TO: NQF Members and Public

FR: NQF Staff

RE: Pre-voting review for National Voluntary Consensus Standards: Cardiovascular

Endorsement Maintenance, 2010: A Consensus Report

DA: July 5, 2011

During the past nine years, the National Quality Forum (NQF) has endorsed a large number of consensus standards to evaluate the quality of care for cardiovascular conditions in the ambulatory and hospital settings. As the quality measurement enterprise has matured and better data systems have become available, the demand for meaningful performance measures has prompted development of more sophisticated measures of healthcare processes and outcomes for cardiovascular disease. An evaluation of all NQF-endorsed cardiovascular measures and consideration of new measures will ensure the currency of NQF's portfolio of voluntary consensus standards.

A <u>20-member Steering Committee</u> representing a range of stakeholder perspectives was appointed to evaluate 26 new measures and 31 previously endorsed measures for maintenance review.

The draft document, <u>National Voluntary Consensus Standards: Cardiovascular Endorsement</u> <u>Maintenance, 2010: A Consensus Report</u>, is posted on the NQF website along with the following additional information:

- Measure submission forms
- Meeting and call summaries from the Steering Committee's discussions.

The draft report presents a summary of the large amount of information considered by the Steering Committee. The draft report includes many links to various NQF documents (measure submission forms, meeting and call summaries, patient-focused episode of care report, eMeasures list, etc.) to assist in navigation among various documents that provide supporting information on the evaluation of the measures.

Pursuant to section II.A of the Consensus Development Process v. 1.8, this draft document, along with the accompanying material, is being provided to you at this time for purposes of review and comment only and is not intended to be used for voting purposes. You may post your comments and view the comments of others on the NQF website.

Because of the close relationship of the topics addressed in Phases I and II of the Cardiovascular Endorsement Maintenance project, NQF prepared a combined draft report rather than a separate report for each phase. To provide sufficient time to review the report **the comment period for this report is 45 days**.

Please note that the organization of this report has been modified, similar to the recent End Stage Renal Disease and Surgery Endorsement Maintenance reports. The intention is to begin with high-level information (e.g., overarching evaluation issues and lists of measures) followed by more detail about the evaluation ratings and rationale in the measure evaluation summary tables.

Hyperlinks are provided to navigate to the detailed measure specifications for the recommended measures in Appendix A and to access all submitted measure information posted on the project web page.

NQF Member comments must be submitted no later than 6:00 pm ET, August 19, 2011.

Public comments must be submitted no later than 6:00 pm ET, August 12, 2011.

Thank you for your interest in NQF's work. We look forward to your review and comments.

NATIONAL VOLUNTARY CONSENSUS STANDARDS: CARDIOVASCULAR ENDORSEMENT MAINTENANCE 2010: A CONSENSUS REPORT

DRAFT REPORT FOR COMMENTING

July 5, 2011

NATIONAL VOLUNTARY CONSENSUS STANDARDS: CARDIOVASCULAR ENDORSEMENT MAINTENANCE 2010:

A CONSENSUS REPORT TABLE OF CONTENTS

EXECUTIVE SUMMARY	3
BACKGROUND	6
STRATEGIC DIRECTIONS FOR NQF	6
NATIONAL PRIORITIES PARTNERSHIP AND THE NATIONAL QUALITY STRATEGY	7
PRIOR NQF WORK RELATED TO CARDIOVASCULAR CONDITIONS	8
Endorsement of Consensus Standards	8
Patient-Focused Episode of Care Measurement Framework	8
NQF'S CONSENSUS DEVELOPMENT PROCESS	9
OVERARCHING MEASURE EVALUATION ISSUES	10
RECOMMENDATIONS FOR ENDORSEMENT	14
CORONARY ARTERY DISEASE -SECONDARY PREVENTION	18
CAD ACUTE PHASE: AMI AND PCI	35
CARDIAC REHABILITATION	61
ATRIAL FIBRILLATION	66
IMPLANTABLE CARDIOVERTER DEFIBRILLATORS (ICD)	71
HEART FAILURE	76
HYPERTENSION	92
RETIRED MEASURES	95
NOTES	96

APPENDIX A: SPECIFICATIONS FOR THE NATIONAL VOLUNTARY CONSENSUS STANDARDS: CARDIOVASCULAR ENDORSEMENT MAINTENANCE, 2010: A CONSENSUS REPORT

APPENDIX B: NATIONAL VOLUNTARY CONSENSUS STANDARDS: CARDIOVASCULAR ENDORSEMENT MAINTENANCE 2010 STEERING COM MITTEE AND NQF STAFF

APPENDIX C: ENDORSED CARDIOVASCULAR CONSENSUS STANDARDS (AFTER 2008)

