

National Cardiac Registry (NCR)

Data Dictionary

Version 1.1

March 2023

Endorsed by the NCR Board 15/03/2023

Data Dictionary Version History

Date	Version	Description of changes
15 March 2023	1.0	Initial version
30 March 2023	1.1	Minor updates to the 'justification' for some elements.

Contents

Data	Diction	ary Version History	. 2
Abb	reviation	s	. 7
Fore	word		. 8
Time	eline of t	he National Cardiac Registry	. 9
Data	Collecti	on	10
Inclu	usion/Exc	clusion Criteria	11
Data	set Defi	nition Sources	12
Guid	le to Me	anings of Categories and Headings	13
App	roval and	d Limitations	14
Data	Definiti	ons	15
1.	Site		16
	1.1.	NCR Hospital ID	16
2.	Patie	nt	17
	2.1.	PatientID NCR	17
	2.2.	Patient ID State Code	18
	2.3.	Sex	19
	2.4.	Indigenous Status	20
3.	Admi	ssion and Preprocedural	21
	3.1.	Admission and Clinical Presentation	21
	3.1.1.	Postcode	21
	3.1.2.	Hospital ID	22
	3.1.3.	Date of arrival at PCI Hospital	23
	3.1.4.	Time of arrival at PCI Hospital	24
	3.1.5.	Patient Age	25
	3.1.6.	Patient Height	26
	3.1.7.	Patient Weight	27
	3.1.8.	Body Mass Index (BMI) (kg/m^2)	28
	3.1.9.	PCI Procedure Date	29
	3.1.10.	PCI Procedure Time	30
	3.1.11.	Primary Operator	31
	3.2.	Pre-Procedural Risk Factors	32
	3.2.1.	Diabetes	32
	3.2.2.	Diabetes Management Type	33
	3.2.3.	Peripheral Vascular Disease History 1 – With Intervention	34
	3.2.4.	Peripheral Vascular Disease history 2 – Broader Definition	35

	3.2.5.	Previous CABG	36
	3.2.6.	When was the most recent CABG?	. 37
	3.2.7.	Previous PCI	38
	3.2.8.	When was the most recent PCI?	39
	3.3. I	Pre-Procedural Renal Status	40
	3.3.1.	Last Pre-Procedure Creatinine (μmol/L)	40
	3.3.2.	Creatinine Result Not Available	41
	3.4. I	eft Ventricular (LV) Function	42
	3.4.1.	LVEF Test Performed	42
	3.4.2.	Date of LVEF Test	43
	3.4.3.	LVEF Test Type	. 44
	3.4.4.	Ejection Fraction (EF) Test Result Digitally Derived	. 45
	3.4.5.	Ejection Fraction (EF) Test Result Estimated	46
	3.4.6.	Estimated Glomerular Filtration Rate (Cockroft-Gault formula)	. 47
	3.4.7.	Estimated Glomerular Filtration Rate Imported	. 48
	3.5.	Clinical Presentation	49
	3.5.1.	Cardiogenic Shock	49
	3.5.2.	Out-of-Hospital Cardiac Arrest (OHCA)	50
	3.5.3.	Pre-Procedural Intubation	51
4.	Proce	dure	52
	4.1.	Clinical Symptoms	52
	4.1.1.	ACS	52
	4.1.2.	Date of ACS Symptom Onset	53
	4.1.3.	Time of ACS Symptom Onset	54
	4.1.4.	Onset Time Not Available	55
	4.1.5.	Type of ACS	56
	4.1.6.	Inter-hospital Transfer	58
	4.1.7.	Pre-hospital Notification	59
	4.1.8.	Balloon/Device Date	60
	4.1.9.	Balloon/Device Time	61
	4.1.10.	Symptom Onset to Reperfusion	62
	4.1.11.	Door to balloon/Device time	63
	4.1.12.	Self-presenter	64
	4.1.13.	Date of First Medical Contact (FMC)	65
	4.1.14.	Time of FMC	66
	4.1.15.	Symptom Onset to FMC	67

4.1	1.16.	FMC to Reperfusion	68
4.1	1.17.	FMC to Door	69
4.2	1.18.	Date of Diagnostic ECG	70
4.1	1.19.	Time of Diagnostic ECG	71
4.1	1.20.	FMC to Diagnostic ECG	72
4.1	1.21.	Diagnostic ECG to Balloon	73
4.1	1.22.	Inpatient at Time of ACS	74
4.2.	PCI I	ndication	75
4.3.	Proc	edure Details	78
4.3	3.1.	Percutaneous Entry Location	78
4.3	3.2.	Procedural Intubation Required	79
4.3	3.3.	Mechanical Circulatory Support	80
4.3	3.4.	Lesion Location	81
4.3	3.5.	In-stent Restenosis (ISR)	82
4.3	3.6.	Stent Thrombosis	83
4.3	3.7.	Lesion Successfully Treated	84
4.3	3.8.	Type of Stent(s) Implanted	85
4.3	3.9.	Total Entrance Radiation Dose (Air Kerma)	87
4.3	3.10.	Total Exam Dose Area Product in mGy/m2	88
4.3	3.11.	Total Fluoroscopy Time	89
5.	Post Pr	ocedure	90
5.1.	Post	-Procedure & In-Hospital Complications	90
5.2	1.1.	In-hospital MI	90
5.2	1.2.	Subsequent PCI (in this admission)	92
5.2	1.3.	Planned Subsequent PCI	93
5.2	1.4.	In-Hospital Target Vessel (PCI)	94
5.2	1.5.	In-Hospital Target Lesion Revascularisation	95
5.2	1.6.	In-Hospital Cardiothoracic Surgery	96
5.2	1.7.	Planned Cardiothoracic Surgery	97
5.2	1.8.	In-Hospital TVR (CABG)	98
5.2	1.9.	In-Hospital Stroke	99
5.2	1.10.	In-Hospital Stroke Type	L 00
5.2	1.11.	In-Hospital Bleeding	l 01
5.2	1.12.	In-Hospital Bleeding Site	L03
5.2	1.13.	In-Hospital Stent Thrombosis	L 0 4
ā	Discha	rge 1	105

	6.1.	Discharge Details	105
	6.1.1	Discharge Status	105
	6.1.2	Date of Discharge/Hospital Mortality	107
	6.1.3	Length of Stay	108
	6.1.4	Cardiac Rehabilitation Referral	109
	6.1.5	Primary Cause of Death	110
	6.2.	Discharge Medications	111
	6.2.1	Aspirin Prescribed at Discharge	111
	6.2.2	Other Antiplatelet Prescribed at Discharge	112
	6.2.3	Statin Prescribed at Discharge	113
	6.2.4	Other Lipid Lowering Therapies Prescribed at Discharge	114
7.	30 Da	y Follow Up	115
	7.1.	30 Day Outcomes	115
	7.1.1	Date of Follow-Up	115
	7.1.2	Follow-Up Status	116
	7.1.3	Date of Death	117
	7.1.4	Primary Cause of Death	118
	7.1.5	New Ml	119
	7.1.6	New Stent Thrombosis	121
	7.1.7	New Stroke	123
	7.1.8	New Stroke Type	124
	7.2.	30-Day Rehospitalisations	125
	7.2.1	Cardiac Rehospitalisation	125
	7.2.2	Cardiac Rehospitalisation Date	127
	7.2.3	Planned Cardiac Rehospitalisation	128
	7.2.4	PCI Rehospitalisation	129
	7.2.5	Target Vessel Revascularisation (PCI)	130
	7.2.6	Target Lesion Revascularisation (TLR)	131
	7.2.7	CABG Rehospitalisation	132
	7.2.8	Target Vessel CABG	133

Abbreviations

ACS Acute Coronary Syndrome
ARC Academic Research Consortium

ARIA Accessibility/Remoteness Index of Australia
BARC Bleeding Academic Research Consortium

BMI Body Mass Index
BMS Bare Metal Stent
BP Blood Pressure
BRS Bioresorbable Stent

CABG Coronary Artery Bypass Graft
CAD Coronary Artery Disease
CCL Cardiac Catheter Laboratory
CCS Canadian Cardiovascular Society

CK-MB Creatine Kinase MB

CT Computerised Tomography
DAPT Dual Antiplatelet Therapy
DBT Door to Balloon Time
DES Drug Eluting Stent
ECG Electrocardiogram

ECMO Extracorporeal Membrane Oxygenation

EF Ejection Fraction
FMC First Medical Contact
GFR Glomerular Filtration Rate
GP General Practitioner

Hg Mercury (Used to measure pressure)

IABP Intra-Aortic Balloon Pump

ICD-10 International Classification of Diseases 10

IV Intravenous

LUCAS Lund University Cardiac Arrest System (mechanical chest compression device)

LV Left Ventricular

LVAD Left Ventricular Assist Device
LVEF Left Ventricular Ejection Fraction

MACCE Major Adverse Cardiac and/or Cerebrovascular Event

MACE Major Adverse Cardiac Event

MI Myocardial Infarction

MR(I) Magnetic Resonance (Imaging)

NSTEMI Non-ST Elevation Myocardial Infarction
PCI Percutaneous Coronary Intervention

POBA Plain Balloon Angioplasty PVD Peripheral Vascular Disease

QI Quality Indicator

SEIFA Socio-Economic Indexes for Areas
STEMI ST Elevation Myocardial Infarction
TIMI Thrombolysis in Myocardial Infarction

TLR Target Lesion Revascularisation
TVR Target Vessel Revascularisation

ULN Upper Limit of Normal

Foreword

The NCR Data Dictionary was created by the National Cardiac Registry Project Team, School of Public Health and Preventative Medicine, Monash University, on behalf of/by National Cardiac Registry Ltd. Members of the following committees, without whom the data dictionary and dataset would not be possible, are named below.

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Timeline of the National Cardiac Registry

November 2016 Ischaemic Heart Disease named as the #1 priority

area in The Australian Commission on Safety and Quality in Health Care's Prioritised list of clinical domains for clinical quality registry development

final report.

March 2016 The Department and existing State registries

agree to form a National Cardiac Registry

October 2018 Engagement of clinicians from States & Territories

and development of strategy

January-March 2019 Development of an initiation and establishment

plan

April 2019 NCR project team employed through Monash

University

August 2019 Funding approved for States and Territories to

help facilitate in participation in NCR

November 2019 Inaugural Steering Committee meeting held with

clinical and jurisdictional representatives from all

eight States & Territories

January 2020 NCR Minimum Dataset and Dictionary Approved

by the NCR Data Subcommittee

May 2020 NCR Ltd incorporated by the Australian Securities

and Investments Commission

July 2020 NCR Ltd. Inaugural board meeting

October 2020 NCR inaugural status report

July 2021 NCR Platform goes live

April 2022 Funding secured within federal budget to

support registry operations

May 2022 Launch of first public facing Annual Status Report

July 2022 All eight jurisdictions contributing data elements into

the NCR Platform

Data Collection

Data for the NCR is collected at state/territory level by the following contributing state and territory registries:

ACT Cardiac Outcomes Registry (ACTCOR)

Coronary Angiogram Database of South Australia (CADOSA)

New South Wales Cardiac Outcomes Registry (NSWCOR)

Northern Territory Top End Coronary Database (NTTCD)

Queensland Cardiac Outcomes Registry (QCOR)

Victorian Cardiac Outcomes Registry (VCOR)

Western Australia Cardiac Outcomes Registry (WACOR)

Tasmania participates via VCOR

Contributing registries upload a CSV file to the NCR platform at agreed time intervals. Data is checked and validated as part of the import process to ensure that the data provided meets the business rules as outlined in the following pages. For more information on this process please see - https://nationalcardiacregistry.org.au/data-flow/.

Inclusion/Exclusion Criteria

Patients aged 18 years or older, who present to a participating hospital in Australia and are treated with percutaneous coronary intervention (PCI).

A PCI is deemed to have taken place if any coronary device approaches, probes or crosses one or more coronary lesions with the intention of performing a coronary intervention. Usually, this device will be a guide wire. The only exception to this is patients who have an adverse cardiac event (during an attempted PCI) that necessitates procedure termination prior to the introduction of a coronary device. This rare type of case will also be defined as a PCI and therefore this will classify as a complication. That is, any patient who crosses the threshold of the cardiac catheter lab for the purpose of PCI as defined above, is recorded in the NCR dataset.

Dataset Definition Sources

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- Australian Commission on Safety and Quality in Health Care 2019. Acute Coronary Syndromes Clinical Care Standard. Australian Commission on Safety and Quality in Health Care.
- Canadian Cardiovascular Society 2012. The Canadian Cardiovascular Society Data Dictionary Core Elements and Demographics Data Elements and Definitions. Canadian Cardiovascular Society (CCS).
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- Lawton, J. S., Tamis-Holland, J. E., Bangalore, S., Bates, E. R., Beckie, T. M., Bischoff, J. M., Bittl, J. A., Cohen, M. G., DiMaio, J. M., Don, C. W., Fremes, S. E., Gaudino, M. F., Goldberger, Z. D., Grant, M. C., Jaswal, J. B., Kurlansky, P. A., Mehran, R., Metkus, T. S., Jr., Nnacheta, L. C., Rao, S. V., Sellke, F. W., Sharma, G., Yong, C. M. & Zwischenberger, B. A. 2022. 2021 ACC/AHA/SCAI Guideline for Coronary Artery Revascularization: A Report of the American College of Cardiology/American Heart Association Joint Committee on Clinical Practice Guidelines. *Circulation*, 145, e18-e114.
- National Institute for Cardiovascular Outcomes Research 2015. National Audit of Percutaneous Coronary Interventions Minimum Data Standard. United Kingdom: National Institute for Cardiovascular Outcomes Research (NICOR).
- National Institute for Health Care and Excellence 2011. Stable Angina: Management. United Kingdom: National Institute for Health Care and Excellence (NICE).
- Swedish Web-system for Enhancement and Development of Evidence-based care in Heart disease Evaluated According to Recommended Therapies (SWEDEHEART) 2021. SWEDEHEART Annual Report 2021. Swedish Web-system for Enhancement and Development of Evidence-based care in Heart disease Evaluated According to Recommended Therapies (SWEDEHEART).
- The National Institute for Cardiovascular Outcomes Research: National Cardiac Audit Programme 2021. National Audit of Percutaneous Coronary Intervention (NAPCI). The National Institute for Cardiovascular Outcomes Research (NICOR): National Cardiac Audit Programme (NCAP).
- Victorian Cardiac Outcomes Registry 2020. PCI Data Definitions. Victorian Cardiac Outcomes Registry (VCOR).

Guide to Meanings of Categories and Headings

Defining Attributes

Name

Name of the collected variable.

Field Name

The actual code used in the platform.

Definition

Concise statement expressing essential nature of a data field and its differentiation from all other data fields. Supporting definition without Codes or Variables.

Justification

The reason for collecting the data field.

Obligation

Mandatory/optional for the purpose of meeting the minimum dataset as predetermined by NCR. It is understood that not all sites and registries collect the minimum dataset.

Representational Attributes

Permitted Values

Set of possible values for the data field. May be a code set, or description of possible values.

Range

The minimum and maximum possible values allowed for the data field.

Guide for Use

Comments to assist in further defining aspects of the data element.

Validation Rules

Assist in reducing input error. Where validation rules are known to exist, they have been included.

Maximum Field Size

The number of characters to represent the data field values. Includes special characters.

Related Data Field

Other data fields in this data dictionary that have some direct relationship with the data fields being described. Specifies fields that may be derived from, or may contribute to deriving the value of the field being defined.

Data Type

Type of symbol or character or other designation used to represent the data field, for example alphanumeric values are text, number or date/time.

Parent / Child

The relationship of the data field to the primary key i.e., if the field is parent there may be one or many values per field per primary key

Behaviour

Data rules as per the platform.

Approval and Limitations

This version of the NCR Data Dictionary has been approved by the NCR Board and Steering Committee. The data dictionary is periodically reviewed and updated to reflect changes. There may be specific limitations in version 1.0 which do not affect the NCR purpose or definitions.

Data Definitions

1. Site

1.1. NCR Hospital ID

Defining Attributes

Name NCR Hospital ID

Field Name ncrhid

Definition

Justification Used to identify a unique hospital.

Obligation Always

Representational Attributes

Permitted Values Values must match the list provided to the NCR by the state registry.

Range NNNNNN

Guide for Use Each Hospital has a set of defining attributes. These include: State/Territory, Public or Private facility, Emergency

Department Y/N and if Hospital perform Coronary Artery Bypass Graft Surgery (CABG) Y/N.

