualitative |
| | | characterization of level of patient's; success/participation in the |
| | | program(s) may be specified |
| | | 5. Other SPECIALIZED clinics (renal failure |
| | | /anemia/geriatrics/pacing/home monitoring) |

ACKNOWLEDGEMENT

The Canadian Cardiovascular Society would like to acknowledge the contributions of the following individuals in the development of the Heart Failure Data Dictionary Chapter:

Data Definitions Heart Failure Chapter Working Group

Justin Ezekowitz (Chair), University of Alberta

Abdul Al-Hesayen, St. Michael's Hospital (Ontario)

Anique Ducharme, Institut de Cardiologie de Montréal (Québec)

Yana Gurevich, Canadian Institute for Health Information

Jonathan G Howlett, Libin Cardiovascular Institute of Alberta

Laurie Lambert, Institut national d'excellence en santé et en services sociaux (Québec)

Douglas Lee, Institute for Clinical Evaluative Studies (Ontario)

Serge LePage, Université de Sherbrooke Sherbrooke (Québec)

Heather Ross, University Health Network (Ontario)

Elizabeth Swiggum, University of British Columbia (British Columbia)

Sean Virani, Vancouver General Hospital, University of British Columbia (British Columbia)

Sulan Dai, Public Health Agency of Canada

Data Definitions Steering Committee

Karin Humphries (Chair), University of British Columbia

Jafna Cox, Cardiovascular Health Nova Scotia

Ross Davies, University of Ottawa Heart Institute (Ontario)

Diane Galbraith, Alberta Provincial Project for Outcome Assessment in Coronary Heart Disease

Kori Kingsbury, Cardiac Care Network of Ontario

Andrew Kmetic, Cardiac Services BC

Dennis Ko, Institute for Clinical Evaluative Services (Ontario)

Laurie Lambert, Institut national d'excellence en santé et en services sociaux (Québec)

Anne McFarlane, Canadian Institute for Health Information

Sulan Dai, Public Health Agency of Canada

Mario Talajic (ex-Officio), Montreal Heart Institute (Québec) President, Canadian Cardiovascular Society Heather Ross (ex-officio), University Health Network (Ontario), Vice-President, Canadian Cardiovascular Society Blair O'Neill (ex-officio), Alberta Health Services Past President, Canadian Cardiovascular Society

Project Support

Anne Ferguson, Chief Executive Officer, Canadian Cardiovascular Society

Nick Neuheimer, Project Director and Director, Health Policy, Advocacy and External Relations, Canadian Cardiovascular Society

Holly Fan, Project Manager (external)

Production of these materials has been made possible by the Canadian Cardiovascular Society and through a financial contribution from the Public Health Agency of Canada.

DISCLAIMER

The views expressed herein do not necessarily represent the views of the Public Health Agency of Canada.

COPYRIGHT

© All rights reserved. No part of this document may be reproduced, stored in a retrieval system or transmitted in any format or by any means, electronic, mechanical, photocopying, recording or otherwise, without the proper written permission of The Canadian Cardiovascular SocietyTM.