APPENDIX D: GAPS IN THE CARDIOVASCULAR PORTFOLIO

1	NATIONAL VOLUNTARY CONSENSUS STANDARDS:
2	CARDIOVASCULAR ENDORSEMENT MAINTENANCE 2010:
3	A CONSENSUS REPORT
4	
5	EXECUTIVE SUMMARY
6	Measuring the quality of care for cardiovascular conditions is critically important. The human
7	and financial costs of cardiovascular disease are enormous. Heart disease is the leading cause of
8	death for men and women in the United States and cost the United States \$316.4 billion in 2010.
9	Hypertension affects 1 in 3 Americans, which increases their risk for heart disease, stroke, or
10	kidney disease and will cost \$76.6 billion in healthcare services, medications, and missed days of
11	work. ¹
12	During the past nine years, the National Quality Forum (NQF) has endorsed a large number of
13	consensus standards to evaluate the quality of care for cardiovascular conditions in the
14	ambulatory and hospital settings. As the quality measurement enterprise has matured, better data
15	systems have become available, electronic health records are closer to reality, and the demand
16	for meaningful performance measures has prompted development of more sophisticated
17	measures of healthcare processes and outcomes for cardiovascular disease. An evaluation of all
18	NQF-endorsed® cardiovascular measures and consideration of new measures will ensure the
19	currency of NQF's portfolio of voluntary consensus standards.
20	This report presents the results of the evaluation of 57 measures considered under NQF's
21	Consensus Development Process (CDP). Thirty-eight measures are recommended for
22	endorsement as voluntary consensus standards suitable for public reporting and quality
23	improvement. Of these, 31 are NQF-endorsed measures that have been reviewed for continued
24	endorsement as part of the maintenance process.
25	
26	CORONARY ARTERY DISEASE - SECONDARY PREVENTION
27	• 0076 optimal vascular care (Minnesota Community Measurement)
28	• 0073 IVD: blood pressure management (NCQA)
29	• 0068 IVD: use of aspirin or another antithrombotic (NCQA)

• 0075 IVD- complete lipid profile and LDL control <100 (NCQA)

0074 Chronic stable coronary artery disease: lipid control (PCPI)

• 0067 CAD: antiplatlet therapy (PCPI)

33 34	• 0066 Chronic stable coronary artery disease: ACE inhibitor or ARB therapy—diabetes or left ventricular systolic dysfunction (LVEF < 40%) (PCPI)
35 36	• 0071 AMI: Persistence of beta blocker therapy after a heart attack (NCQA)
37	CORONARY ARTERY DISEASEACUTE PHASE: ACUTE MYOCARDIAL
38	INFARCTION AND PERCUTANEOUS CORONARY INTERVENTION
39	
40	• 0289 Median time to ECG (CMS)
41	• 0286 Aspirin at arrival [for patients being transferred] (CMS)
42	• 0288 Fibrinolytic therapy received within 30 minutes of ED arrival and
43	Median time to fibrinolysis [for patients being transferred] (CMS)
44	• 0290 Median time to transfer to another facility for acute coronary intervention (CMS)
45	• 0132 Aspirin at arrival for acute myocardial infarction (AMI) (CMS)
46	• 0163 Primary PCI within 90 minutes of hospital arrival (CMS)
47	• 0164 Fibrinolytic therapy received within 30 minutes of hospital arrival (CMS)
48	• 0137 ACEI or ARB for left ventricular systolic dysfunction- acute myocardial infarction
49	(AMI) patients (CMS)
50	 0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute
51	myocardial infarction (AMI) hospitalization (CMS)
52	 0355 Bilateral cardiac catheterization rate (IQI 25) (AHRQ)
53	 0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge (ACCF)
54	• 0133 PCI mortality (risk-adjusted) (ACC)
55	 0160 Beta blocker prescribed at discharge* (CMS)
56	• 0142 Aspirin prescribed at discharge for AMI* (CMS)
57	

ATRIAL FIBRILLATION

- 1524 Assessment of thromboembolic risk (CHADS 2) (ACCF/AHA/PCPI)
- 1525 Chronic anticoagulation therapy (ACCF/AHA/PCPI)

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IMPLANTABLE CARDIOVERTER DEFIBRILLATOR (ICD)

- 1522 ACE/ARB therapy at discharge for ICD implant patients with LVSD (ACCF)
- 1528 Beta blocker at discharge for ICD implant patients with a previous MI (ACCF)
- 1529 Beta blocker at discharge for ICD implant patients with LVSD (ACCF)
- 0965 Patients with an ICD implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge (ACCF)

67 68 69

HEART FAILURE

70	• 0079 Heart failure: Left ventricular ejection fraction assessment (outpatient setting)
71	(PCPI)
72	• 0081 Heart failure: ACEI or ARB therapy for left ventricular systolic dysfunction (PCPI)
73	• 0083 Heart Failure: Beta-blocker therapy for left ventricular systolic dysfunction (PCPI)
74	 0135 Evaluation of left ventricular systolic dysfunction (CMS)*
75 76	 0162 ACEI or ARB for left ventricular systolic dysfunction – heart failure patients (CMS)
77	 0358 Congestive heart failure (CHF) mortality rate (IQI 16)(AHRQ)
78	• 0277 CHF admission (PQI 8) (AHRQ)
79 80	 0229 Hospital 30-day, all-cause, risk standardized mortality rate (RSMR) following heart failure hospitalization (CMS)
81	• 0330 Hospital 30-day, all-cause, risk standardized readmission rate following heart
82	failure hospitalization (risk adjusted) (CMS)
83	
84	HYPERTENSION
85	• 0018 Controlling high blood pressure (NCQA)
86	
87	
88	
89	
90	*Endorsement with placement in reserve status
91	