Validation Rules Field cannot be blank

Maximum Field Size 6
Related Data Field hid
Data Type Text

Parent / Child Behaviour

2. Patient

2.1. PatientID NCR

Defining Attributes

Name Patient ID NCR Field Name ncrpatientid

Definition State generated ID, must be unique to each patient

Justification Used to identify a unique patient record.

Obligation Always

Representational Attributes

Permitted Values N

Range

Guide for Use System generate unique ID number = site state code + site public/private code + PatientID

Validation Rules Field cannot be blank

Maximum Field Size 18

Related Data Field

Data Type Text/ Number

Parent / Child

Behaviour Reject file if NULL

2.2. Patient ID State Code

Defining Attributes

Name PatientID State Code

Field Name patientid

Definition Unique patient identification number, generated by each State.

Justification Collected to allow contributing registries to identify a record within their own registry system.

Obligation Always

Representational Attributes

Permitted Values N

Range

Guide for Use Values must match or be able to be mapped back to the state level patient ID.

Validation Rules Field cannot be blank

18

Maximum Field Size

Related Data Field

Data Type Text/ Number

Parent / Child

Behaviour Reject file if NULL

2.3. Sex

Defining Attributes

Name Sex Field Name sex

Definition The sex of an individual.

Justification Collected to determine differences in incidence, management and mortality associated with biological sex and risk

adjustment modelling.

Obligation Always

Representation	nal Attributes		
Permitted	l Values	Code	Description
		1	Male
		2	Female
Range		1-2	
Guide for	llse		

Guide for Use

Field cannot be blank Validation Rules

Maximum Field Size 1

Related Data Field

Data Type Number

Parent / Child

Reject file if NULL Behaviour

2.4. Indigenous Status

Defining Attributes

Name Indigenous Status

Field Name inds

Definition Australian Indigenous Status, including Aboriginal, Torres Strait Islander, Both, or Neither.

Justification Collected to determine differences in incidence, management and mortality associated with Indigenous Status.

Obligation Always

Obligation	Always		
Representational Attributes	_		
Permitted Values	Code	Description	
	0	Neither	
	1	Aboriginal	
	2	Torres Strait Islander	
	3	Both Aboriginal and Torres Strait Islander	
	-1	Unknown	
Range	-1 - 3		
Guide for Use	Is the patient Aboriginal or Torres Strait Islander, Both, Neither, or is their Indigenous status Unknown?		
Validation Rules	Field cannot be blank		
Maximum Field Size	2		
Related Data Field			
Data Type	Number		
Parent / Child			
Behaviour	Reject file if NULL		

3. Admission and Preprocedural

3.1. Admission and Clinical Presentation

3.1.1. Postcode

Defining Attributes

Name Postcode Field Name pcode

Definition Primary residential postal code, if known. Numeric descriptor for postal delivery area by suburb or locality.

Justification Collected for demographic statistical reporting at time of admission and SEIFA and ARIA status

Obligation Optional

Representational Attributes

Permitted Values NNNN Range 0-9999

Guide for Use What was the patient's postcode at the time of admission?

Validation Rules

Maximum Field Size 4

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL

3.1.2. Hospital ID

Defining Attributes

Name Hospital ID

Field Name hid

Definition Hospital identification number allocated by State/Territory Staff when setting up a new hospital in their system.

Justification Collected to monitor clinical guideline compliance.

Obligation Always

Representational Attributes

Permitted Values NNNNNN Range 0-999999

Guide for Use Enter the unique code associated with the hospital at which the PCI was undertaken.

Validation Rules Field cannot be blank

Maximum Field Size 6

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL

3.1.3. Date of Arrival at PCI Hospital

Defining Attributes

Name Date of arrival at PCI hospital

Field Name doa

Definition Date the patient was admitted or transferred to the hospital in which their PCI was performed.

Justification Collected to identify a unique PCI procedure. Collected to allow reporting of NCR QIs – Time from door to PCI-mediated

reperfusion.

Obligation Always

Representational Attributes

Permitted Values DD/MM/YYYY Range NN/NN/NNNN

Guide for Use On what date did the patient arrive at the PCI hospital?

Validation Rules Field cannot be blank. Field cannot be the same as Time of Arrival, Date of Procedure, Time of Procedure and Patient ID

across multiple entries.

Maximum Field Size 10

Related Data Field

Data Type Date/Time

Parent / Child

Behaviour Reject file if NULL

Reject file if doa, toa, dop, top and patientID are all the same across multiple entries.

3.1.4. Time of Arrival at PCI Hospital

Defining Attributes

Name Time of arrival at PCI hospital

Field Name toa

Definition Time the patient was admitted or transferred to the hospital in which their PCI was performed. This is the time the patient

was triaged (entered into the hospital administrative system).

Justification Collected to identify a unique PCI procedure. Collected to allow reporting of NCR QIs – Time from door to PCI-mediated

reperfusion.

Obligation Always

Representational Attributes

Permitted Values HH:MM Range 00:00-23:59

Guide for Use At what time did the patient arrive at the PCI hospital? This is the time when they were triaged (entered into the hospital

administrative system).

Validation Rules Field cannot be blank. Field cannot be the same as Time of Arrival, Date of Procedure, Time of Procedure and Patient ID

across multiple entries.

Maximum Field Size

Related Data Field

Data Type Date/Time

Parent / Child

Behaviour Reject file if NULL

5

Reject file if doa, toa, dop, top and patientID are all the same across multiple entries.

3.1.5. Patient Age

Defining Attributes

Name Patient Age

Field Name age

Definition Age of the patient at the date of procedure, measured as number of years.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always

Representational Attributes

Permitted Values Code Description 18-110 Valid Age

0-18 Invalid Age

Range 18-110

Guide for Use How old is the patient at the date of procedure?

Validation Rules Field cannot be blank. Field cannot <18.

Maximum Field Size

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL. Reject file if <18

3

3.1.6. Patient Height

Defining Attributes

Name Patient Height

Field Name htm

Definition Height in centimeters in bare or stockinged feet. Can be self-reported, estimated, or measured.

Justification Collected to calculate BMI.

Obligation Optional

Representational Attributes

Permitted Values NNN Range 111-231

Guide for Use How tall is the patient in centimeters?

Validation Rules

Maximum Field Size 3

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL

3.1.7. Patient Weight

Defining Attributes

Name Patient Weight

Field Name wkg

Definition Weight in kilograms in light clothing. Can be self-reported, estimated or measured.

Justification Collected to calculate BMI.

Obligation Optional

Representational Attributes

Permitted Values NNN Range 35-300

Guide for Use How much does the patient weigh in kilograms?

Validation Rules

Maximum Field Size 3

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if value entered is >300

3.1.8. Body Mass Index (BMI) (kg/m^2)

Defining Attributes

Name Body Mass Index [kg/m^2]

Field Name bmi

Definition Body Mass Index gives an indication as to a patient's body size.

Justification A characteristic as to the risk profile of the patient.

Obligation Derived Variable

Representational Attributes

Permitted Values NN.N (kg/m²)

Range

Guide for Use What is the patient's BMI?

Validation Rules

Maximum Field Size 3

Related Data Field

Data Type Number

Parent / Child

Behaviour Derived - No import on this field. BMI = wkg/(htm/100)^2

3.1.9. PCI Procedure Date

Defining Attributes

Name PCI Procedure Date

Field Name dop

Definition Date on which the patient crossed the threshold of entry into the cardiac catheter laboratory for the current PCI in

DD/MM/YYYY format.

Note: Where multiple procedures occur during the index hospitalisation, a separate entry must be recorded for every

relevant procedure.

Justification Collected to identify a unique PCI procedure and to benchmark performance and timeliness of treatment.

Obligation Always

Representational Attributes

Permitted Values DD/MM/YYYY Range NN/NN/NNNN

Guide for Use On what date was the PCI performed?

10

Validation Rules Field cannot be blank. PCI Procedure Date must be < date of file import. PCI Procedure Date must be > Date of Arrival at

PCI Hospital. Date of Arrival at PCI Hospital, Time of Arrival at PCI Hospital, PCI Procedure Date and Patient ID cannot be

the same across multiple entries.

Maximum Field Size

Related Data Field

Data Type Date/Time

Parent / Child

Behaviour Reject file if dop is > date of file import

Reject file if dop is NULL Reject file if dop is <doa

Reject file if doa, toa, dop, top and patientID are all the same across multiple entries within this import

3.1.10. PCI Procedure Time

Defining Attributes

Name PCI Procedure Time

Field Name top

Definition Time the patient crossed the threshold of entry into the cardiac catheter laboratory for the current PCI.

Note: Where multiple procedures occur during the index hospitalisation, a separate entry must be recorded for every

relevant procedure.

Justification Collected to identify a unique PCI procedure and to benchmark performance and timeliness of treatment.

Obligation Always

Representational Attributes

Permitted Values DD/MM/YYYY
Range NN/NN/NNN

Guide for Use On what date was the PCI performed?

Validation Rules Field cannot be blank. PCI Procedure Date cannot = Date of Arrival when PCI Procedure Time < Time of Arrival at PCI

Hospital. Date of Arrival at PCI Hospital, Time of Arrival at PCI Hospital, PCI Procedure Date and Patient ID cannot be the

same across multiple entries.

Maximum Field Size

Related Data Field

Data Type Date/Time

Parent / Child

Behaviour Reject file if dop=doa and top<toa

10

Reject file if top is NULL

Reject file if doa, toa, dop, top and patientID are all the same across multiple entries within this import

3.1.11. Primary Operator

Defining Attributes

Name Primary Operator

Field Name

Definition Unique code of the cardiologist/clinician who is ultimately responsible for the PCI as allocated by the State Registry.

Justification To be used in the future for clinician level reporting. Note: The NCR does not currently undertake any clinician level

reporting.

Obligation Optional

Representational Attributes

Permitted Values

Range

Guide for Use Who was the interventional cardiologist responsible for the PCI?

Validation Rules Maximum Field Size

Related Data Field

Data Type

Parent / Child Behaviour

Text

3.2. Pre-Procedural Risk Factors

3.2.1. Diabetes

Defining Attributes

Name Diabetes Field Name db

Definition Indicate if the patient has been diagnosed with diabetes mellitus regardless of duration of disease and, this includes a

medical diagnosis made during the current admission.

Justification Collected to determine the patient's pre-existing risk profile.

Obligation Always

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Does the patient have a diagnosis of diabetes?

Validation Rules Field cannot be blank.

Maximum Field Size 1
Related Data Field dbm
Data Type Number
Parent / Child Parent

Behaviour Reject file if NULL.

3.2.2. Diabetes Management Type

Defining Attributes

Name Diabetes Management Type

Field Name dbm

Definition The diabetes management type for patients with a diagnosis of diabetes.

Oral: Patient requires oral hypoglycaemic medication to control their condition.

Insulin: Patient requires insulin to control their condition, with or without oral therapy.

Diet: Patient has received dietary advice appropriate to their condition but is not taking medication to lower blood sugar

Justification Collected to determine the patient's pre-existing risk profile.

Obligation Always, if Diabetes = Yes

	, ,		
Representational Attributes			
Permitted Values	Code	Description	
	1	Oral	
	2	Insulin	
	3	Diet	
Range	1-3		
Guide for Use	Where Diabetes =Yes, indicate the management type		
Validation Rules	Field cannot be blank if Diabetes = Yes. Field cannot be complete if Diabetes = No.		
Maximum Field Size	1		
Related Data Field	db		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if db=0, and data is in dbm		
	Reject file if db=1 AND dbm is NULL		

3.2.3. Peripheral Vascular Disease History 1 – With Intervention

Defining Attributes

Name Peripheral Vascular Disease History 1 – With Intervention

Field Name pvd1

Definition The patient displays evidence of either chronic or acute PVD. The presence of PVD must be demonstrated by vascular

reconstruction or amputation for arterial insufficiency, bypass surgery or percutaneous intervention.

Note: This definition of PVD must be accompanied with intervention such as reconstruction, amputation, surgery or percutaneous intervention. Includes the aorta, extremities and carotid vessels. If no intervention, select the broader

option of PVD 2.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always, if pvd2 is blank.

	, , ,		
Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
Range	0-1		
Guide for Use	Does the patient have a history of PVD with intervention?		
Validation Rules	Field cannot be complete if pvd2 is complete. Field cannot be blank if pvd2 is blank.		
Maximum Field Size	1		
Related Data Field	pvd2		
Data Type	Number		
Parent / Child			
Behaviour	Reject file if pvd1 and pvd2 both have data		
	Reject file if p	vd1 and pvd2 are both NULL	

3.2.4. Peripheral Vascular Disease history 2 – Broader Definition

Defining Attributes

Name Peripheral Vascular Disease History 2 – Broader Definition

Field Name pvd2

Definition The patient has a current or previous history of peripheral vascular disease (includes subclavian, iliac, femoral, and upper-

and lower-extremity vessels; excludes renal, coronary, cerebral, and mesenteric vessels and aneurysms.

This can include: Claudication on exertion, amputation for arterial vascular insufficiency, vascular reconstruction, bypass surgery, or percutaneous revascularisation in the arteries of the extremities, positive non-invasive test (e.g., ankle brachial index <=0.9, ultrasound, MR, CT imaging of >50% diameter stenosis in any peripheral artery (i.e., subclavian,

femoral, iliac) or angiographic imaging.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always, if pvd1 is blank.

Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
Range	0-1		
Guide for Use	Does the patient have a history of PVD with or without intervention?		
Validation Rules	Field cannot be complete if pvd1 is complete. Field cannot be blank if pvd1 is blank.		
Maximum Field Size	1		
Related Data Field	pvd1		
Data Type	Number		
Parent / Child			
Behaviour	Reject file if pvd1 and pvd2 both have data		
	Reject file if p	pvd1 and pvd2 are both NULL	

3.2.5. Previous CABG

Defining Attributes

Name Previous CABG

Field Name pcabg

Definition Previous Coronary Artery Bypass Graft (CABG) Surgery prior to the current PCI procedure

Justification Collected to determine the patient's pre-existing risk profile.

Obligation Always

Representational Attributes

Permitted Values Code Description

1

0 No 1 Yes

Range 0-1

Guide for Use Has the patient previously undergone a CABG?

Validation Rules Field cannot be blank.

Maximum Field Size

Related Data Field

Data Type Number Parent / Child Parent

Behaviour Reject file if NULL

3.2.6. When was the most recent CABG?

Defining Attributes

Name When was the most recent CABG?

Field Name dpcabg

Definition The date on which patient had their most recent CABG in DD/MM/YYYY format. Note: If only the year is known or

estimated, this will suffice, enter as 01/01/YYYY.

Justification Collected to determine the patient's pre-existing risk profile.

Obligation Always, if pcabg=1.

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use If Previous CABG is coded 1=Yes, on what date?

Validation Rules Field cannot >Date of PCI Procedure. Field cannot be complete if Previous CABG = No. Field cannot be empty if Previous

CABG = Yes.

Maximum Field Size 10
Related Data Field pcabg
Data Type Date/Time
Parent / Child Child

Behaviour Reject file if dpcabg is >dop

Reject file if pcabg = 0 AND dpcabg has data Reject file if pcabg = 1 AND dpcabg is NULL

3.2.7. Previous PCI

Defining Attributes

Name Previous PCI

Field Name ppci

Definition Indicate if the patient has had a prior Percutaneous Transluminal Coronary Angioplasty, Coronary Atherectomy, and/or

coronary stent done at any time prior to the current PCI procedure. Note: This may include a PCI performed during the

current admission.

Justification Collected to determine the patient's pre-existing risk profile.

Obligation Always

0.0.1.04.1011	, , .			
Representational Attributes				
Permitted Values	Code	Description		
	0	No		
	1	Yes		
Range	0-1			
Guide for Use	Has the patient had a prior PCI?			
Validation Rules	Field cannot be blank.			

Maximum Field Size

Related Data Field

Data Type Number Parent / Child Parent

Behaviour Reject file if NULL

1

3.2.8. When was the most recent PCI?

Defining Attributes

Name When was the most recent PCI?

Field Name dppci

Definition The date on which the patient had their most recent PCI in DD/MM/YYYY format. Note: if only the year is known or

estimated, this will suffice, enter as 01/01/YYYY.

Justification Collected to determine the patient's pre-existing risk profile.

Obligation Always, if Previous PCI = Yes

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use If Previous PCI is coded 1=Yes, on what date?

Validation Rules Field cannot be blank if Previous PCI = Yes. Date of Previous PCI must ≤ PCI Procedure Date. Field cannot be complete if

Previous PCI = No.

Maximum Field Size 10 Related Data Field ppci

Data Type Date/Time Parent / Child Child

Behaviour Reject file if dpcabg is >dop

Reject file if pcabg =0 AND dpcabg has data Reject file if pcabg = 1 AND dpcabg is NULL

3.3. Pre-Procedural Renal Status

3.3.1. Last Pre-Procedure Creatinine (µmol/L)

Defining Attributes

Name Last Pre-Procedure Creatinine (μmol/L)

Field Name pcr

Definition The last serum creatinine levels recorded within 60 days prior to the current PCI (in µmol/L). Note: To convert from

mmol/L to µmol/L, multiply by 1000 or move decimal point 3 spaces to the right.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always, if npcr is blank

Representational Attributes

Permitted Values NN-NNNN μmol/L

Range 20-2000

Guide for Use What was the last pre-procedure creatinine?

Validation Rules Field must be complete, if Creatinine Result Not Available is blank. Field must be blank, if Creatinine Result Not Available

is complete.

Maximum Field Size 4
Related Data Field npcr
Data Type Number

Parent / Child

Behaviour Reject file if npcr=1 and pcr have data

Reject file if both pcr AND ncpr are NULL

3.3.2. Creatinine Result Not Available

Defining Attributes

Name Creatinine Result Not Available

Field Name npcr

Definition Last pre-procedure creatinine results not available.

Justification Used to determine if no creatinine test was undertaken.

Obligation Always, if pcr is blank

Representationa	l Attributes

Permitted Values Code Description

1 Yes, not available

Range

Guide for Use Is the last pre-procedure creatinine not available?

Validation Rules Field cannot be complete if Last Pre-Procedure Creatinine is complete. Field cannot be blank if Last Pre-Procedure

Creatinine is blank.

Maximum Field Size 1
Related Data Field pcr
Data Type Number

Parent / Child

Behaviour Reject file if there is a value in pcr and npcr=1

Reject file if both pcr AND npcr are NULL

3.4. Left Ventricular (LV) Function

3.4.1. LVEF Test Performed

Defining Attributes

Name LVEF Test Performed

Field Name eftp

Definition For all patients (excluding STEMIs – see note below) indicate whether the patient's ventricular ejection fraction (EF) was

measured (or estimated) within 6 months prior to the current procedure up to four weeks post-discharge. This includes the period leading up to and including the cardiac catheter lab visit, after the lab visit and up to 30 days after the patient was discharged. If multiple test results are available during this period, select the test result closest to the date/time of

the index PCI.

Note: For STEMI patients a LVEF test must have been recorded during the index admission or up to 30 days post-discharge for this item to be coded 'yes'. For these patients, where no LVEF test was performed during the index admission, code

'no'.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use For STEMI patients, was a LVEF test performed during the index admission or up to 30 days post-discharge, and for other

patients was a LVEF test performed within 6 months prior, or up to 30 days post-discharge?

Validation Rules Field cannot be blank.

Maximum Field Size 1

Related Data Field

Data Type Number Parent / Child Parent

3.4.2. Date of LVFF Test

Defining Attributes

Name Date of LVEF test

def Field Name

Definition The date of the most recent LVEF test. Note: Where multiple LVEF test results exist within the 6-month period leading

up to the index admission or anytime during the index admission but within 30 days post-discharge, record the test result

closest to the index PCI date/time.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always, if LVEF Test Performed = Yes

Representational Attributes Permitted Values NN/NN/NNNN Range DD/MM/YYYY Guide for Use Where LVEF test performed= Yes, on what date was the most recent LVEF test? Validation Rules If Type of ACS is not coded STEMI, then Date of LVEF test must be between [6 months prior to Date of Procedure] and [Date of Discharge/Hospital Mortality+30 days] If Type of ACS is STEMI, then Date of LVEF test must be between Date of Arrival and [Date of Discharge/Hospital Mortality+30 Days]

Field must be blank, if LVEF Test Performed = No. Field must be complete if LVEF Test Performed = Yes. Date of LVEF Test

must <Date of Discharge/Hospital Mortality+30 days.

Maximum Field Size 10 Related Data Field eftp Date/Time Data Type Parent / Child Child

Reject file if eftp=0 and def has data Behaviour

Reject file if eftp = 1 and def is NULL Reject file if def is >dod+30 days Reject file if acst=3 AND def <dop

Reject file if acst is NOT = 3 and def is <dop-6 months

3.4.3. LVEF Test Type

Defining Attributes

Name LVEF Test Type

Field Name eft

Definition Select ONE of the following: Echocardiography; Angiography Gated Cardiac Blood Pool Scan; Magnetic resonance

imaging (MRI) Myocardial Perfusion Scan; Not Stated/Inadequately described.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always, if LVEF Test Performed = Yes

Opligation	Aways, it Even reserve in ordined these		
Representational Attributes			
Permitted Values	Code	Description	
	1	Echocardiography	
	2	Angiography	
	3	Gated Cardiac Blood Pool Scan	
	4	Magnetic Resonance Imaging (MRI)	
	5	Myocardial Reperfusion Scan	
	6	Not Stated/Inadequately Described	
Range	1-6		
Guide for Use	Where LVEF test performed is coded 'yes', what was the most recent LVEF test type?		
Validation Rules	Field must be blank if LVEF Test Performed = No. Field must be complete if LVEF Test Performed = Yes.		
Maximum Field Size	1		
Related Data Field	eftp		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if eftp=0 and eft has data		
	Reject file if eftp =1 and eft is NULL		

3.4.4. Ejection Fraction (EF) Test Result Digitally Derived

Defining Attributes

Name Ejection Fraction (EF) Test Result Digitally Derived %

Field Name ef

Definition The digitally derived EF result (where the EF test result is a computed, digital EF percentage).

Note: The patient's digitally derived EF result should ONLY be expressed as a whole number (between 10 and 80)

expressed as a percentage of blood emptied from the left ventricle at the end of contraction.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always, if LVEF Test Performed = Yes, and Ejection Fraction (EF) Test Result Estimated = Blank

Representational Attributes

Permitted Values Must be reported as a percentage (%). If a percentage range was reported, report the lowest number of the range BELOW

(i.e., 50-55%, is reported as 50%).

Range 01-99

Guide for Use Where LVEF test performed is coded 'yes', what was the digitally derived EF test result? Only answer digitally derived or

estimated not both.

Validation Rules Field must be blank if LVEF Test Performed = No. Field must be blank if Ejection Fraction Test Result Estimated is complete.

Field must be complete if Ejection Fraction Test Result Estimated is blank, and LVEF Test Performed = Yes.

Maximum Field Size 2
Related Data Field eftp

Data Type Number (%)

Parent / Child Child

Behaviour Reject file if eftp=0 and ef has data

Reject file if ef has data & efes has data

Reject file if eftp =1 and both efes and ef are NULL

3.4.5. Ejection Fraction (EF) Test Result Estimated

Defining Attributes

Name Ejection Fraction (EF) Test Result Estimated

Field Name efect

Definition The patient's ejection fraction (EF) is the amount of blood emptied from the left ventricle at the end of contraction. EF

is estimated where the test itself has not computed a digital EF percentage to express ventricular function.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Always, if LVEF Test Performed = Yes, and Ejection Fraction (EF) Test Result Digitally Derived % = No.

Obligation	, ,	Et lest ellottied les, and Ejection l'action (Et) lest nesalt digitally delived 70 No.	
Representational Attributes			
Permitted Values	Code	Description	
	1	Normal Function (>50%)	
	2	Mild Dysfunction/Impairment (45-49%)	
	3	Moderate Dysfunction/Impairment (35-44%)	
	4	Severe Dysfunction/Impairment (<35%)	
Range	1-4		
Guide for Use	Where LVEF test performed is coded 'yes', what was the estimated EF test result? Only answer digitally derived or estimated not both.		
Validation Rules	Field must be blank if Ejection Fraction (EF) Test Digitally Derived is complete. Field must be complete if LVEF test performed = Yes. Field must be blank If LVEF test performed = No.		
Maximum Field Size	1		
Related Data Field	eftp		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if eftp=0 and efes has data		
	Reject file if ef has data & efes has data		
Reject file if eftp =1 and both efes and ef are NULL		eftp =1 and both efes and ef are NULL	

3.4.6. Estimated Glomerular Filtration Rate (Cockroft-Gault formula)

Defining Attributes

Name Estimated Glomerular Filtration Rate (Cockroft-Gault formula)

Field Name egfr

Definition An overall measure of kidney function, the glomerular filtration rate measures how well kidneys filter the wastes from

the blood. It is calculated using age, weight (in kg) and last pre-procedure creatinine.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Derived Variable

Representational Attributes

Permitted Values NNN.NN Range 1-999.00

Guide for Use

Validation Rules Estimated Glomerular Filtration Rate = (140 – age) x Weight x Constant / Pre-Procedure Creatinine (in μmol/L)

[Constant = 1.23 for men and Constant=1.04 for women]

Maximum Field Size

Related Data Field

Data Type Number (include 2 decimal places)

5

Parent / Child

Behaviour Derived – no import

3.4.7. Estimated Glomerular Filtration Rate Imported

Defining Attributes

Name Estimated Glomerular Filtration Rate Imported

Field Name egfri

Definition Where a hospital system calculates an egfr this can be included here as per the hospital's definition.

Justification Collected to determine the patient's pre-existing risk profile and risk adjustment models.

Obligation Optional

Representational Attributes

Permitted Values NNN.NN Range 1.00-999.00

Guide for Use What was the hospital system calculated EGFR? As per hospital definition.

Validation Rules

Maximum Field Size 5

Related Data Field

Data Type Number (include 2 decimal places)

Parent / Child Behaviour

3.5. Clinical Presentation

3.5.1. Cardiogenic Shock

Defining Attributes

Name Cardiogenic Shock

Field Name shock

Definition Indicate if the patient was in cardiogenic shock at the time of index PCI. Cardiogenic supporting definition: Transient

episodes of hypotension reversed with IV fluid or atropine do not constitute cardiogenic shock. Cardiogenic shock is coded as 'yes' if all of the following apply: 1. Sustained (>30 minutes) episode of systolic blood pressure <90 mm Hg (or vasopressors required to maintain BP >90 mm Hg); AND 2. Evidence of elevated filling pressures (e.g. pulmonary congestion on examination or chest radiograph); AND 3. Evidence of end organ hypoperfusion (e.g. urine output

30mL/hour; or cold/diaphoretic extremities; or altered mental status, etc.).

Justification Collected to determine patient risk profile and risk adjustment models.

Obligation Always

Representational Attributes
Permitted Values
Code
Description

0 No 1 Yes

Range 0-1

Guide for Use Was the patient in cardiogenic shock at the time of index PCI?

Validation Rules Field cannot be blank.

Maximum Field Size 1

Related Data Field

Data Type Number

Parent / Child

3.5.2. Out-of-Hospital Cardiac Arrest (OHCA)

Defining Attributes

Name Out-of-Hospital Cardiac Arrest

Field Name oca

Definition The patient has experienced an out of hospital cardiac arrest (i.e. the lack of effective cardiac output) including if the

person was under cardiac arrest at the time of presentation to the hospital.

Justification Collected to determine patient risk profile and risk adjustment models.

Obligation Always

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Has the patient had an Out-of-Hospital Cardiac Arrest?

Validation Rules Field cannot be blank.

Maximum Field Size 1

Related Data Field

Data Type Number

Parent / Child

3.5.3. Pre-Procedural Intubation

Defining Attributes

Name Pre-Procedure Intubation

Field Name pint

Definition The patient received intubation prior to the PCI procedure.

Justification Collected to identify high risk patients.

Obligation Always

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Was the patient intubated before the PCI procedure commenced?

Note: Intubation during the procedure should be coded under "Procedural intubation required: pintr"

Validation Rules Field cannot be blank.

Maximum Field Size 1

Related Data Field

Data Type Number

Parent / Child

4. Procedure

4.1. Clinical Symptoms

4.1.1. ACS

Defining Attributes

Name Acute Coronary Syndrome (ACS)

Field Name acs

Definition ACS encompasses clinical features comprising chest pain or overwhelming shortness of breath, defined by accompanying

clinical, ECG and biochemical features. Specifically, ACS refers to unstable angina, non-ST-Elevation Myocardial Infarction

(NSTEMI) and/or ST-Elevation Myocardial Infarction (STEMI).

Justification Collected to determine patient risk profile and risk adjustment models.

Obligation Always

Representational Attributes
Permitted Values
Code
Description
O
No

1 Yes

Range 0-1

Guide for Use Was the patient suffering from an acute coronary syndrome (ACS) in the 7-day period leading up to and including the

index PCI?

Validation Rules Field cannot be blank.

Maximum Field Size

Related Data Field

Data Type Number Parent / Child Parent

4.1.2. Date of ACS Symptom

Defining Attributes

Name Date of ACS Symptom Onset

Field Name dso

Definition In the event of stuttering symptoms, ACS symptom onset is the date when symptoms became constant in quality or

intensity. ACS symptoms may include: tightness, pressure, heaviness, fullness or squeezing in the chest which may spread to the neck and throat, jaw, shoulders, back, upper abdomen, either or both arms and even into the wrists and hands, dyspnoea, nausea/vomiting, cold sweat or syncope. Seeking medical attention could include the person presenting to their GP who then refers them to hospital or the person presenting directly to hospital (via ambulance or own form of transport). If the person is already a patient at the hospital for another reason then the time recorded would be when they advised hospital staff of their symptoms. Medical attention is defined as either at hospital or from a general

practitioner. Restrict coding to 7 days prior to current procedure.

Justification Collected to determine if patient is an inpatient at the time of ACS, and to benchmark performance and timeliness of

treatment.

Obligation Always, if ACS = Yes

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

[Date of Procedure]-7 Days — [Date of Procedure]

Guide for Use Where item Acute Coronary Syndrome = Yes, on what date did the ACS symptoms start that prompted them to seek

medical treatment?

Validation Rules Field cannot be blank if ACS = Yes. Date of symptom onset must be ≤Date of Procedure, and >[Date of Procedure-7

Days]

Maximum Field Size 10 Related Data Field acs

Data Type Date/Time
Parent / Child Child

Behaviour Reject file if dso is < dop-7 Reject file if dso is > dop

Reject file if acs=1 and dso is NULL

4.1.3. Time of ACS Symptom Onset

Defining Attributes

Name Time of ACS Symptom Onset

Field Name tso

Definition The time at which a person experienced ACS symptoms that prompted them to seek medical attention (on the date

outlined in 'Date of ACS symptom onset' in 24-hour clock (HH:MM) format. If the symptom onset time is not specified in the medical record, it may be recorded as 0700 for morning; 1200 for lunchtime; 1500 for afternoon; 1800 for

dinnertime; 2200 for evening and 0300 if awakened from sleep

Justification Collected to determine if patient is an inpatient at the time of ACS, and to benchmark performance and timeliness of

treatment.

Obligation Always, if ACS = Yes

Representational Attributes

Permitted Values HH:MM Range 00:00-23:59

Guide for Use Where item Acute Coronary Syndrome = Yes, at what time did the ACS symptoms start that prompted them to seek

medical treatment?

Validation Rules Field cannot be blank if ACS = Yes and Onset Time Not Available is blank. Field cannot be complete if Onset Time Not

Available = Yes.

Maximum Field Size 5
Related Data Field acs

Data Type Date/Time Parent / Child Child

Behaviour Reject file if acs=1 and tso AND ntso is NULL

Reject file if tso has data and ntso =1

4.1.4. Onset Time Not Available

Defining Attributes

Onset Time Not Available Name

Field Name ntso

Definition Where time of symptom onset is not available, specify here. Only applicable if time is missing.

Justification Used to determine that the ACS onset time is not available.

Obligation Always, if ACS Symptom Onset Time Not Available

Representational Attributes

Permitted Values Code Description 1

Yes

Range

Guide for Use Is the time of symptom onset not available?

Validation Rules Field cannot be complete if Time of ACS Symptom Onset is complete.

Maximum Field Size Related Data Field tso Data Type Number Parent / Child Child

Reject file if tso has data and ntso =1 Behaviour

4.1.5. Type of ACS

Defining Attributes

Definition

Name Type of ACS

Field Name acst

Unstable Angina: Symptoms must include at least one of the following: 1. Angina that occurred at rest and was prolonged, usually lasting >20 mins; 2. New-onset angina of at least CCS class III severity; 3. Recent acceleration of angina reflected by an increase in severity of at least 1 CCS class (to at least CCS class III).