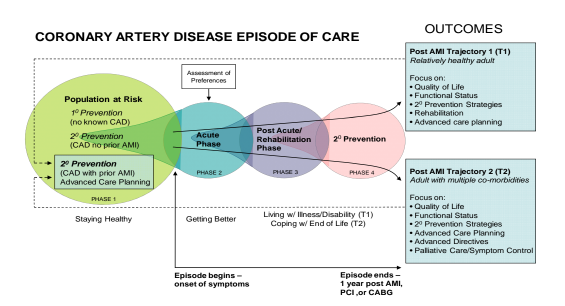
92	NATIONAL VOLUNTARY CONSENSUS STANDARDS:
93	CARDIOVASCULAR ENDORSEMENT MAINTENANCE 2010:
94	A CONSENSUS REPORT
95	
96	BACKGROUND
97	Measuring the quality of care for cardiovascular conditions is critically important. The human
98	and financial costs of cardiovascular disease are enormous. Heart disease is the leading cause of
99	death for men and women in the United States and cost the United States \$316.4 billion in 2010.
100	Hypertension affects 1 in 3 Americans, which increases their risk for heart disease, stroke, or
101	kidney disease and will cost \$76.6 billion in healthcare services, medications, and missed days of
102	work. ¹
103	
104	During the past nine years, the National Quality Forum (NQF) has endorsed a large number of
105	consensus standards to evaluate the quality of care for cardiovascular conditions in the
106	ambulatory and hospital settings. As the quality measurement enterprise has matured, better data
107	systems have become available, electronic health records are closer to reality, and the demand
108	for meaningful performance measures has prompted development of more sophisticated
109	measures of healthcare processes and outcomes for cardiovascular disease. Evaluation of NQF-
110	endorsed® cardiovascular measures and consideration of new measures will ensure the currency
111	of NQF's portfolio of consensus standards.
112	
113	STRATEGIC DIRECTIONS FOR NQF
114	NQF's mission includes three parts: 1) setting national priorities and goals for performance
115	improvement, 2) endorsing national consensus standards for measuring and publicly reporting on
116	performance, and 3) promoting the attainment of national goals through education and outreach
117	programs. As greater numbers of quality measures are developed and brought to NQF for
118	consideration of endorsement, NQF must assist stakeholders in measuring "what makes a
119	difference" and addressing what is important to achieve the best outcomes for patients and
120	populations.

121	
122	Several strategic issues have been identified to guide consideration of candidate consensus
123	standards:
124	DRIVE TOWARD HIGH PERFORMANCE. Over time, the bar of performance expectations
125	should be raised to encourage achievement of higher levels of system performance.
126	EMPHASIZE COMPOSITES. Composite measures provide much-needed summary
127	information pertaining to multiple dimensions of performance and are more comprehensible to
128	patients and consumers.
129	MOVE TOWARD OUTCOME MEASUREMENT. Outcome measures provide information
130	of keen interest to consumers and purchasers, and when coupled with healthcare process
131	measures, they provide useful and actionable information to providers. Outcome measures also
132	focus attention on much-needed system-level improvements because achieving the best patient
133	outcomes often requires a carefully designed care process, teamwork, and coordinated action on
134	the part of many providers.
135	CONSIDER DISPARITIES IN ALL WE DO. Some of the greatest performance gaps relate to
136	care of minority populations. Particular attention should be focused on identifying disparities-
137	sensitive performance measures and on identifying the most relevant
138	gender/race/ethnicity/language/socioeconomic strata for reporting purposes.
139	
140	NATIONAL PRIORITIES PARTNERSHIP AND THE NATIONAL QUALITY
141	STRATEGY
142	The National Priorities Partnership, a multi-stakeholder collaborative of 48 organizations
143	convened by NQF, plays a key role in identifying strategies for achieving national goals for
144	quality healthcare and facilitating coordinated, multi-stakeholder action. The Department of
145	Health and Human Services has asked the Partnership for its collective, multi-stakeholder input
146	on the National Quality Strategy (NQS) framework, which includes three inextricably linked
1/17	domains_better care, affordable care, and healthy people/healthy communities_around which

148	priorities, goals, measures, and strategic opportunities for improvement are to be identified
149	and/or refined.
150	
151	When the NQS was announced in March 2011, one of the initial priorities identified was
152	"Promoting the Most Effective Prevention and Treatment of the Leading Causes of Mortality,
153	Starting with Cardiovascular Disease." The NQF cardiovascular portfolio contains endorsed
154	process and outcome measures that are being used to track performance and monitor
155	improvements in the priority area of cardiovascular disease.
156	
157	PRIOR NQF WORK RELATED TO CARDIOVASCULAR CONDITIONS
158	Endorsement of Consensus Standards
159	The 31 measures undergoing maintenance review were originally evaluated and endorsed in
160	several projects:
161 162 163 164 165 166 167 168	 National Voluntary Consensus Standards for Hospital Care – An Initial Performance Measure Set 2003 National Voluntary Consensus Standards for Hospital Care: Additional Priority Areas 2005-2006 National Voluntary Consensus Standards for Hospital Care 2007: Additional Performance Measures National Voluntary Consensus Standards for Emergency Care National Voluntary Consensus Standards for Ambulatory Care
170	Patient-Focused Episode of Care Measurement Framework
171	NQF has endorsed a measurement framework for patient-focused episodes of care. The
172	definition for an episode of care is "a series of temporally contiguous healthcare services related
173	to the treatment of a given spell of illness or provided in response to a specific request by the
174	patient or other relevant entity." An episode perspective is required to determine if the delivery
175	system is indeed achieving its intended purpose. This approach allows for care to be analyzed
176	over time and offers a better assessment of the patient's resultant health status. This

Cardiovascular Endorsement Maintenance project used the patient-focused episode of care framework for coronary artery disease (Figure 1) to consider measures in the topic areas of coronary artery disease (CAD), acute myocardial infarction (AMI), and percutaneous coronary intervention (PCI).