Non ST-Elevation Myocardial Infarction (NSTEMI): <u>At least one</u> of the following biomarkers for detecting myocardial necrosis <u>must</u> be present: 1. Troponin T or I: Maximal concentration of Troponin T or I greater than the MI diagnostic limit on at least one occasion within 24 hours from the index clinical event; 2. CK-MB: Maximal value of CK-MB >2x the upper limit of normal (ULN) on one occasion during the first hours after the index clinical event; OR Maximal value of CK-MB (preferable CK-MB mass) > ULN on two successive samples. 3. Total CK: Only where Troponin or CK-MB assays are unavailable, total CK >2x the ULN (or the B fraction of CK) may be employed.

Note: The preferred assay to use as biomarkers for myocardial necrosis is Troponin. In the absence of troponin, CK-MB is the next best alternative but total CK can be used in the absence of troponin and CK-MB and one of the following: 1. Either ST segment depression or T wave abnormalities in the ECG; or 2. Ischaemic symptoms in the presence or absence of chest discomfort. Ischaemic symptoms may include: Unexplained nausea and vomiting; or Persistent shortness of breath secondary to left ventricular failure; or Unexplained weakness, dizziness, light-headedness, or Syncope.

ST-Elevation Myocardial Infarction (STEMI): At least one of the following biomarkers for detecting myocardial necrosis MUST be present (refer to Note regarding Reference Control Limits): 1. Troponin T or I: Maximal concentration of Troponin T or I > the MI diagnostic limit on at least one occasion within 24 hours from the index clinical event; 2. CK-MB: Maximal value of CK-MB >2x the upper limit of normal (ULN) on one occasion during the first hours after the index clinical event; OR Maximal value of CK-MB (preferable CK-MB mass) > ULN on two successive samples. 3. Total CK: Only where Troponin or CK-MB assays are unavailable, total CK >2x the ULN (or the B fraction of CK) may be employed. Note: The preferred assay to use as biomarkers for myocardial necrosis is Troponin. In the absence of troponin, CK-MB is the next best alternative but total CK can be used in the absence of troponin and CK-MB and one of the following ECG changes: 1. ST-segment elevation: New or presumed new ST segment elevation at the J point in two or more contiguous leads with the cut-off points \geq 0.2mV in leads V1, V2, or V3, or \geq 0.1 mV in other leads. 2. Development of any Q wave in leads V1 through V3, or the development of a Q-wave \geq 30ms (0.03s) in leads I, II, aVL, aVF, V4, V5, or V6. (Q wave changes must be present in any two contiguous leads, and be \geq 1mm in depth).

Collected to assess clinical appropriateness of procedure.

Always, if ACS = Yes

Justification Obligation

Permitted Values Code Description

1 Unstable Angina

2 NSTEMI 3 STEMI

Range 1-3

Guide for Use What is the type of ACS? Validation Rules Field cannot be blank.

Maximum Field Size 1
Related Data Field acs
Data Type Number
Parent / Child Child

Behaviour Reject file if acs=1 and acst is NULL

4.1.6. Inter-Hospital Transfer

Defining Attributes

Name Inter-Hospital Transfer

Field Name iht

Definition The patient was admitted to the PCI hospital following transfer from another acute care facility.

Justification Collected to allow reporting of NCR QI – Time from door to PCI-mediated reperfusion

Obligation Always, if Type of ACS = 3, STEMI

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Where patient 'Type of ACS' coded as 'STEMI', was the patient transferred from another acute care facility?

Validation Rules Field cannot be left blank if ACS type = STEMI

Maximum Field Size 1
Related Data Field acst
Data Type Number
Parent / Child Child

Behaviour Reject file if acst=3 and iht is NULL

4.1.7. Pre-Hospital Notification

Defining Attributes

Name Pre-Hospital Notification

Field Name phn

Definition PCI hospital was notified via electronic ECG transmission (or equivalent) from a paramedic field triage service prior to

their arrival at the PCI hospital.

Justification Collected to allow reporting of NCR QI – Time from door to PCI-mediated reperfusion.

Obligation Always, if Type of ACS = 3, STEMI

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Where patient 'Type of ACS' coded as 'STEMI', Indicate if the PCI hospital was notified via electronic ECG transmission

(or equivalent) from a paramedic field triage service prior to their arrival at the PCI hospital.

Validation Rules Field cannot be left blank if ACS type = STEMI

Maximum Field Size 1
Related Data Field acst
Data Type Number
Parent / Child Child

Behaviour Reject file if acst=3 and phn is NULL

4.1.8. Balloon/Device Date

Defining Attributes

Name Balloon/Device Date

Field Name dbd

Definition The date the first device was used to treat the target lesion. Examples of the first device used include, but are not

limited to: balloon; thrombectomy device; atherectomy device or stent. If the lesion cannot be crossed by the

guidewire or device, use the time that the guide catheter was introduced.

Note: This is a process measure, not a clinical outcomes measure and, as such, is not related to the timing of or

whether Thrombolysis in Myocardial Infarction (TIMI) 3 (Complete Perfusion) flow was/was not restored.

Justification Collected to allow reporting of NCR QI – Time from door to PCI-mediated reperfusion and Time from diagnostic

electrocardiogram to PCI mediated reperfusion.

Obligation Derived Variable

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use Where patient 'Type of ACS' coded as 'STEMI', on what date was the first device used?

Validation Rules If [Balloon/Device Time ≤ PCI Procedure Time] then Balloon/Device Date = PCI Procedure Date + [1 day]

If [Balloon/Device Time > PCI Procedure Time] then Balloon/Device Date = PCI Procedure Date

Maximum Field Size 10
Related Data Field acst
Data Type Date/Time

Parent / Child Child

Behaviour If [tbd=<top] then dbd=dop + [1 day]

If [tbd>top] then dbd=dop

4.1.9. Balloon/Device Time

Defining Attributes

Name Balloon/Device Time

Field Name tbd

Definition The time the first device was used to treat the target lesion. Examples of the first device used include, but are not

limited to: balloon; thrombectomy device; atherectomy device or stent. If the lesion cannot be crossed by the

guidewire or device, use the time that the guide catheter was introduced.

Note: This is a process measure, not a clinical outcomes measure and, as such, is not related to the timing of or

whether Thrombolysis in Myocardial Infarction (TIMI) 3 (Complete Perfusion) flow was/was not restored.

Justification Collected to allow reporting of NCR QI – Time from door to PCI-mediated reperfusion and Time from diagnostic

electrocardiogram to PCI mediated reperfusion.

Obligation Always, if Type of ACS = 3, STEMI

Representational Attributes

Permitted Values HH:MM Range 00:00-23:59

Guide for Use Where 'Type of ACS' coded as 'STEMI', at what time was the first device used?

Validation Rules Field cannot be left blank if ACS type = STEMI

Maximum Field Size 5
Related Data Field acst

Data Type Date/Time
Parent / Child Child

Behaviour Reject file if acst=3 and tbd is NULL

4.1.10. Symptom Onset to Reperfusion

Defining Attributes

Name Symptom Onset to Reperfusion

Field Name sor

Definition Time between symptom onset, and PCI mediated reperfusion.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Derived Variable

Representational Attributes

Permitted Values NNNN Range 0-9999

Guide for Use What is the time between Symptom Onset and Reperfusion of the Occluded Vessel?

Validation Rules Symptom Onset to Reperfusion = [Balloon/Device Time – Time of ACS Symptom Onset] + [Balloon/Device Date – Date of

ACS Symptom Onset]

Maximum Field Size

Related Data Field

Data Type Integer (Duration in Minutes)

4

Parent / Child

Behaviour sor = [tbd-tso] + [dbd-dso] (expressed in minutes)

4.1.11. Door to Balloon/Device Time

Defining Attributes

Name Door to Balloon/Device Time

Field Name dbdt

Definition Time between a patient arriving at the PCI hospital to inflation of the balloon/device insertion.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Derived Variable

Representational Attributes

Permitted Values NNNN Range 0-9999

Guide for Use What is the time between the patient's entry into the hospital's administrative system and the first device used to treat

the target lesion?

Validation Rules Door to Balloon Time = [Balloon/Device Time - Time of Arrival at PCI Hospital] + [Balloon/Device Date = Date of Arrival at

PCI Hospital]

Maximum Field Size 4
Related Data Field acst

Data Type Integer (Duration in Minutes)

Parent / Child Child

Behaviour dbdt = [tbd-toa] + [dbd-doa]

4.1.12. Self-Presenter

Defining Attributes

Name Self-Presenter

Field Name spr

Definition The patient presented to hospital by way of their own means e.g., not in an ambulance or other emergency vehicle.

Justification Collected to assess method of presentation to hospital

Obligation Always, if Type of ACS = 3, STEMI

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Where patient 'Type of ACS' coded as 'STEMI', did the patient present to the hospital without an ambulance?

Validation Rules Field cannot be left blank if ACS type = STEMI

Maximum Field Size 1
Related Data Field acst
Data Type Number
Parent / Child Child

Behaviour Reject file if acst=3 and spr is NULL

4.1.13. Date of First Medical Contact

Defining Attributes

Name Date of First Medical Contact

Field Name dfmc

Definition The date on which the patient first received medical attention (either by ambulance arriving, attendance at hospital or

from a general practitioner) for their ACS symptoms.

Note: Seeking medical attention could include the person presenting to their GP who then refers them to hospital or the person presenting directly to hospital, or the arrival of the ambulance service. If the person is already a patient at

the hospital for another reason then the date recorded would be when they advised hospital staff of their symptoms.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Always, if Type of ACS = 3, STEMI

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use Where 'Type of ACS' coded as 'STEMI', what was the date of First Medical Contact?

Validation Rules Field cannot be blank if Type of ACS = STEMI. Date of First Medical Contact cannot > Date of Procedure, Date of Symptom

Onset +7, Date of Diagnostic ECG. Date of First Medical Contact cannot < Date of Symptom Onset.

Maximum Field Size 10 Related Data Field acst

Data Type Date/Time (Minutes)

Parent / Child Child

Behaviour Reject file if acst = 3 and dfmc is NULL

Reject file if dfmc>dop Reject file if dfmc>dso+7 Reject file if dfmc<dso Reject file if dfmc>decgd

4.1.14. Time of FMC

Defining Attributes

Name Time of First Medical Contact

Field Name tfmc

Definition The time at which the patient first received medical attention (either by ambulance arriving, attendance at hospital or

from a general practitioner) for their ACS symptoms.

Note: Seeking medical attention could include the person presenting to their GP who then refers them to hospital or the person presenting directly to hospital, or the arrival of the ambulance service. If the person is already a patient at

the hospital for another reason then the time recorded would be when they advised hospital staff of their symptoms.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Always, if Type of ACS = 3, STEMI

Representational Attributes

Permitted Values HH:MM Range 00:00-23:59

Guide for Use Where 'Type of ACS' coded as 'STEMI', what was the time of First Medical Contact?

Validation Rules Field cannot be blank if Type of ACS = STEMI. Time of First Medical Contact cannot < Time of Symptom Onset when Date

of First Medical Contact = Date of Symptom Onset. Time of First Medical Contact cannot > Time of Procedure when Date

of First Medical Contact = Date of Procedure

Maximum Field Size 5

Related Data Field acst
Data Type Date/Time

Parent / Child Child

Behaviour Reject file if acst = 3 and tfmc is NULL

Reject file when tfmc<tso and dfmc=dso Reject file when tfmc>top and dfmc=dop

4.1.15. Symptom Onset to FMC

Defining Attributes

Name Symptom Onset to FMC

Field Name sofmc

Definition Derived variable to determine the time between symptom onset and first medical contact.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Derived Variable

Representational Attributes

Permitted Values NNNNN Range 0-99999

Guide for Use

Validation Rules Symptom Onset to First Medical Contact = [Time of First Medical Contact – Time of Symptom Onset] + [Date of First

Medical Onset- Date of Symptom Onset]

Maximum Field Size

Related Data Field

Data Type Integer (Duration in Minutes)

5

Parent / Child

Behaviour sofmc= [tfmc-tso] + [dfmc-dso]

4.1.16. FMC to Reperfusion

Defining Attributes

Name FMC to Reperfusion

Field Name fmctr

Definition Derived variable to determine the time between first medical contact to reperfusion.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Derived Variable

Representational Attributes

Permitted Values NNNN Range 0-9999

Guide for Use Derived variable to determine the time between first medical contact to reperfusion.

Validation Rules First Medical Contact to Reperfusion = [Balloon/Device Time – Time of First Medical Contact] + [Balloon/Device Date –

Date of First Medical Contact]

Maximum Field Size

Related Data Field

Data Type Integer (Duration in Minutes)

Parent / Child

Behaviour fmctr= [tbd-tfmc] + [dbd-dfmc]

4.1.17. FMC to Door

Defining Attributes

Name FMC to Door

Field Name fmctd

Definition Derived variable to determine the time between first medical contact to hospital door.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Derived Variable

Representational Attributes

Permitted Values NNNN Range 0-9999

Guide for Use Derived variable to determine the time between first medical contact to reperfusion.

Validation Rules First Medical Contact to Door= [Time of Arrival -Time of First Medical Contact] + [Date of Arrive – Date of First Medical

Contact]

Maximum Field Size

Related Data Field

Data Type Integer (Duration in Minutes)

4

Parent / Child

Behaviour fmctd= [toa-tfmc] + [doa-dfmc]

4.1.18. Date of Diagnostic ECG

Defining Attributes

Name Date of Diagnostic ECG

Field Name decgd

Definition The date of the diagnostic electrocardiogram (ECG). The ECG recorded that shows evidence of ECG abnormalities

consistent with a STEMI or suspected STEMI event.

Note: The diagnostic ECG may have been before admission to this hospital.

Justification Collected to allow reporting of NCR QI – Time from diagnostic electrocardiogram to PCI mediated reperfusion.

Obligation Always, if ACS Type = STEMI

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use Where 'Type of ACS' coded as 'STEMI', what was the date of diagnostic ECG?

Validation Rules Field cannot be blank if Type of ACS = STEMI

Maximum Field Size 10 Related Data Field acst

Data Type Date/Time Parent / Child Child

Behaviour Reject file if acst = 3 and decgd is NULL

4.1.19. Time of Diagnostic ECG

Defining Attributes

Name Time of Diagnostic ECG

Field Name tecgd

Definition The time of the diagnostic ECG. The ECG recorded that shows evidence of ECG abnormalities consistent with a STEMI

or suspected STEMI event.

Note: The diagnostic ECG may have been before admission to this hospital.

Justification Collected to allow reporting of NCR QI – Time from diagnostic electrocardiogram to PCI mediated reperfusion.

Obligation Always, if ACS Type = STEMI

Representational Attributes

Permitted Values HH:MM Range 00:00-23:59

Guide for Use Where 'Type of ACS' coded as 'STEMI', what was the time of diagnostic ECG?

Validation Rules Field cannot be blank if ACS Type = STEMI. Time of Diagnostic ECG cannot > PCI Procedure Time when Date of Diagnostic

 $\mathsf{ECG} = \mathsf{Date} \ of \ \mathsf{PCI} \ \mathsf{Procedure}. \ \mathsf{Time} \ of \ \mathsf{Diagnostic} \ \mathsf{ECG} \ \mathsf{cannot} \\ \leq \mathsf{Time} \ of \ \mathsf{ACS} \ \mathsf{Symptom} \ \mathsf{Onset} \ \mathsf{when} \ \mathsf{Date} \ \mathsf{of} \ \mathsf{Diagnostic} \ \mathsf{ECG} \ \mathsf{Constant} \\ \mathsf{Diagnostic} \ \mathsf{$

– Date of Symptom Onset. Time of Diagnostic ECG cannot ≤ Time of First Medical Contact when Date of Diagnostic ECG

= Date of First Medical Contact

Maximum Field Size 5

Related Data Field acst

Data Type Date/Time
Parent / Child Child

Behaviour Reject file if acst = 3 and tecgd is NULL

Reject file when tecgd > top and decgd = dop Reject file when tecgd ≤ tso and decgd=dso Reject

file when tecgd ≤ tfmc when decgd = dfmc

4.1.20. FMC to Diagnostic ECG

Defining Attributes

Name FMC to Diagnostic ECG

Field Name fmctecg

Definition The time between First Medical Contact to Diagnostic ECG

Note: The diagnostic ECG may have been before admission to this hospital.

Justification Collected to benchmark performance and timeliness of treatment.