Figure 1: Patient-focused episode of care applied to patients with coronary artery disease



NQF'S CONSENSUS DEVELOPMENT PROCESS

NQF's 2010 Cardiovascular Endorsement Maintenance project seeks to endorse measures suitable for both public reporting and quality improvement. This project evaluated both newly submitted measures as well as measures endorsed prior to June 2008 for maintenance review. Within NQF's portfolio of endorsed cardiovascular measures, 41 measures were endorsed after June 2008 (Appendix C) and will undergo maintenance review in 2013.

Evaluating Potential Consensus Standards

New candidate consensus standards were solicited through a Call for Measures in September 2010. Cardiovascular measures endorsed prior to June 2008 were evaluated as part of NQF's

193	routine maintenance processes. Because of the number of measures, the evaluation process was
194	conducted in two phases:
195	Phase 1—coronary artery disease, AMI, and PCI, including treatments, diagnostic studies,
196	interventions, or procedures associated with these conditions
197	Phase 2—hypertension, heart failure, atrial fibrillation, and other heart disease and
198	treatments, diagnostic studies, interventions, or procedures associated with these conditions
199	
200	Using NQF's standard evaluation criteria, the Steering Committee evaluated 57 measures for
201	suitability as voluntary consensus standards for quality improvement and public reporting.
202	Steering Committee work groups initially rated each measure for compliance with the sub-
203	criteria. The entire Steering Committee evaluated each measure based on the four main criteria—
204	importance to measure and report, scientific acceptability of the measure properties, usability,
205	and feasibility—to determine whether the measure met NQF's criteria for endorsement.
206	Measure developers were available during Committee discussions to respond to questions and
207	clarify any issues or concerns. Steering Committee recommendations were determined in a
208	stepwise process:
209	Step 1: Evaluate each measure individually to determine whether it meets the
210	endorsement criteria;
211	Step 2: For measures that meet the endorsement criteria:
212	 Evaluate measure harmonization among related measures, and
213	 Select best measure from among competing measures; and
214	Step 3: Determine final recommendation for endorsement.
215	
216	OVERARCHING MEASURE EVALUATION ISSUES
217	During the Steering Committee's discussion of the measures, several overarching issues emerged
218	and were factored into the Committee's ratings and recommendations for many measures
219	
220	Disparities

NQF REVIEW DRAFT—DO NOT CITE, QUOTE, REPRODUCE, OR CIRCULATE NQF MEMBER comments due August 19, 2011, by 6:00 PM ET; PUBLIC comments due August 12, 2011 by 6:00 PM ET

221	Most initial measure submissions did not provide data addressing disparities or did not
222	sufficiently respond to the disparities questions on the measure submission form. The Committee
223	established disparities as a major priority in the evaluation of measures and required information,
224	and preferably data, on how each measure addresses disparities in order to be recommended for
225	endorsement. Developers submitted additional data stratified by disparities when available.
226	
227	Measures demonstrating very high current performance
228	The Committee noted that several measures have been publicly reported for several years and
229	demonstrate very high performance and little variation such that there is no longer much
230	opportunity for improvement. The Committee believed that removing endorsement from these
231	evidence-based, reliable, and valid measures would send the wrong message and asked if there
232	was an alternative designation.
233	
234	In response to the Committee's concern, the NQF Board of Directors approved a policy in May
235	2011 that established a special category of endorsed measures with "reserve status." To be put
236	on reserve status a measure must be highly credible, reliable, and valid and have high levels of
237	performance with little opportunity for improvement. These measures meet all of the NQF
238	criteria except for one sub-criterion, (1b) relating to an opportunity for improvement.
239	Performance can be reassessed in the future if necessary to ensure that performance does not
240	decline.
241	
242	Related and Competing Measures
243	The Committee noted that multiple measures addressed similar aspects of care, such as use of
244	aspirin or beta blockers for secondary prevention of ischemic vascular disease, and repeatedly
245	suggested that similar measures be consolidated into a single measure that can be used across
246	settings and stratified into populations of interest. The Committee also noted that similar
247	measures are not harmonized. The Committee used NQF's guidance for evaluating related and
248	competing measures to further evaluate similar measures that meet NQF's evaluation criteria.
249	The Committee reviewed side-by-side tables of related measures to select "best-in-class" among
	The committee to the field of blac moles of folded modeling to belief best in class among

250	competing measures and to identify a need for harmonization for related measures. However, the
251	Committee struggled with determining which measures were truly competing or just related,
252	such as several measures had similar numerator specifications but related but different
253	denominators (coronary artery disease or ischemic vascular disease), and whether endorsing an
254	all-or-none composite measure was preferred to endorsing individual measures for the
255	components as well as the composite. Endorsing the composite measure only would reduce the
256	need for harmonization of multiple individual measures, though many of the individual measures
257	are in wide use and retooled for EHRs.
258	
259	Harmonization
260	Because of the large number of similar and related measures, the Committee identified the need
261	for harmonization for the majority of measures under review.
262	However, discussions with measure developers revealed significant challenges in achieving
263	harmonization:
264	• Developers have different approaches and philosophies about measurement.
265	• Review and approval of all changes by a developer's technical panel and organizational
266	leadership take significant time (sometimes several months).
267	• When there are several related measures, the determination of which measure should be
268	the basis for harmonization may be difficult.
269	• Individual measures may be part of a group in use by the developer and changes may
270	cause a measure to be out of alignment with that group.
271	 Trending data may be affected by changes in specifications.
272	• There may be disagreement as to what degree of alignment is needed to achieve
273	harmonization.
274	As noted in the recent NQF Task Force on Harmonization report, harmonization is optimally
275	achieved during development of measures rather than after they have been in use.
276	