Obligation Derived Variable

Representational Attributes

Permitted Values NNN Range 0-100

Guide for Use

Validation Rules First Medical Contact to Diagnostic ECG = [Time of Diagnostic ECG – Time of First Medical Contact] + [Date of Diagnostic

ECG – Date of First Medical Contact] (expressed in minutes)

Maximum Field Size

Related Data Field

Data Type Integer (Duration in Minutes)

3

Parent / Child

Behaviour fmctecg= [tecgd-tfmc] + [decgd-dfmc] (expressed in minutes)

4.1.21. Diagnostic ECG to Balloon

Defining Attributes

Name Diagnostic ECG to Balloon

Field Name ecgdb

Definition The time between Diagnostic ECG and inflation of Balloon/Device

Note: The diagnostic ECG may have been before admission to this hospital

Justification Collected to allow reporting of NCR QI – Time from diagnostic ECG to PCI-mediated reperfusion

Obligation Derived Variable

Representational Attributes

Permitted Values NNN Range 0-100

Guide for Use

Validation Rules Diagnostic ECG to Balloon= [Time of Diagnostic ECG – Balloon/Device Time] + [Date of Diagnostic ECG – Date of

Balloon/Device]

Maximum Field Size

Related Data Field

Data Type Integer (Duration in Minutes)

3

Parent / Child

Behaviour ecgdb = [tecgd-tbd] + [decgd-dbd] (expressed in minutes)

4.1.22. Inpatient at Time of ACS

Defining Attributes

Name Inpatient at Time of ACS

Field Name inp

Definition Was the patient an inpatient in the hospital at the Time of ACS onset?

Justification Collected to allow reporting of NCR QI — Time from door to PCI-mediated reperfusion

Obligation Derived Variable

Representational Attributes Values Code Description 0 No 1 Yes 0-1 Range Guide for Use Where patient 'Type of ACS' coded as 'STEMI', did the patient present to the hospital without an ambulance? Validation Rules Inpatient = No if ACS = Yes & [Date of Symptom Onset & Time of Symptom Onset] ≤ [Date of Arrival & Time of Arrival] Inpatient = Yes if ACS = Yes & [Date of Symptom Onset & Time of Symptom Onset] > [Date of Arrival & Time of Arrival] Inpatient = NULL if ACS = 0 Maximum Field Size 1

Maximum Field Size 1
Related Data Field acs
Data Type Number
Parent / Child Child

4.2. PCI Indication

Defining Attributes

Name Field Name

Definition

PCI Indication

pci

- 1. Primary PCI for patient with STEMI (or STEMI equivalent) within 12 hours of symptom onset.
- 2. PCI for patient with STEMI (or STEMI equivalent) between 12-24 hours of symptom onset
- 3. Rescue PCI for STEMI (or STEMI equivalent) after a failed full-dose lysis
- 4. PCI for STEMI (or STEMI equivalent) where the patient is stable after receiving fill-dose lysis
- 5. PCI for STEMI (1-7 days no prior lysis)
- 6. PCI for STEMI (1-7 days following lysis)
- 7. PCI for cardiac arrest/cardiogenic shock (Not STEMI) PCI performed post-cardiac arrest or cardiogenic shock but without ECG or biomarker evidence of acute myocardial infarction.

Note: This does not include any PCI cases for Myocardial Infarction (MI).

- 8. PCI for NSTEMI, including patients who have high risk features for short-term risk of death or nonfatal MI. High risk features include at least one of the following:
 - History accelerating tempo of ischaemic symptoms in preceding 48 hours
 - Character of pain prolonged ongoing (greater than 20 minutes) rest pain
 - Clinical findings: Pulmonary oedema, most likely due to ischemia
 - New or worsening mitral regurgitation murmur
 - S3 or new worsening crackles/crepitations
 - Hypotension, bradycardia, tachycardia
 - Age greater than 75 years
 - ECG: Angina at rest with transient ST-segment changes greater than 0.5 mm
 - Bundle-branch block, new or presumed new
 - Sustained ventricular tachycardia.
 - Cardiac markers NSTEMI patients with elevated cardiac Tn-T, Tn-I, or CK-MB.
 - 9. PCI for unstable angina (<7 days) Includes patients with unstable angina who have high risk features for short-term risk of death or nonfatal MI.

High risk features include at least one of the following

- History accelerating tempo of ischaemic symptoms in preceding 48 hours
- Character of pain prolonged ongoing (greater than 20 minutes) rest pain
- Clinical findings pulmonary oedema, most likely due to ischaemia

Justification Collected to allow reporting of NCR QI – Time from door to PCI-mediated reperfusion and procedures are undertaken for appropriate reasons and for clinical presentation reporting.

Obligation Always

Obligation	Aiways	
Representational Attributes		
Permitted Values	Code	Description
	1	Primary PCI for STEMI<12 hours
	2	PCI for STEMI (12-24 hours)
	3	Rescue PCI for STEMI (<24 hours – unstable following lysis)
	4	PCI for STEMI (<24 hours – stable following lysis)
	5	PCI for STEMI (1-7 days no prior lysis)
	6	PCI for STEMI (1-7 days following lysis)
	7	PCI for cardiac arrest/cardiogenic shock (without evidence of STEMI)
	8	PCI for NSTEMI (<7 days)
	9	PCI for unstable angina (<7 days)
	10	PCI for recent ACS (7-30 days)
	11	Staged PCI
	12	Angina/angina equivalent symptoms
	13	No angina/angina equivalent symptoms
Range	1-13	
Guide for Use	What was the	reason for the current PCI?
Validation Rules	PCI Indication cannot = Primary PCI for STEMI 12 hours, PCI for STEMI (12-24 hours), Rescue PCI for STEMI (<24 hours) unstable following lysis), PCI for STEMI (<24 hours – stable following lysis), PCI for STEMI (1-7 days no prior lysis) or for STEMI (1-7 days following lysis) if Acute Coronary Syndrome = No. PCI Indication cannot = PCI for cardiac arrest/cardiogenic shock (without evidence of STEMI), PCI for NSTEMI (<7 days), PCI for unstable angina (<7 days), PCI for recent ACS (7-30 days), Staged PCI, Angina/angina equivalent symptoms, angina/angina equivalent symptoms if Type of ACS = STEMI. PCI Indication must = 8, if Type of ACS = NSTEMI.	
	PCI Indication	must = 9 if Type of ACS = Unstable Angina
Maximum Field Size Related Data Field	2	
Data Type Parent / Child	Number	
Behaviour	Reject file if ac	s = 0 and pci = 1, 2, 3, 4, 5 or 6.

Reject file if acst=3 and pci = 7, 8, 9, 10, 11, 12 or 13. Reject file if acst = 2 and pci does not =8 Reject file is acst=1 and pci does not =9 Reject file if NULL

4.3. Procedure Details

4.3.1. Percutaneous Entry Location

Defining Attributes

Name Percutaneous Entry Location

Field Name pel

Definition The percutaneous entry location used to provide arterial vascular access for the procedure.

Note: If crossover occurred, select the percutaneous entry location that successfully facilitated the index PCI

procedure.

Justification Collected for use in risk adjustment models and trends of best practice.

0.0.1641.011	,	
Representational Attributes		
Permitted Values	Code	Description
	1	Brachial
	2	Radial
	3	Femoral
Range	1-3	
Guide for Use	Which entry l	location was used to provide access for the procedure?
Validation Rules	Field cannot be blank.	
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		
Behaviour	Reject file if N	NULL
Additional Information		
References		
Related Metadata		

4.3.2. Procedural Intubation Required

Defining Attributes

Name Procedural Intubation Required

Field Name pintr

Definition Intubation required during the index PCI procedure.

Justification Collected to identify high risk patients.

Obligation Always

Representational Attributes

Permitted Values Code Description

1

0 No 1 Yes

Range 0-1

Guide for Use Was the patient intubated during the procedure?

Validation Rules Field cannot be blank.

Maximum Field Size

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL

4.3.3. Mechanical Circulatory Support

Defining Attributes

Name Mechanical Circulatory Support

Field Name vrs

Definition Mechanical ventricular support required prior to or during the index PCI procedure.

Note: Mechanical ventricular support includes: intra-aortic balloon pump (IABP), cardiopulmonary bypass, left ventricular assist device (LVAD), extracorporeal membrane oxygenation (ECMO), mechanical chest compression device (e.g. Lund University Cardiac Arrest System - LUCAS) and/or catheter based cardiac assist device (e.g., IMPELLA heart

pump).

Justification Collected to identify high risk patients and risk adjustment models.

Obligation Always

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Was ventricular support required during the procedure?

Validation Rules Field cannot be blank.

Maximum Field Size 1

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL

4.3.4. Lesion Location

Defining	Attri	butes

Name Lesion Location Field Name lr1_lesion

Definition The coronary segment that applies for EACH coronary lesion attempted during the current PCI. Every coronary lesion attempted during the current PCI must be recorded separately (add new lesion). Up to 5 lesions per PCI can be

recorded. Select ONE segment for each lesion treated.

Justification Collected to determine the lesion treated.

Obligation	Aiways	
Representational Attributes		
Permitted Values	Code	Description
	1	Left anterior descending artery (LAD)
	2	Diagonal artery
	3	Left circumflex (LCx)
	4	Left main coronary artery
	5	Posterior descending artery
	6	Right coronary artery
	7	Internal mammary graft
	8	Radial artery graft
	9	Saphenous vein graft
	10	Not stated/inadequately described
Range	1-10	
Guide for Use	What was the l	esion location?
Validation Rules	Field cannot be	e blank.
Maximum Field Size	2	
Related Data Field		
Data Type	Number	
Parent / Child		
Behaviour	Reject file if N LR5_LESION.	ULL. Up to 5 lesions can be entered. Fields are coded in data extract as LR1_LESION, LR2_LESION

4.3.5. In-stent Restenosis (ISR)

Defining Attributes

Name In-stent Restenosis (ISR)

Field Name lr1_isr

Definition The current lesion location has had a pre-existing stent implanted from a prior PCI to the same site OR is within 5mm of

the proximal or distal stent edges of a pre-existing stent from a previous PCI.

Justification Collected to identify trends and safely and efficacy of the treatment of clinical subsets.

Obligation Always

Representational Attributes
Permitted Values
Code
Description

1

0 No 1 Yes

Range 0-1

Guide for Use

Does the current lesion location have a pre-existing stent implanted from a prior PCI to the same site?

Validation Rules

Field cannot be blank. Field cannot be coded 1=Yes if Previous PCI = No and In-stent Restenosis = Yes

Maximum Field Size

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL

Reject file if ppci =0 AND lr1_isr=1

Up to 5 lesions can be entered. Fields are coded in data extract as LR1 LESION, LR2 LESION ... LR5 LESION.

4.3.6. Stent thrombosis

Defining Attributes

Name Stent Thrombosis

Field Name lr1_isrst

Definition The current lesion was treated due to the presence of a thrombus in the pre-existing stent OR within 5mm of stent

edges.

Justification Collected to identify trends and safely and efficacy of the treatment of clinical subsets.

Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
Range	0-1		
Guide for Use	Was the curi	rent lesion treated due to the presence of a thrombus in a pre-existing stent?	
Validation Rules	Field cannot be blank. Field cannot be coded 1=Yes if Previous PCI = No and Stent Thrombosis = Yes		
Maximum Field Size	1		
Related Data Field			
Data Type	Number		
Parent / Child			
Behaviour	Reject file if	NULL	
	Reject file if	PPCI=0 AND Ir1_isrst=1	
	Up to 5 lesio	ons can be entered. Fields are coded in data extract as LR1_LESION, LR2_LESION LR5_LESION.	

4.3.7. Lesion Successfully Treated

Defining Attributes

Name Lesion Successfully Treated

Field Name Ir1_lst

Definition The lesion was successfully treated. Successful dilation of a lesion is considered where: Residual stenosis is less than

10% following coronary stenting; OR Residual stenosis is less than 50% after POBA (balloon angioplasty/atherectomy)

alone AND Coronary blood perfusion rates have reached TIMI 2 or TIMI 3 flow.

Note: Thrombolysis in Myocardial Infarction (TIMI) 2 flow (partial reperfusion) is evidenced by delayed or sluggish

antegrade flow with complete filling of the distal territory. TIMI 3 flow (complete perfusion) is normal flow which fills

the distal coronary bed completely.

Justification Collected to identify trends and safely and efficacy of the treatment of clinical subsets.

Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
Range	0-1		
Guide for Use	Was the lesion successfully treated?		
Validation Rules	Field cannot be blank. Field cannot be coded 1=Yes if Previous PCI = No and Stent Thrombosis = Yes		
Maximum Field Size	1		
Related Data Field			
Data Type	Number		
Parent / Child			
Behaviour	Reject file if I	NULL	
	Up to 5 lesio	ns can be entered. Fields are coded in data extract as LR1_LESION, LR2_LESION LR5_LESION.	

4.3.8. Type of Stent(s) Implanted

Defining Attributes

Name Type of Stent(s) Implanted

Field Name lr1_sit

Definition Select ONE of the following options: <u>Bare metal stents (BMS):</u> Where one or more bare metal coronary stent(s) were

implanted in the current lesion (and no other type of stent/scaffold was implanted). Drug eluting stents (DES): Where one or more drug eluting coronary stent(s) were implanted in the current lesion (and no other type of stent was implanted). Mixed stents (combined BMS & DES): Where at least one BMS and at least one DES were implanted in the current lesion. Bioresorbable stent(s) (BRS): Where one or more bioresorbable coronary stent(s) were implanted in the current lesion (and no other type of stent/scaffold was implanted). Other: Where stent(s) or scaffold(s) were implanted into the current lesion, but the available options listed above do not apply. This would include combinations of metal stents and BVS(s) whether they were drug eluting and/or non-drug eluting devices. Enter the details of the stent/scaffold in the free text provided. POBA (Plain Balloon Angioplasty): Where the only treatment for this lesion was balloon inflation to recanalise coronary arteries; this includes both balloons/catheters coated in a drug (e.g. paclitaxel, etc.) or NOT coated with any drug. No Stent/No Balloon.

Justification Collected to identify trends and safely and efficacy of the treatment of clinical subsets.

Representational Attributes		
Permitted Values	Code	Description
	1	Bare metal stent(s) only (BMS)
	2	Drug-eluting stent(s) (DES)
	3	Mixed stent(s) (Combined BMS & DES)
	4	Bioresorbable stent(s) (BRS)
	5	Other stents/scaffolds
	6	POBA (Plain Balloon Angioplasty)
	7	No Stent/No Balloon
Range	1-7	
Guide for Use	What type of	stent/balloon was used?
Validation Rules	Field cannot b	pe blank.
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		

Behaviour

Reject file if NULL

Up to 5 lesions can be entered. Fields are coded in data extract as LR1_LESION, LR2_LESION ... LR5_LESION.

4.3.9. Total Entrance Radiation Dose (Air Kerma)

Defining Attributes

Name Total Entrance Radiation Dose (Air Kerma)

Field Name tak

Definition The total radiation dose (Cumulative Air Kerma, or Reference Air Kerma) recorded to the nearest milligray (mGy). The

value recorded should include the total dose for the lab visit. Cumulative Air Kerma is the total Air Kerma accrued from

the beginning of an examination or procedure and includes all contributions from fluoroscopic and radiographic

irradiation.

Justification Collected to assess radiation risk

Obligation Optional

Representational Attributes

Permitted Values NNN Range 0-999

Guide for Use What was the total radiation dose?

Validation Rules Field cannot >999

Maximum Field Size 3

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if value entered >999.

4.3.10. Total Exam Dose Area Product in mGy/m2

Defining Attributes

Name Total Exam Dose Area Product in mGy/m2

Field Name dap

Definition The total fluoroscopy dose to the nearest integer. The value recorded should include the total dose for the lab visit.

Justification Collected to assess radiation risk.

Obligation Optional

Representational Attributes

Permitted Values NNN Range 0-999

Guide for Use What was the Dose Area Product?

Validation Rules Field cannot >999

Maximum Field Size 3

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if value entered >999.

4.3.11. Total Fluoroscopy Time

Defining Attributes

Name Total Fluoroscopy Time

Field Name tft

Definition The total fluoroscopy time in mins. Report to the nearest 0.1 min

Justification Collected to assess radiation risk.

Obligation Optional

Representational Attributes

Permitted Values NNN.N Range 0.1-300.0

Guide for Use What was the total fluoroscopy time?

Validation Rules Field cannot >300.0

Maximum Field Size

Related Data Field

Data Type Integer (Duration in Minutes)

Parent / Child

Behaviour Reject file if value entered >300.0

5. Post Procedure

5.1. Post-Procedure & In-Hospital Complications

5.1.1. In-Hospital MI

Defining Attributes

Name In-Hospital MI

Field Name ihmi

Definition The patient experienced the new presence of myocardial infarction (MI) either during the index cardiac catheter lab

visit, after the lab visit or any subsequent lab visits, but prior to being discharged from the hospital.

There may be some minor variation in the way participating registries define this item. Supporting definition: The inhospital, PCI-related MI must be a distinct clinical event that can be identified as a separate event to the index PCI. As such, the in-hospital new or recurrent MI will differ among patients presenting with normal cardiac biomarker levels and those patients whose pre-procedure baseline biomarker levels are elevated. 1. Patients with normal baseline biomarker levels, the NEW or recurrent MI must include: Elevated biomarkers* >5 the upper limit of normal (ULN) ≤48 hours of index PCI and at least one of the following: Evolutionary ST segment elevation (≥0.1mV) distinct from the index event; Development of new Q-waves (≥40ms duration) in 2 or more contiguous ECG leads; Ischaemic symptoms (with or without chest discomfort) lasting ≥20mins; Angiographic evidence flow limiting complications (embolism, persistent slow-flow or no re-flow, etc.); Imaging evidence of new loss of myocardium or new regional wall flow abnormality. Note: CABG-related peri-procedural MI (within 48 hours of CABG) requires CK-MB ≥10x ULN. 2. Patients with elevated baseline cardiac biomarker levels.

If cardiac biomarker levels are still rising (have not peaked) at the time of the index PCI procedure, an in-hospital NEW or recurrent MI cannot be diagnosed. For patients whose elevated cardiac biomarker level have peaked (no longer rising) by the current PCI procedure time, the in-hospital MI must include: A NEW elevation of biomarkers ≥20% from the pre-procedural cardiac biomarker levels and at least one of the following: Evolutionary ST segment elevation (≥0.1mV) distinct from the index event; Development of new Q-waves (≥40ms duration) in 2 or more contiguous ECG leads; Ischaemic symptoms (with or without chest discomfort) lasting >20mins; Angiographic evidence flow limiting complications (embolism, persistent slow-flow or no re-flow, etc.); Imaging evidence of new loss of myocardium or new regional wall flow abnormality.

Note: CABG-related peri-procedural MI (within 48 hours of CABG) requires CK-MB ≥10x ULN. * Troponin (T or I) is the preferred cardiac biomarker for defining the presence of MI. In the absence of Troponin, CK-MB is the best alternative (but total CK can be used, where CK-MB and Troponin are both unavailable).

Collected to allow reporting of in-hospital and 30-day MACE and MACCE.

Obligation Always

Justification

Representational Attributes

Permitted Values Code Description

0 No 1 Yes

Range 0-1

Guide for Use Did the patient experience a myocardial infarction while an in-patient?

Validation Rules Field cannot be blank.

Maximum Field Size 1

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if NULL

5.1.2. Subsequent PCI (in this admission)

Defining Attributes

Name Subsequent PCI (in this admission)

Field Name ihpci

Definition The patient underwent a subsequent PCI (distinct from the index PCI) after the cardiac catheter lab visit, but prior to

discharge. Note: This includes emergent PCI revascularisations and/or planned, staged PCI visits to the cardiac catheter

lab during the index admission, but prior to discharge.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation	Aiways		
Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
Range	0-1		
Guide for Use	Did the patient undergo a subsequent PCI during the same admission?		
Validation Rules	Field cannot be blank.		
Maximum Field Size	1		
Related Data Field			
Data Type	Number	Number	
Parent / Child	Parent		
Behaviour	Reject file if I	NULL	

5.1.3. Planned Subsequent PCI

Defining Attributes

Name Planned Subsequent PCI

Field Name ihpcip

Definition The subsequent PCI was planned. Note: A 'planned in-hospital PCI' is defined as a staged or scheduled elective PCI only.

For emergent PCI interventions that were not previously scheduled, code as 'no'.

Note: Any target vessel revascularisation of a successfully treated lesion should be coded as 'unplanned'. Subsequent

revascularisation of a previous unsuccessfully treated lesion is situation dependant.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation Always, if In-hospital PCI = Yes

Obligation	Aiways, ii iii ii	103pitari Ci – 1C3	
Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
Range	0-1		
Guide for Use	Where item S	Subsequent PCI was coded 'yes', was the subsequent PCI planned?	
Validation Rules	Field cannot k	be blank if In-hospital PCI = Yes. Field cannot be complete if In-hospital PCI = No.	

Maximum Field Size 1
Related Data Field ihpci
Data Type Number
Parent / Child Child

Behaviour Reject file if ihpci=0 and ihpcip has data Reject file if ihpci=1 and ihpcip is NULL

5.1.4. In-Hospital Target Vessel (PCI)

Defining Attributes

Behaviour

Name In-Hospital Target Vessel (PCI)

Field Name ihtvr

Definition The target vessel was revascularised during the subsequent in-hospital PCI.

Reject file if ihpci=0 and intvr has data

Reject file if ihpci =1 and ihtvr is NULL

Note: A PCI TVR is any revascularisation due to restenosis/occlusion within the target coronary artery and/or the same arterial branch that was treated during the index PCI. This includes any percutaneous revascularisation within the same

arterial branch treated during the index PCI, regardless of whether the index PCI was successful.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days

Obligation Always, if In-hospital PCI = Yes

Obligation	Aiways, II III-	Always, il ili-ilospitai r Ci – 1es		
Representational Attributes				
Permitted Values	Code	Description		
	0	No		
	1	Yes		
Range	0-1			
Guide for Use	Where item Subsequent PCI was coded 'yes', was the target vessel revascularised?			
Validation Rules	Field cannot be blank if In-hospital PCI = Yes. Field cannot be complete if In-hospital PCI = No.			
Maximum Field Size	1			
Related Data Field	ihpci			
Data Type	Number			
Parent / Child	Child			

5.1.5. In-Hospital Target Lesion Revascularisation

Defining Attributes

Name In-Hospital Target Lesion Revascularisation

Field Name ihtlr

Definition Indicate whether the target lesion was also revascularised during the subsequent in-hospital PCI.

Note: A target lesion revascularisation (TLR) is any revascularisation of the same lesion treated during the index PCI. This includes treatment of a restenosis/occlusion within a stent implanted during the index PCI (or within 5-mm of the proximal and distal margins of the stent edges). It also includes attempts to revascularise the target lesion where a stent was not successfully implanted and/or plain old balloon angioplasty (POBA) was employed during the index PCI.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation Always, if In-hospital Target Vessel (PCI) = Yes

Representational Attributes				
Permitted Values	Code	Description		
	0	No		
	1	Yes		
Range	0-1			
Guide for Use	Where item S	Where item Subsequent PCI was coded 'yes', was the target vessel revascularised?		
Validation Rules	Field cannot be blank if In-hospital PCI = Yes. Field cannot be complete if In-hospital PCI = No.			
Maximum Field Size	1			
Related Data Field	ihtvr			
Data Type	Number			
Parent / Child	Child			
Behaviour	Reject file if i	htvr=0 and ihtlr has data		
	Reject file if i	htvr=1 and ihtlr is NULL		

5.1.6. In-Hospital Cardiothoracic Surgery

Defining Attributes

Name In-Hospital Cardiothoracic Surgery

Field Name ihcab

Definition The patient underwent or was transferred for cardiothoracic surgery (whether or not this actually involved the placing

of bypass grafts) either during the cardiac catheter lab visit, after the lab visit, but prior to discharge and/or any

subsequent lab visits.

Note: The surgery should be cardiothoracic (i.e. not related to peripheral vascular complications). The surgical indication may have been, but are not exclusive to: Prompted/indicated by a need to correct an emergency complication of the index PCI (e.g. abrupt vessel closure, cardiac vessel perforation, dissection of a thoracic great vessel, etc.); OR CABG to revascularise the target vessel treated during the index PCI; either during or after the catheter lab visit, but prior to discharge and/or any subsequent lab visits; OR For a patient who underwent a successful PCI (e.g. for an ACS) but required a CABG for anatomical or pre-existing CAD indications (e.g. left-main disease, triple-

vessel disease, etc.).

Note: If the patient was transferred from the index hospital to a tertiary facility for cardiothoracic surgery as part of

ongoing treatment related to the index PCI (e.g. emergency CABG) then code as 'yes'.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation Always

Representational Attributes
Permitted Values
Code
Description

0 No 1 Yes

Range 0-1

Guide for Use Did the patient undergo or was transferred for cardiothoracic surgery prior to discharge?

Validation Rules Field cannot be blank.

Maximum Field Size 1

Related Data Field

Data Type

Parent / Child

Number Parent

Behaviour Reject file if NULL

5.1.7. Planned Cardiothoracic Surgery

Defining Attributes

Name Planned Cardiothoracic Surgery

Field Name ihpcab

Definition The cardiothoracic surgery was planned.

Note: A 'planned cardiothoracic surgery' is defined as a scheduled surgical event only. For emergent surgeries that

were not scheduled or planned, code as 'no'.

Note: Any target vessel revascularisation of a successfully treated lesion should be coded as 'unplanned'. Subsequent

revascularisation of a previous unsuccessfully treated lesion is situation dependant Please contact NCR project team if

unsure.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
Range	0-1		
Guide for Use	Where item	In-hospital cardiothoracic surgery was coded 'yes', was the cardiothoracic surgery was planned?	
Validation Rules	Field cannot be blank if In-hospital Cardiothoracic Surgery = Yes. Field cannot be complete if In-hospital Cardiothoraci		
	Surgery = No.		
Maximum Field Size	1		
Related Data Field	ihcab		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if ihcab=0 and ihpcab has data		
	Reject file if	ihcab=1 and ihpcab is NULL	

5.1.8. In-Hospital TVR (CABG)

Defining Attributes

Name In-Hospital TVR (CABG)

Field Name ihtvcab

Definition The surgery involved the placing of coronary artery bypass grafts to revascularise the target vessel that was treated

during the index PCI.

Note: A TVR CABG is a bypass revascularisation due to restenosis/occlusion within the target coronary artery and/or

the same arterial branch treated during the index PCI.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Reject file if ihcab=1 and ihtvcab is NULL

Obligation Always, if In-hospital Cardiothoracic Surgery = Yes

, a, e,	Toophan caraternorable sarbery res
Code	Description
0	No
1	Yes
0-1	
	spital cardiothoracic surgery was coded 'yes', did the surgery involve the placing of coronary artery bypass iscularise the target vessel?
Field cannot be blank if In-hospital Cardiothoracic Surgery = Yes. Field cannot be complete if In-hospital Cardiothoracic Surgery = No.	
1	
ihcab	
Number	
Child	
Reject file if ihcab=0 and ihtvcab has data	
	Code 0 1 0-1 Where In-horgrafts to reval Field cannot Surgery = No 1 ihcab Number Child

5.1.9. In-Hospital Stroke

Defining Attributes

Name In-Hospital Stroke

Field Name ihstr

Definition The patient experienced a stroke or new central neurologic deficit (persisting for > 72 hours) during the cardiac

catheter lab visit, after the lab visit, but prior to discharge and/or any subsequent lab visits.

Note: Stroke is evidenced by persistent loss of neurological function caused by an ischaemic or haemorrhagic event.

Justification Collected to allow reporting of NCR QI Peri-PCI stroke, and in-hospital MACCE.

Opligation	7 (1 Ways	
Representational Attributes		
Permitted Values	Code	Description
	0	No
	1	Yes
Range	0-1	
Guide for Use	Did the patient experience a stroke whilst an in-patient?	
Validation Rules	Field cannot be blank.	
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child	Parent	
Behaviour	Reject file if NULL	

5.1.10. In-Hospital Stroke Type

Defining Attributes

Name In-Hospital Stroke Type

Field Name ihstrt

Definition Indicate whether the type of stroke, either haemorrhagic or ischaemic. Select ONE of the following options:

Haemorrhagic: Imaging evidence (MRI or CT), lumbar puncture, neurosurgery or autopsy evidence confirms haemorrhage in the cerebral parenchyma, or subdural/subarachnoid haemorrhage; Ischaemic: In the absence of haemorrhagic evidence, inadequate cerebral blood supply is the suspected cause of stroke. Cerebrovascular occlusion may be caused by (but is not limited to) thrombus, embolism or vasoconstriction, etc.; Unknown: Stroke type could not be conclusively

determined.

Justification Collected to allow reporting of NCR QI Peri-PCI stroke.

Obligation Always, if In-hospital Stroke = Yes

Representational Attributes		
Permitted Values	Code	Description
	1	Haemorrhagic
	2	Ischaemic
	-1	Unknown
Range	-1 – 2	
Guide for Use	Where In-hospital stroke was coded 'yes', specify the type of stroke.	
Validation Rules	Field cannot be bl	ank if In-hospital Stroke = Yes. Field cannot be complete if In-hospital Stroke = No.
Maximum Field Size	1	
Related Data Field	ihstr	
Data Type	Number	
Parent / Child	Child	
Behaviour	Reject file if ihstr=	=O and ihstrt has data
	Reject file if ihstr=	=1 and ihstrt is NULL

5.1.11. In-Hospital Bleeding

Defining Attributes

Name In-Hospital Bleeding

Field Name ihbl

Definition The patient experienced a NEW bleeding event either during this cardiac catheter lab visit, after this lab visit but prior

to any subsequent lab visits and prior to discharge. Bleeding Academic Research Consortium (BARC): Type 0: No bleeding; Type 1: Bleeding that is not actionable and does not cause the patient to seek treatment; Type 2: Any clinically overt sign of haemorrhage that "is actionable" and requires diagnostic studies, hospitalization, or treatment by a health care professional; Type 3a: Overt bleeding plus haemoglobin drop of 3 to < 5 g/dL (provided haemoglobin drop is related to bleed); transfusion with overt bleeding Type 3b: Overt bleeding plus haemoglobin drop < 5 g/dL (provided haemoglobin drop is related to bleed); cardiac tamponade; bleeding requiring surgical intervention for control; bleeding requiring IV vasoactive agents; Type 3c: Intracranial haemorrhage confirmed by autopsy, imaging, or lumbar puncture; intraocular bleed compromising vision; Type 4: CABG-related bleeding within 48 hours; Type 5a: Probable fatal bleeding Type 5b: Definite fatal bleeding (overt or autopsy or imaging confirmation).

Justification Collected to allow reporting of NCR QI In-hospital major bleeding.

Representational Attributes			
Permitted Values	Code	Description	
	0	Type 0	
	1	Type 1	
	2	Type 2	
	3	Type 3a	
	4	Type 3b	
	5	Type 3c	
	6	Type 4	
	7	Type 5a	
	8	Type 5b	
Range	0-8		
Guide for Use	Did the bleeding occur during the admission?		
Validation Rules	Field cannot be blank. Field cannot be coded Type 4 if In-Hospital Cardiothoracic Surgery = No. Field cannot be coded 5a or 5b if Discharge Status is not coded as Hospital Mortality.		
Maximum Field Size Related Data Field	1		

Data Type Parent / Child Behaviour Number Parent

Reject file if NULL

Reject file if ihbl=6 and ihcab=0 Reject file if ihbl>6 and dis<6

5.1.12. In-Hospital Bleeding Site

Defining Attributes

Name In-Hospital Bleeding Site

Field Name ihblsite

Definition The bleeding site from the following: Retroperitoneal: Indicate whether retroperitoneal bleeding occurred during or after

the Cardiac Catheter Laboratory (CCL) visit and until discharge. Percutaneous entry site: Indicate whether bleeding occurred at the percutaneous entry site during or after the CCL visit and until discharge. Bleeding at the percutaneous entry site can be external or a haematoma >10cm for femoral access or >2cm for radial access or >5cm for brachial access. Gastrointestinal tract: Indicate whether gastrointestinal bleeding occurred during or after the CCL visit and until discharge. Cerebral: Indicate whether cerebral bleeding occurred during or after the CCL visit and until discharge. Genital/Urinary: Indicate whether genital or urinary bleeding occurred during or after the CCL visit and until discharge.

Other: Other bleeding site not indicated above.

Justification Collected to allow reporting of NCR QI In-hospital major bleeding.