Conflicting Guidelines

277

278	The Committee noted that similar measures for intermediate outcomes such as blood pressure
279	(BP) targets, may be based on conflicting guidelines. The Committee recommended that all
280	NQF-endorsed measures align to a single national guideline, such as the Joint National
281	Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure (JNC)
282	for blood pressure measures and the National Heart, Lung, and Blood Institute's Expert Panel or
283	the Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult Treatment
284	Panel [ATP]) for lipids.
285	
286	Composite Measures
287	During this project several new composite measures were submitted for consideration. The
288	Committee encouraged the development of more "all-or-none" composite measures, particularly
289	for groups of processes of care applicable to most patients, such as discharge medications for
290	acute myocardial infarction (AMI), percutaneous coronary intervention (PCI), and heart failure.
291	In response to the Committee's suggestion, the American College of Cardiology Foundation
292	quickly developed and tested two new all-or-none composite measures that were favorably
293	reviewed by the Committee. The Committee identified additional potential composite measures
294	that would enhance the cardiovascular portfolio.
295	
296	Medication Management Measures
297	Committee members noted that medication management measures that evaluate adherence, such
298	as medication possession ratio, are more meaningful measures of medication use for chronic
299	conditions compared to those that capture a single prescription or dispensing of a medication.
300	
301	Outcomes measures
302	The Committee supported NQF's move to more outcome measures and voiced support to
303	broaden the denominator populations to include the largest number of appropriate patients
304	whenever possible
305	
306	Gaps in NQF's Cardiovascular Portfolio

307 During its discussion the Steering Committee identified important gap areas in the cardiovascular care episodes of care framework for further measure development: 308 measures that assess functional status, stability, and symptom control based on patient 309 reported data, particularly those that are likely to reduce emergency department (ED) 310 visits and readmissions and improve quality of life; 311 • better measures of patient education and comprehension of self-management prior during 312 transitions of care; 313 measures of appropriate referral, care coordination and transitions of care; 314 patient safety measures such as diverse reactions to cardiac medications, for example, 315 316 aspirin and warfarin use in patients with coronary artery disease (CAD) and atrial fibrillation (AF); upstream use of clopidogrel in sicker patients who then have 317 complications at surgery; and angioedema with ACEI medications; and 318 measures for effectiveness and outcomes of cardiac rehabilitation that are independent of 319 320 linkage to a certifying organization. Additionally the Committee offered approaches that would focus the cardiovascular portfolio on 321 important aspects of care with fewer measures: 322 • expand the denominator populations whenever appropriate; e.g., ACEI/ARBs for all 323 patients with LVSD, not just AMI+LVSD or HF+LVSD; 324 consolidate measures, for example, a single measure for BP control that can be applied to 325 326 a variety of settings and can be stratified into populations of interest such as CAD or diabetes; and 327 more all-or-none composite measures. 328 RECOMMENDATIONS FOR ENDORSEMENT 329 330 This report presents the results of the evaluation of 57 measures considered under the NQF CDP. Thirtyeight measures are recommended for new or continued endorsement as voluntary consensus standards 331

candidate consensus standards. Hyperlinks are provided:

suitable for public reporting and quality improvement. Evaluation summary tables follow the list of

measures and summarize the results of the Steering Committee's evaluation of and voting on the

332333

334

335	 from each listed measure to the evaluation summary table (control + click on title); 	
336	• from each summary table to the detailed measure specifications and measure submission	
337	information:	
338	• from each summary table to the web page where all materials submitted by the developer or	
339	steward are posted; and	
340	• from each summary table to the web page where the meeting and call summaries, transcripts,	and
341	recordings can be accessed.	
342	As this is a new format for NQF reports, comments/suggestions for improved navigation are	
343	welcome.	
344		
345	CORONARY ARTERY DISEASE -SECONDARY PREVENTION	18
346	Recommended for Endorsement:	18
347	0076 Optimal vascular care	18
348	0073 IVD: Blood Pressure Management	20
349	0068 Ischemic vascular disease (IVD): use of aspirin or another antithrombotic	22
350	0067 Chronic stable coronary artery disease: antiplatelet therapy	24
351	0075 IVD: Complete lipid profile and LDL control <100	26
352	0074 Chronic stable coronary artery disease: lipid control	27
353 354	0066 Chronic stable coronary artery disease: ACE inhibitor or ARB therapy—diabetes or left ventricular systolic dysfunction (LVEF <40%)	
355	0071 Acute Myocardial Infarction (AMI): Persistence of Beta-Blocker Treatment After a Heart Attack	30
356	Not Recommended for Endorsement:	31
357	1486 Chronic stable coronary artery disease: blood pressure control	31
358 359	0070 Chronic Stable Coronary Artery Disease: Beta-Blocker TherapyPrior Myocardial Infarction (MI) (Left Ventricular Systolic Dysfunction (LVEF <40%)	
360	0065 Chronic stable coronary artery disease: symptom and activity assessment	34
361	CAD ACUTE PHASE: AMI AND PCI	35
362	Recommended for endorsement:	35
363	0289 Median time to ECG	35
364	0286 Aspirin at arrival	37
365	0288 Fibrinolytic therapy received within 30 Minutes of ED arrival	38
366	0290 Median time to transfer to another facility for acute coronary intervention	39