Representational Attributes		
Permitted Values	Code	Description
	1	Retroperitoneal
	2	Percutaneous Entry Site
	3	Gastrointestinal
	4	Cerebral
	5	Genital/Urinary
	6	Other
Range	1-6	
Guide for Use	Where in-hos	spital bleeding was coded 'Type 1, 2, 3a, 3b, 3c, 4, 5a, or 5b', specify the site.
Validation Rules	Field cannot be blank if In-hospital Bleeding ≠ No. Field cannot be complete if In-hospital Bleeding = No.	
Maximum Field Size	1	
Related Data Field	ihbl	
Data Type	Number	
Parent / Child	Child	
Behaviour	Behaviour Reject file if ihbl = 0 and ihblsite has data	
	Reject file if il	nbl >=1 and ihblsite is NULL

5.1.13. In-Hospital Stent Thrombosis

Defining Attributes

Justification

Name In-Hospital Stent Thrombosis

Field Name ihst

Definition A stent thrombosis occurred during hospitalisation, after the index PCI lab visit, but prior to discharge and/or any

subsequent lab visits. Stent thrombosis is defined as the presence of a thrombus or angiographic documentation of vessel occlusion within a pre-existing stent OR within 5mm of the proximal or distal stent edges. Stent thromboses are classified by the Academic Research Consortium (ARC) as definite, probable, or silent. Select ONE of the following options that apply: No stent thrombosis: Where there are no acute coronary syndrome (ACS) symptoms or death after the index PCI lab visit and/or prior to discharge OR ACS symptoms occur during the defined period, but angiographic evidence does not indicate thrombus or occlusion. Definite stent thrombosis: The presence of an ACS with angiographic or autopsy evidence of thrombus or occlusion. Probable stent thrombosis: Includes unexplained deaths during the index admission after the procedure or acute myocardial infarction involving the target-vessel territory without angiographic or autopsy confirmation. Silent stent thrombosis: The incidental angiographic documentation of stent occlusion in the absence of

clinical signs or symptoms is not considered stent thrombosis.

Collected to allow reporting of in-hospital and 30-day MACE and MACCE.

Representational Attributes		
Permitted Values	Code	Description
	0	No Stent Thrombosis
	1	Definite Stent Thrombosis
	2	Probable Stent Thrombosis
	3	Silent Stent Thrombosis
Range	0-3	
Guide for Use	Did a stent thrombosis occur during the admission?	
Validation Rules	Field cannot be blank.	
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		
Behaviour	Reject file if NULL	

6. Discharge

6.1. Discharge Details

6.1.1. Discharge Status

Defining Attributes

Name Discharge Status

Field Name dis

Definition

The conditions under which the patient was discharged from the hospital following the index admission. Select one of the following options: Note: Patients who died during their index admission without being discharged from the acute care setting should be coded as hospital mortality. Home: Discharged to home, with no planned contact before routine

review; Hospital in the Home: Discharged to home, with planned visits to home by medical or paramedical staff; Rehabilitation Unit/Hospital: Discharged for inpatient rehabilitation; Local or referring hospital: Discharged to another hospital for continuing care not related to any complications arising from the index PCI; Tertiary referral centre: Discharged to another hospital or centre for tertiary care related to a complication arising from the index PCI (e.g. patient had PCI complications that required advanced medical treatment/investigation that could not be performed by the PCI

hospital); Hospital Mortality: Patient died in-hospital during or after the index PCI procedure, but prior to discharge.

Justification Collected to allow reporting of NCR QIs In-hospital and 30-day mortality.

Representational Attributes		
Permitted Values	Code	Description
	1	Home
	2	Hospital in the Home
	3	Rehab Unit/Hospital
	4	Local or Referring Hospital
	5	Tertiary Referral Centre
	6	Hospital Mortality
Range	1-6	
Guide for Use	Under what c	onditions was the patient discharged from the hospital?
Validation Rules	Field cannot be blank. If In-hospital Bleeding is 5a = Probable fatal bleeding or 5b = Definite fatal bleeding (overt or autopsy or imaging confirmation), then Discharge Status must = Hospital Mortality.	
Maximum Field Size Related Data Field	1	
Data Type	Number	

Parent / Child Behaviour

Reject file if ihbl>6 and dis=1-5

Reject file if dis = NULL

6.1.2. Date of Discharge/Hospital Mortality

Defining Attributes

Name Date of Discharge/Hospital Mortality

Field Name dod

Definition The date the patient was discharged from the PCI hospital admission (or the date of death where the patient died during

the index admission). Note: For a patient who dies in-hospital without being discharged from the acute care setting the

discharge and death date are the same.

Justification Collected to allow reporting of NCR QIs In-hospital and 30-day mortality.

Obligation Always

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use On what date was the patient discharged or date of in hospital mortality?

Validation Rules Field cannot be blank. Date of Discharge/Hospital Mortality must be ≥ Date of Procedure

Maximum Field Size

Related Data Field

Data Type Date/Time
Parent / Child Parent

Behaviour Reject file if dod < dop Reject file if dod = NULL

10

6.1.3. Length of Stay

Defining Attributes

Name Length of Stay

Field Name los

Definition To determine the length of hospital stay and patient discharged on the same day as their PCI

Justification Collected allow reporting on patients length of stay.

Obligation Derived Variable

Representational Attributes

Permitted Values NNN

Range

Guide for Use How long did the patient remain an inpatient in the PCI hospital?

Validation Rules Derived Variable

Maximum Field Size

Related Data Field

Data Type Number (Days)

Parent / Child

Behaviour los=[dod-doa]

6.1.4. Cardiac Rehabilitation Referral

Defining Attributes

Name Cardiac Rehabilitation Referral

Field Name crehab

Definition Written documentation of a patient referral to an outpatient cardiac rehabilitation program, or a documented reason

why referral was not made (by physician, nurse, or other hospital personnel).

Justification Collected to allow reporting of NCR QI Patients referred to cardiac rehabilitation or other secondary prevention program.

Obligation Always, if Discharge Status/Hospital Mortality ≠ Hospital Mortality

Representational Attributes $\begin{array}{ccc} \text{Permitted Values} & \text{Code} & \text{Description} \\ & 0 & \text{No} \\ & 1 & \text{Yes} \\ & -1 & \text{Unknown} \\ & \text{Range} & -1-1 \end{array}$

Guide for Use Was the patient referred to cardiac rehabilitation program?

Validation Rules Field cannot be blank if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be complete if Discharge

Status/Hospital Mortality = Hospital Mortality.

Maximum Field Size

Related Data Field

Data Type Number

Parent / Child

Behaviour Reject file if dis=6 and crehab has data

Reject file if dis < 6 crehab is NULL

6.1.5. Primary Cause of Death

Defining Attributes

Name Primary Cause of Death

Field Name mortc

Definition The cause of death. Primary cause of death is the first significant abnormal event which ultimately led to death. Select

ONE of the following options: **Non-cardiac**: The primary cause of death was clearly non-cardiac related AND there is documented evidence of a non-cardiac primary cause of death (e.g. cancer, renal failure, major trauma, etc.). **Cardiac**: The primary cause of death was diagnosed as clearly cardiac-related and includes, but is not exclusive to any of the following circumstances: chronic heart failure; acute coronary syndrome (e.g. STEMI, NSTEMI; recurrent or unstable

angina, etc.); arrhythmia; and/or bleeding or other complications following a cardiac procedure.

Justification Collected to allow reporting of NCR QI In-hospital mortality and 30-day mortality after PCI.

Representational Attributes			
Permitted Values	Code	Description	
	1	Cardiac	
	2	Non-Cardiac	
Range	1-2		
Guide for Use	Where Disch	narge status is coded "Hospital Mortality", what was the Primary Cause of Death?	
Validation Rules	Field cannot be complete if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be blank if Discharge		
	Status/Hosp	ital Mortality = Hospital Mortality.	
Maximum Field Size	1		
Related Data Field	dis		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if	dis < 6 and mortc has data	
	Reject file if	dis=6 and mortc is NULL	

6.2. Discharge Medications

6.2.1. Aspirin Prescribed at Discharge

Defining Attributes

Name Aspirin Prescribed at Discharge

Field Name dasp

Definition The patient is prescribed and taking aspirin at discharge from the PCI admission. Aspirin agents include: aspirin, astrix,

cardiprin, cartia, assasantin, aspro, disprin and solprin.

Justification Collected to allow reporting of NCR QI Proportion of patients without a clear and documented contraindication for Aspirin

and/or P2Y12 inhibitor discharged on DAPT.

	, ,	onar Be stated from the tarrey from the tarrey
Representational Attributes		
Permitted Values	Code	Description
	0	No
	1	Yes
	2	Contraindicated
	3	Not Collected
Range	0-3	
Guide for Use	Was aspirin prescribed at discharge?	
Validation Rules	Field cannot be blank if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality.	
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		
Behaviour	Reject file if o	dis=6 and dasp has data
	Reject file if o	dis = 1-5 and dasp is NULL

6.2.2. Other Antiplatelet Prescribed at Discharge

Defining Attributes

Name Other Antiplatelet Prescribed at Discharge

Field Name doap

Definition The patient is prescribed and taking other antiplatelets at discharge from the PCI admission. Antiplatelets agents include:

Thienopyridine agents (clopidogrel, ticlopidine, prasugrel), Ticagrelor.

Justification Collected to allow reporting of NCR QI Proportion of patients without a clear and documented contraindication for Aspirin

and/or P2Y12 inhibitor discharged on DAPT.

0.0.1.0.1.1	, - ,	
Representational Attributes		
Permitted Values	Code	Description
	0	No
	1	Yes
	2	Contraindicated
	3	Not Collected
Range	0-3	
Guide for Use	Were other antiplatelets (excluding aspirin) prescribed at discharge?	
Validation Rules	Field cannot be blank if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality.	
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		
Behaviour	Reject file if a	dis=6 and doap has data
	Reject file if a	dis = 1-5 and doap is NULL

6.2.3. Statin Prescribed at Discharge

Defining Attributes

Name Statin Prescribed at Discharge

Field Name dstp

Definition The patient is prescribed and taking a statin at discharge from the PCI admission. Statin agents include: atorvastatin,

fluvastatin, pravastatin, rosuvastatin and simvastatin.

Justification Collected to allow reporting of NCR QI Patients without contraindication discharged on lipid-lowering therapy.

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Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	2	Contraindicated	
	3	Not Collected	
Range	0-3		
Guide for Use	Was a statin prescribed at discharge?		
Validation Rules	Field cannot be blank if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality.		
Maximum Field Size	1		
Related Data Field			
Data Type	Number		
Parent / Child			
Behaviour	Reject file if	dis=6 and dstp has data	
	Reject file if	dis = 1-5 and dstp is NULL	

6.2.4. Other Lipid Lowering Therapies Prescribed at Discharge

Defining Attributes

Name Other Lipid Lowering Therapies Prescribed at Discharge

Field Name doll

Definition The patient is prescribed and taking other Lipid Lowering Therapies at discharge from the PCI admission.

Other lipid lowering agents include: ezetimibe and fibrates (gemfibrozil, fenofibrate and clofibrate).

Justification Collected to allow reporting of NCR QI Patients without contraindication discharged on lipid-lowering therapy.

	, ,	
Representational Attributes		
Permitted Values	Code	Description
	0	No
	1	Yes
	2	Contraindicated
	3	Not Collected
Range	0-3	
Guide for Use	Were other Lipid Lowering Therapies (excluding statins) prescribed at discharge?	
Validation Rules	Field cannot be blank if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality.	
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		
Behaviour	Reject file if d	lis=6 and doll has data
	Reject file if o	lis = 1-5 and doll is NULL

7. 30 Day Follow Up

7.1. 30 Day Outcomes

7.1.1. Date of Follow-Up

Defining Attributes

Name Date of Follow-Up

Field Name dfu30

Definition Date the follow-up was undertaken. All patients must be followed up (or attempted to be followed up) at 30 days post

discharge from the PCI admission.

Justification Collected to allow reporting of 30-day outcomes.

Obligation Always

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use On what date was the follow-up undertaken? All patients must be followed up (or attempted to be followed up) at 30

days post discharge from the PCI admission.

Validation Rules Field cannot be blank if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be complete if Discharge

Status/Hospital Mortality = Hospital Mortality. Date of Follow-Up cannot < Date of Discharge.

Maximum Field Size

Related Data Field

Data Type Date/Time

Parent / Child

Behaviour Reject file if dfu30 is NULL AND dis=1-5

10

Reject file if dis=6 and dfu30 contains data

Reject file if dfu30 <dod

7.1.2. Follow-Up Status

Defining Attributes

Name Follow-Up Status

Field Name stat30

Definition The patient's mortality status. I.e., whether the patient was alive or dead at 30 days after discharge from the index

admission.

Justification Collected to allow reporting of 30-day outcomes and NCR QI 30-day mortality.

Danna aantatianal Attributaa	•		
Representational Attributes			
Permitted Values	Code	Description	
	0	Alive	
	1	Deceased	
	-1	Lost to Follow-Up	
Range	-1 - 1		
Guide for Use	Was the patient alive or dead at follow-up?		
Validation Rules	Field cannot be blank if Discharge Status/Hospital Mortality ≠ Hospital Mortality. Field cannot be complete if Discharge		
	Status/Hospi	ital Mortality = Hospital Mortality.	
Maximum Field Size	1		
Related Data Field			
Data Type	Number		
Parent / Child	Parent		
Behaviour	Reject file if	stat30 is NULL AND dis=1-5	
	Reject file if	dis=6 and stat30 contains data	

7.1.3. Date of Death

Defining Attributes

Name Date of Death Field Name dmort30

Definition The date on which the patient died.

Justification Collected to allow reporting of NCR QI 30-day mortality after PCI.

Obligation Always, if Follow-Up Status = Deceased

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use Where the Follow-up status is coded as 'Deceased', what was the date of death?

Validation Rules Field cannot be blank if Follow-Up Status = Deceased. Field cannot be complete if Follow-Up Status = Alive. Field cannot

< Date of Discharge. Field cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality.

Maximum Field Size 10
Related Data Field stat30
Data Type Date/Time
Parent / Child Child

Behaviour Reject file if dmort30 < dod

Reject file if stat30=1 and dmort30 is NULL Reject file if stat30=0 and dmort30 contains data Reject file if dis=6 and dmort30 contains data

7.1.4. Primary Cause of Death

Defining Attributes

Behaviour

Name Primary Cause of Death

Field Name mort30r

Definition Primary cause of death is the first significant abnormal event which ultimately led to death. Select ONE of the following

options: Non-cardiac: The primary cause of death was clearly non-cardiac related AND there is documented evidence of a non-cardiac primary cause of death (e.g. cancer, renal failure, major trauma, etc.). Cardiac: The primary cause of death was diagnosed as clearly cardiac-related and includes, but is not exclusive to any of the following circumstances: chronic heart failure; acute coronary syndrome (e.g. STEMI, NSTEMI; recurrent or unstable angina, etc.); arrhythmia; and/or bleeding or other complications following a cardiac procedure. Uncertain: The primary cause of death is not definitively

known.

Justification Collected to allow reporting of NCR QI 30-day mortality.

Obligation Always, if Follow-Up Status = Deceased

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Representational Attributes		
Permitted Values	Code	Description
	1	Cardiac
	2	Non-Cardiac
	3	Uncertain
Range	1-3	
Guide for Use	Where the Fo	ollow-up status is coded as 'Deceased', what was the cause of death?
Validation Rules	Field cannot	be blank if Follow-Up Status = Deceased. Field cannot be complete if Follow-Up Status = Alive.
Maximum Field Size	1	
Related Data Field	stat30	
Data Type	Number	
Parent / Child	Child	

Reject file if stat30=0 or -1 AND mort30r contains data

Reject file if stat30=1 and mort30r is NULL

7.1.5. New MI

Defining Attributes

Name New MI
Field Name mi30
Definition The pati

The patient experienced a new myocardial infarction (MI) within 30 days from the date of discharge from the index admission. The new MI is defined by the following criteria:

Non ST-Elevation Myocardial Infarction (NSTEMI): AT LEAST ONE of the following biomarkers for detecting myocardial necrosis MUST be present (refer to note regarding Reference Control Limits): 1. Troponin T or I: Maximal concentration of Troponin T or I > the MI diagnostic limit on at least one occasion within 24 hours from the index clinical event; 2. CK-MB: Maximal value of CK-MB >2x the upper limit of normal (ULN) on one occasion during the first hours after the index clinical event; OR Maximal value of CK-MB (preferable CK-MB mass) > ULN on two successive samples. 3. Total CK: Only where Troponin or CK-MB assays are unavailable, total CK >2x the ULN (or the B fraction of CK) may be employed. Note: The preferred assay to use as biomarkers for myocardial necrosis is Troponin. In the absence of troponin, CK-MB is the next best alternative but total CK can be used in the absence of troponin and CK-MB. AND ONE of the following: 1. Either ST segment depression or T wave abnormalities in the ECG; or 2. Ischaemic symptoms in the presence or absence of chest discomfort. Ischaemic symptoms may include: Unexplained nausea and vomiting; or Persistent shortness of breath secondary to left ventricular failure; or Unexplained weakness, dizziness, light-headedness or syncope.