367	0132 Aspirin at arrival for acute myocardial infarction (AMI)	41
368	0163 Primary PCI received within 90 minutes of hospital arrival	42
369	0164 Fibrinolytic therapy received within 30 minutes of hospital arrival	43
370	0137 ACEI or ARB for left ventricular systolic dysfunction- acute myocardial infarction (AMI) patients	44
371 372	0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization	46
373	0355 Bilateral cardiac catheterization rate (IQI 25)	49
374 375	0964 Composite Measure: Therapy with aspirin, P2Y12 inhibitor and statin at discharge following PCI in eligible patients	
376	0133 PCI mortality (risk-adjusted)©	.511
377	Recommended for endorsement and placement in reserve status:	54
378	0160 Beta-blocker prescribed at discharge for AMI	
379	0142 Aspirin prescribed at discharge for AMI	55
380	Not recommended:	56
381	961 Composite measure of hospital quality for acute myocardial infarction (AMI)	56
382	0282 Angina without procedure (PQI 13)	58
383 384	1495 P2Y12 Inhibitor at discharge for patients with percutaneous coronary intervention (PCI) (with ster	
385	1493 Aspirin at discharge for patients with percutaneous coronary intervention (PCI)	59
386	1498 Statins at discharge for patients with percutaneous coronary intervention (PCI)	60
387	CARDIAC REHABILITATION	61
388	Not recommended:	61
389 390	1496 Cardiac rehabilitation/secondary prevention (CR) program structure-based measurement set to safety standards for CR programming	
391 392	1494 Cardiac rehabilitation/secondary prevention (CR) program measurement set related to monitoring response to therapy and documenting program effectiveness	-
393 394	1497 Cardiac rehabilitation/secondary prevention (CR) program measurement set to assess risk for adverse cardiovascular events	64
395	960 Cardiac rehabiltation composite	65
396	ATRIAL FIBRILLATION	66
397	Recommended for endorsement:	66
398 399	1524 Assessment of thromboembolic risk factors (CHAD	
100	1525 Chronic anticoagulation therapy	68

401	Not recommended:	69
402 403	1505 Adult patient(s) with atrial fibrillation taking amiodarone that had serum ALT or AST test in last reported months	
404	IMPLANTABLE CARDIOVERTER DEFIBRILLATORS (ICD)	71
405	Recommended for endorsement:	71
406	1522 ACE/ARB therapy at discharge for ICD implant patients with LVSD	71
407	1528 Beta Blocker at discharge for ICD implant patients with a previous MI	72
408	1529 Beta blocker at discharge for ICD implant patients with LVSD	73
409 410	0965 Patients with an ICD implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge	74
411	Not recommended:	75
412	1530 Prophylactic antibiotics prior to ICD (lead or implant) procedure	75
413	HEART FAILURE	76
414	Recommended for endorsement:	76
415	0079 Heart failure: Left ventricular ejection fraction assessment (outpatient setting)	76
416 417	0081 Heart Failure: Angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (therapy for left ventricular systolic dysfunction	
418	0083 Heart failure: Beta-blocker therapy for left ventricular systolic dysfunction	78
419	0162 ACEI or ARB for left ventricular systolic dysfunction—Heart failure (HF) patients	80
420	0358 Congestive heart failure (CHF) mortality rate (IQI 16)	81
421	0277 Congestive heart failure admission rate (PQI 8)	82
422 423	0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization	
424 425	0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure hospitaliza	
426	Recommended for endorsement and placement in reserve status:	86
427	0135 Evaluation of left ventricular systolic function (LVS)	89
428	Not recommended:	90
429	0077 Heart failure: Symptom and activity assessment	90
430	962 Composite measure of hospital quality for heart failure (HF)	91
431	HYPERTENSION	92
432	Recommended for endorsement:	92
433	0018 Controlling high blood pressure	92

134	Not recommended:	93
435	0013 Hypertension: Blood pressure management	93
436	0276 Hypertension admission rate (PQI 7)	95
137		
138	EVALUATION SUMMARY TABLES	
139	CORONARY ARTERY DISEASE –SECONDARY PREVENTION	

440 Measures Recommended for Endorsement:

0076 Optimal vascular care

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of adult patients ages 18 to 75 who have ischemic vascular disease with optimally managed modifiable risk factors (LDL, blood pressure, tobacco-free status, daily aspirin use).

Numerator Statement: Patients ages 18 to 75 with ischemic vascular disease (IVD) who meet all of the following targets from the most recent visit during the measurement period: LDL less than 100, Blood Pressure less than 140/90, Tobacco-Free Status, Daily Aspirin Use (unless contraindicated).

<u>Please note</u>: On July 27,2010, the blood pressure component of this measure was changed for patients with a co-morbidity of diabetes (target less than 140/90). MNCM's technical advisory group recommended this change based on ACCORD results, ICSI's most recent guideline changes (July 2010), and the national meaningful use measures for diabetes blood pressure control. A target of less than 140/90 allows for individualization of patient goals.

On March 9, 2011, the measurement and reporting committee reviewed recent ICSI guideline changes for blood pressure targets for stable coronary artery disease and hypertension and additionally considered the request of the NQF cardiovascular committee and decided to change the blood pressure target to < 140/90 for all IVD patients.

Values are collected as the most recent during the measurement period (January 1 through December 31), with the exception of the LDL value which is collected over a 15 month time span to allow a greater window of time for patients that may not complete a cholesterol test within the 12 month time frame, but do complete a cholesterol test within 15 months (October 1 of the previous year through December 31 of the measurement year).

Denominator Statement: Patients ages 18 to 75 with ischemic vascular disease who have at least 2 visits for this condition over the last 2 years (established patient) with at least 1 visit in the last 12 months.

Exclusions: Valid exclusions include patients who only had one coded visit to the clinic during the last two years, patients who had died during the measurement period, patients who were in hospice during the measurement period, patients who were permanent nursing home residents during the measurement period, or patients who were coded with IVD in error.

Adjustment/Stratification: Risk adjustment for this measure is based on case mix (health plan product). Health plan product was selected because it can serve as a proxy for socioeconomic status, if more specific variables are not available. Socioeconomic status can be a variable in a patient's ability to comply with a treatment plan for achieving the intermediate outcomes that can postpone or prevent the long term complications of cardiovascular disease.

The overall average state-wide distribution of patients across three major insurance types (Commercial, Medicare and MN Healthcare Programs plus Self-pay/Uninsured) is calculated and then each reporting site's patient distribution is adjusted to match the average mix. Rates are re-weighted based on the new distribution of patients and then rates are re-calculated.

Background and Evolution of Risk Adjustment:

MN Community Measurement has been publicly reporting unadjusted ambulatory outcome rates at the clinic site level for several years dating back to 2004. Currently, the lowest level of reporting is at the clinic site and we do not publicly report any practitioner level information. As our state begins moving towards utilizing cost and quality measures to demonstrate value and utilizing these measures for incentive based payment and tiering by health plans, we began to explore risk adjustment of measures used for these purposes.

0076 Optimal vascular care

Our subcommittee of the Board of Directors, the Measurement and Reporting Committee (MARC) has reviewed several methods for risk adjusting these measures. Part of their discussion included the potential use of the risk adjusted measures for public reporting to consumers on our MN HealthScores website. The group agreed that risk adjustment would be more beneficial for tiering and incentive based programs and that there was value in reporting the unadjusted clinic site level rate for consumers for the following reasons: rates reflect actual performance, confusion for consumers in terms of explaining risk adjustment or displaying two rates (adjusted and unadjusted), or creating a mindset that it is acceptable for patients in public programs to have different treatment standards than those with commercial insurance.

There are no current plans to report risk adjusted data on our consumer facing website; however we will provide both adjusted and unadjusted clinic site level rates on our corporate website (pdf format).

Level of Analysis: Clinicians: Group/Practice Type of Measure: Outcome

Data Source: Paper medical record/flowsheet; Electronic Health/Medical Record; Registry data. Many medical groups extract the information from their EMR. Registries can be used as a source of information to create the data file; however, groups must ensure that all of their eligible patients are included. Paper abstraction forms are provided for those clinics who wish to use them as an interim step to creating their data file. All data is uploaded in electronic format (.csv file) to a HIPAA-secure, encrypted, and password-protected data portal.

Measure Steward: MN Community Measurement

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- All-or-none-composite of important care processes and intermediate outcomes.
- Patient-oriented measure; assesses whether an individual patients is meeting important targets.
- In use in Minnesota.

2. Scientific Acceptability of Measure Properties: C-1; P-13; M-5; N-2

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

BP target values have been changing due to recent studies but seem to be <140/90 for most patients. New JNC 8 guidelines
to be released early 2012, at which time the developer will modify the measure specifications accordingly.

3. Usability: C-14; P-7; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Measure in use in Minnesota, reported by a large number of practices and patients.
- There is a need for harmonizaton with measures that address the component elements...

4. Feasibility: C-18; P-3; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Data are generated from the process of care and are easily extracted.
- Very few exclusions and contraindications have been rolled into the definitions.
- Data are carefully audited for naccuracies, errors, and unintended consequences.

Does the Measure Meet Criteria for Endorsement: As submitted: Y-5, N-16

If developer changes BP target to <140/90: Y-19; N-1; A-0

Rationale: This measure meets criteria with conditions --if the specifications are changed to target BP<140/90.

If Applicable, Conditions/Questions for Developer:

Change the BP target to <140/90. Response: MN Community Measurement agrees to align measures to JNC8 going forward.
 We took the Cardiovascular E&M Steering Committee's recommendation to modify the blood pressure target to <140/90 to our

0076 Optimal vascular care

Measurement and Reporting Committee on March 9, and they approved this change. This modification is supported by the 2009 European Hypertension update (cited during the February 15 call), as well as ICSI Guidelines on Hypertension Diagnosis and Treatment, released in November 2010.

Evaluation of Competing and Related Measures

- 0073 IVD: Blood pressure management (NCQA)
- 0068 IVD: Use of aspirin or antithrombotics (NCQA)
- 0067 CAD: Antiplatelet therapy (PCPI)
- 0075 IVD: Complete lipid profile and LDL control <100 (NCQA)
- 0074 Chronic stable coronary artery disease: Lipid control (PCPI)

Several Committee members suggested that the composite measure 0076 would be sufficient to address the outcomes and processes of care for secondary prevention rather than endorsing multiple measures addressing the components that would need harmonization. The Committee discussed the pros and cons of recommending the composite measure only versus the composite measure and individual component measures:

PROS

- The composite focuses on several factors that are all important to the individual patient in a single measure. This is a more challenging, but important, patient-focused goal.
- Reduces the number of measures in this topic area and eliminates redundancy.
- Eliminates the need for harmonization of multiple measures.
- Conserves opportunity/measurement costs.
- The Consensus Standards Approval Committee (CSAC) has been pushing for more challenging, broad, patient-focused measures, rather than continuing with numerous narrowly focused measures.
- Significant harmonization is needed among the individual measures.