ST-Elevation Myocardial Infarction (STEMI): AT LEAST ONE of the following biomarkers for detecting myocardial necrosis MUST be present (refer to note regarding Reference Control Limits): 1. Troponin T or I: Maximal concentration of Troponin T or I > the MI diagnostic limit on at least one occasion within 24 hours from the index clinical event; 2. CK-MB: Maximal value of CK-MB >2x the upper limit of normal (ULN) on one occasion during the first hours after the index clinical event; OR Maximal value of CK-MB (preferable CK-MB mass) > ULN on two successive samples.

- 3. Total CK: Only where Troponin or CK-MB assays are unavailable, total CK >2x the ULN (or the B fraction of CK) may be employed. Note: The preferred assay to use as biomarkers for myocardial necrosis is Troponin. In the absence of troponin, CK-MB is the next best alternative but total CK can be used in the absence of troponin and CK-MB.

 AND ONE of the following ECG changes. 1. ST-segment elevation: New or presumed new ST segment elevation at the J point in two or more contiguous leads with the cut-off points ≥ 0.2mV in leads V1, V2, or V3, or ≥0.1 mV in other leads.
- 2. Development of any Q wave in leads V1 through V3, or the development of a Q-wave ≥ 30ms (0.03s) in leads I, II, aVL, aVF, V4, V5, or V6. (Q wave changes must be present in any two contiguous leads, and be ≥1mm in depth). Collected to allow reporting of 30-day MACE and MACCE.

Always, if Follow-Up Status ≠ Lost to Follow-Up.

Justification Obligation

Representational Attributes

Permitted Values	Code	Description
	0	No
	1	Yes
	-1	Unknown
Range	-1 - 1	
Guide for Use	Did the patie	ent experience a myocardial infarction during the 30-day follow-up period?
Validation Rules	Field cannot be blank if Follow-Up Status ≠ Deceased. Field cannot be complete if Follow-Up Status = Lost to Follow-Up.	
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		
Behaviour	Reject file if I	NULL and stat30≥0
	Reject file if s	stat30=-1 and mi30 contains data
	Reject file if	dis=6 and mi30 contains data

7.1.6. New Stent Thrombosis

Defining Attributes

Name New Stent-Thrombosis

Field Name st30

Definition A stent thrombosis occurred within 30 days from the date of discharge from the current admission. Stent thrombosis is

defined as the presence of a thrombus or angiographic documentation of vessel occlusion within a pre-existing stent OR within 5mm of the proximal or distal stent edges. Stent thromboses are classified by the Academic Research Consortium (ARC) as definite, probable, or silent. Select ONE of the following options that apply: No stent thrombosis: Where there are no acute coronary syndrome (ACS) symptoms or death after the index PCI lab visit and/or prior to discharge OR ACS symptoms occur during the defined period, but angiographic evidence does not indicate thrombus or occlusion. Definite stent thrombosis: The presence of an ACS with angiographic or autopsy evidence of thrombus or occlusion. Probable stent thrombosis: Includes unexplained deaths during the index admission after the procedure or acute myocardial infarction involving the target-vessel territory without angiographic or autopsy confirmation. Silent stent thrombosis: The incidental angiographic documentation of stent occlusion in the absence of clinical

signs or symptoms is not considered stent thrombosis. Unknown.

Justification Collected to allow reporting of 30-day MACE and MACCE.

Obligation Always, if Follow-Up Status ≠ Lost to Follow-Up.

Representational Attributes		
Permitted Values	Code	Description
	0	No Stent Thrombosis
	1	Definite Stent Thrombosis
	2	Probable Stent Thrombosis
	3	Silent Stent Thrombosis
	-1	Unknown
Range	-1 - 3	
Guide for Use	Did a stent th	rombosis occur during the 30-day follow-up period?
Validation Rules	Field cannot b	be blank if Follow-Up Status ≠ Lost to Follow-Up. Field cannot be complete if Follow-Up Status = Lost to
	Follow-Up. Fie	eld cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality
Maximum Field Size	1	
Related Data Field		
Data Type	Number	
Parent / Child		

Behaviour	Reject file if NULL and stat30≥0	
	Reject file if stat30=-1 and st30 contains data	
	Reject file if dis=6 and st30 contains data	

7.1.7. New Stroke

Defining Attributes

Name New Stroke

Field Name nstr

Definition The patient experienced a stroke or new central neurologic deficit (persisting for > 72 hours) in the 30-days post

discharge. Stroke is evidenced by persistent loss of neurological function caused by an ischaemic or haemorrhagic event.

Justification Collected to allow reporting on 30-day stroke and 30-day MACCE.

Obligation Always, if Follow-Up Status ≠ Lost to Follow-Up.

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Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	-1	Unknown	
Range	-1 - 1		
Guide for Use	Did the patie	ent experience a stroke during the 30-day follow-up period?	
Validation Rules	Field cannot be blank if Follow-Up Status ≠ Lost to Follow-Up. Field cannot be complete if Follow-Up Status = Lost to		
	Follow-Up. F	Field cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality	
Maximum Field Size	1		
Related Data Field			
Data Type	Number		
Parent / Child	Parent		
Behaviour	Reject file if	NULL and stat30≥0	
	Reject file if	stat30=-1 and nstr contains data	
	Reject file if	dis=6 and nstr contains data	

7.1.8. New Stroke Type

Defining Attributes

Name New Stroke Type

Field Name nstrt

Definition Type of new stroke. Either haemorrhagic or ischaemic. Select ONE of the following options: Haemorrhagic: Imaging

evidence (MRI or CT), lumbar puncture, neurosurgery or autopsy evidence confirms haemorrhage in the cerebral parenchyma, or subdural/subarachnoid haemorrhage; Ischaemic: In the absence of haemorrhagic evidence, inadequate cerebral blood supply is the suspected cause of stroke. Cerebrovascular occlusion may be caused by (but is not limited

to) thrombus, embolism or vasoconstriction, etc. Unknown: Stroke type could not be conclusively determined.

Justification Collected to allow reporting on 30-day stroke and 30-day MACCE.

Obligation Always, if Follow-Up Status ≠ Lost to Follow-Up.

Representational Attributes			
Permitted Values	Code	Description	
	1	Haemorrhagic	
	2	Ischaemic	
	-1	Unknown	
Range	-1 - 2		
Guide for Use	Where new stroke was coded 'yes', specify the type of stroke.		
Validation Rules	Field cannot be blank if New Stroke = Yes. Field cannot be complete if New Stroke = No.		
Maximum Field Size	1		
Related Data Field	nstr		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if n	nstr=1 and nstrt is NULL	
		nstr = 0 or -1 and nstrt contains data	

7.2. 30-Day Rehospitalisations

NOTE: Up to 6 rehospitalisations can be entered. Fields are coded: NAME; NAME 2; NAME 3-6.

7.2.1. Cardiac Rehospitalisation

Defining Attributes

Name Cardiac Rehospitalisation

Field Name crh30

Definition The patient had a cardiac rehospitalisation, where a cardiac rehospitalisation is defined as an in-patient admission that

includes, but is not limited to any of the following circumstances: heart failure; acute coronary syndrome (e.g STEMI, NSTEMI; recurrent or unstable angina); arrhythmia; elective (non-emergency) PCI or CABG etc. Include rehospitalisations with a primary diagnosis that has the following ICD-10 AM codes: I00-I02: Acute rheumatic fever; I05-I09: Chronic rheumatic heart disease; I10-I15: Hypertensive diseases; I20-I25: Ischaemic heart disease; I30-I52: Other forms of heart disease; I70-I72: Atherosclerosis, aortic aneurysm and dissection; I74: Arterial embolism and thrombosis; I77.0: Arteriovenous fistula, acquired; R00: Abnormalities of heart beat; R01: Cardiac murmurs and other cardiac sounds; R03: Abnormal blood pressure reading; R07: Pain in throat and chest; All

other ICD-10 codes should be classified as non-cardiac.

Justification Collected to allow reporting of NCR QI 30-day unplanned cardiac readmission rate after PCI.

Obligation Always

Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	-1	Unknown	
Range	-1 - 1		
Guide for Use	Did the patient have a cardiac rehospitalisation within 30 days?		
Validation Rules	Field cannot be blank if Follow-Up Status ≠ Lost to Follow-Up. Field cannot be complete if Follow-Up Status = Lost to Follow-Up. Field cannot be complete if Discharge Status/Hospital Mortality = Hospital Mortality.		
Maximum Field Size	1		
Related Data Field			
Data Type Parent / Child	Number		

Behaviour	Reject file if crh30 is NULL and stat30≥0
	Reject file if crh30 contains data and stat30=-1
	Reject file if dis=6 and crh30 contains data

7.2.2. Cardiac Rehospitalisation Date

Defining Attributes

Name Cardiac Rehospitalisation Date

Field Name rhdte

Definition The date of cardiac rehospitalisation

Justification Collected to allow reporting of NCR QI 30-day unplanned cardiac readmission rate after PCI.

Obligation Always, if Cardiac Rehospitalisation = Yes.

Representational Attributes

Permitted Values NN/NN/NNNN Range DD/MM/YYYY

Guide for Use Where cardiac rehospitalisation is coded as 'Yes', on what date was the rehospitalisation?

Validation Rules Field cannot be blank if Cardiac Rehospitalisation = Yes. Field cannot be complete if Cardiac Rehospitalisation = No.

Cardiac Rehospitalisation Date cannot > Date of Discharge + 30. Cardiac Rehospitalisation Date cannot < Date of

Discharge.

Maximum Field Size 10
Related Data Field crh30
Data Type Date/Time
Parent / Child Child

Behaviour Reject file if rhdte >dod + 30

Reject file if rhdte <dod

Reject file if crh30 = 1 and rhdte is NULL Reject file if crh30=0 and rhdte has data

7.2.3. Planned Cardiac Rehospitalisation

Defining Attributes

Name Planned Cardiac Rehospitalisation

Field Name pc30

Definition The cardiac rehospitalisation was planned.

Note: A 'planned cardiac rehospitalisation' is defined as an in-patient

admission to hospital for a <u>scheduled</u> visit only. For emergency admissions, code as 'no'.

Justification Collected to allow reporting of NCR QI 30-day unplanned cardiac readmission rate after PCI.

Obligation Always, if Cardiac Rehospitalisation = Yes.

	Always, il Cardiac (Criospitalisation – 163.		
Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	-1	Unknown	
Range	-1 - 1		
Guide for Use	Where Cardiac rehospitalisation is coded as 'Yes', was the hospitalisation planned?		
Validation Rules	Field cannot be blank if Cardiac Rehospitalisation = Yes. Field cannot be complete if Cardiac Rehospitalisation = No.		
Maximum Field Size	1		
Related Data Field	crh30		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if c	crh30=1 AND pc30 is NULL	
	Reject file if o	crh30=0 and pc30 has data	

7.2.4. PCI Rehospitalisation

Defining Attributes

Name PCI Rehospitalisation

Field Name pci30

Definition Indicate whether Percutaneous Coronary Intervention (PCI) was performed during the cardiac rehospitalisation

admission

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation Always, if Cardiac Rehospitalisation = Yes.

	, ,		
Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	-1	Unknown	
Range	-1 - 1		
Guide for Use	Where Cardiac Rehospitalisation is coded as 'Yes', was PCI performed during this admission?		
Validation Rules	Field cannot be blank if Cardiac Rehospitalisation = Yes. Field cannot be complete if Cardiac Rehospitalisation = No.		
Maximum Field Size	1		
Related Data Field			
Data Type	Number		
Parent / Child	Parent		
Behaviour	Reject file if c	rh30=1 AND pci30 is NULL	
	Reject file if c	rh30=0 and pci30 has data	

7.2.5. Target Vessel Revascularisation (PCI)

Defining Attributes

Name Target Vessel Revascularisation (PCI)

Field Name tvr30

Definition The target vessel was revascularised during the post-discharge subsequent PCI. Note: A PCI TVR is any repeated

percutaneous revascularisation due to restenosis/occlusion within the target coronary artery and/or the same arterial branch that was treated during the index PCI. This includes any percutaneous revascularisation within the same arterial

branch treated during the index PCI, regardless of whether the index PCI was successful.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation Always, if PCI Rehospitalisation = Yes.

Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	-1	Unknown	
Range	-1 - 1		
Guide for Use	Guide for Use Where PCI rehospitalisation is coded as 'Yes', was the target vessel revascularised during the		
	PCI?		
Validation Rules	Field cannot be blank if PCI Rehospitalisation = Yes. Field cannot be complete if PCI Rehospitalisation = No.		
Maximum Field Size	1		
Related Data Field	pci30		
Data Type	Number		
Parent / Child	Child		
Behaviour Reject file if pci30 =0 or -1 and tvr30 has data		pci30 =0 or -1 and tvr30 has data	
	Reject file if	pci30=1 and tvr30 is NULL	

7.2.6. Target Lesion Revascularisation (TLR)

Defining Attributes

Name Target Lesion Revascularisation (TLR) Vessel Revascularisation (PCI)

Field Name tlr30

Definition The target lesion was revascularised during the post-discharge subsequent PCI. Note: A target lesion revascularisation

(TLR) is any repeated percutaneous revascularisation of the same lesion treated during the index PCI. This includes treatment of a restenosis/occlusion within a stent implanted during the index PCI (or within 5-mm of the proximal and distal margins of the stent edges). It also includes attempts to revascularise the target lesion where a stent was not

successfully implanted and/or plain old balloon angioplasty (POBA) was employed during the index PCI.

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation Always, if Target Vessel Revascularisation = Yes.

Representational Attributes		
Permitted Values	Code	Description
	0	No
	1	Yes
	-1	Unknown
Range	-1 - 1	
Guide for Use	Where PCI rehospitalisation is coded as 'Yes', was the target lesion revascularised during the post-discharge subsequent PCI?	
Validation Rules	Field cannot be blank if Target Vessel Revascularisation = Yes. Field cannot be complete if Target Vessel Revascularisation = No.	
Maximum Field Size	1	
Related Data Field	tvr30	
Data Type	Number	
Parent / Child	Child	
Behaviour	Reject file if tvr30 =0 or -1 and tlr30 has data	
	Reject file if t	vr30=1 and tlr30 is NULL

7.2.7. CABG Rehospitalisation

Defining Attributes

Name CABG Rehospitalisation

Field Name cab30

Definition Coronary Bypass Grafting (CABG) was performed during the cardiac rehospitalisation admission

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Obligation Always, if Cardiac Rehospitalisation = Yes.

Obligation	Always, it caldide heriospitalisation – res.		
Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	-1	Unknown	
Range	-1 - 1		
Guide for Use	Where Cardiac Rehospitalisation is coded as 'Yes', was CABG performed during this admission?		
Validation Rules	Field cannot be blank if Cardiac Rehospitalisation = Yes. Field cannot be complete if Cardiac Rehospitalisation = No.		
Maximum Field Size	1		
Related Data Field	crh30		
Data Type	Number		
Parent / Child	Child		
Behaviour Reject file if crh30=1 AND cab30 is NULL		crh30=1 AND cab30 is NULL	
	Reject file if	crh30=0, or -1, or NULL AND cab30 has data	

7.2.8. Target Vessel CABG

Defining Attributes

Name Target Vessel CABG

Field Name tvcab30

Definition The surgery involved the placing of coronary artery bypass grafts to revascularise the target vessel treated during the

index PCI

Justification Collected to allow reporting of NCR QI Unplanned revascularisation within 30 days.

Reject file if cab30=1 and tvcab30 is NULL

Obligation Always, if CABG Rehospitalisation = Yes.

Representational Attributes			
Permitted Values	Code	Description	
	0	No	
	1	Yes	
	-1	Unknown	
Range	-1 - 1		
Guide for Use	Where CABG rehospitalisation is coded as 'Yes', did the surgery involved the placing of coronary artery bypass grafts to revascularise the target vessel treated during the index PCI?		
Validation Rules	Field cannot be blank if CABG Rehospitalisation = Yes. Field cannot be complete if CABG Rehospitalisation = No.		
Maximum Field Size	1		
Related Data Field	cab30		
Data Type	Number		
Parent / Child	Child		
Behaviour	Reject file if cab30 =0, or -1, or NULL and tvcab30 has data		

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