CONS

- The individual measures, such as blood pressure control or aspirin use, may be important for end users as stand-alone measures.
- The individual measures that form the Minnesota Community measurement composite have not been evaluated as standalone measures and are not available for multiple users for public reporting or payment programs.
- The lack of uniform availability of an electronic platform necessitates maintenance of measures that can be obtained from different data sources (e.g., claims, EHRs, registries).
- The competing individual measures have been endorsed for several years and are in use in many large programs such as CMS's Physicians Quality Reporting System (PQRS) and NCQA HEDIS.
- Some of the individual measures have been re-tooled as eMeasures for meaningful use.

The Committee did not reach consensus on whether to recommend the composite measure 0076 only: Yes – 10, No-9

RECOMMENDATION: MAINTAIN ENDORSEMENT

441

442

0073 IVD: Blood pressure management

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The percentage of patients 18 years of age and older who were discharged alive with acute myocardial infarction (AMI), coronary artery bypass graft (CABG) or percutaneous coronary interventions (PCI) from January 1–November 1 of the year prior to the measurement year, or who had a diagnosis of ischemic vascular disease (IVD) during the measurement year and the year prior to the measurement year and who had BP reported as under control <140/90.

Numerator Statement: The numerator is the number of patients in the denominator whose most recent blood pressure is adequately controlled during the measurement year. For a patient's BP to be controlled, both the systolic and the diastolic BP must meet the desired threshold of <140/90 mm Hg.

Denominator Statement: Patients 18 years or older as of December 31 of the measurement year who were discharged alive for AMI, CABG or PCI on or between January 1 and November 1 of the year prior to the measurement year or who had a diagnosis of IVD during

0073 IVD: Blood pressure management

both the measurement year and the year prior to the measurement year.

Exclusions: All patients with ESRD, who are pregnant or who had an admission to a non-acute inpatient setting during the

measurement year.

Adjustment/Stratification: No risk adjustment necessary NA

Level of Analysis: Clinicians: Individual; Clinicians: Group

Type of Measure: Intermediate Outcome

Data Source: Paper medical record/flow-sheet; Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical

Record NA; retooled eMeasure

Measure Steward: National Committee for Quality Assurance | 1100 13th Street NW, Suite 1000 | Washington | District Of Columbia | 20005

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Intermediate outcome measure.
- Extensive evidence of benefit for achieving blood pressure control in patients with ischemic vascular disease.
- What is the evidence for BP target of < 140/80?
- Evidence base for elderly population and benefit of taking their systolic to less than 140 is lacking.
- Gap demonstrated with the 10th percentile being 28% and the 90th being 62%.

2. Scientific Acceptability of Measure Properties: C-0; P-16; M-4; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Concern regarding lack of inclusion of home blood pressure measurements
- Measure submission included evidence supporting importance of excluding end stage renal disease patients from this measure; however, they are not listed as an exclusion in the measure specifications.

3. Usability: C-4; P-15; M-1; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Measuring blood pressure only once in the year after a procedure may not be very meaningful in patients with fluctuating BP.
- Step-wise process for identifying patients in medical records; this submission is a hybrid specification and a physician-level measure.

4. Feasibility: C-5; P-13; M-2; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

Burden for public reporting purposes as a hybrid measure if only 50 percent of physicians' offices use electronic health records.

Does the Measure Meet Criteria for Endorsement: <u>Deferred</u> (Based on measure as submitted: Yes -8; No-12) Rationale:

The Steering Committee deferred final evaluation of this measure citing several concerns:

- Remove 140/80 lack of evidence for this target. (140/90 only is in retooled EHR specifications)
- Exclusions for elderly patients or patient's intolerance of lower BP.
- Home monitoring BP not included.
- Specifications for exclusion of ESRD not clear.

If Applicable, Conditions/Questions for Developer:

- NCQA withdraws the <140/80 threshold
- NCQA is very open to reconsideration when JNC8 guidelines are released.
- Will discuss home BP monitoring with NCQA's Committee on Performance Measures (CPM) again.

0073 IVD: Blood pressure management

After reviewing the measure developer's responses, does the measure meet NQF's criteria for endorsement?

The Committee is very concerned with the lack of an upper age limit for this measure. Since NCQA indicated an openness to harmonization with measure 0076 that has an upper age limit of 75 years, the Committee considered harmonization as a condition on recommendation for endorsement:

Recommend as currently specified (BP <140/90, no age limits): Yes-3; No-9

Recommend ONLY IF the measure is harmonized with 0076 as to age (18-75 years): Yes-12; No-1

Developer response: NCQA is agreeable to harmonization, however, they point out that JNC8 guidelines are due in early 2012 and it doesn't make sense to make several changes in a short timeframe. They will discuss the upper age limit with the CPM with an overall good faith attempt to achieve harmonization in 2012.

RECOMMENDATION: MAINTAIN ENDORSEMENT and expect harmonization by mid-summer of 2012.

443

444

0068 Ischemic vascular disease (IVD): use of aspirin or another antithrombotic

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The percentage of patients 18 years and older with ischemic vascular disease who were discharged alive for acute myocardial infarction (AMI), coronary artery bypass graft (CABG), or percutaneous coronary interventions (PCI) from January 1-November 1 of the year prior to the measurement year, or who had a diagnosis of ischemic vascular disease (IVD) during the measurement year and the year prior to the measurement year and who had the following during the measurement year.

-Use of aspirin or another antithrombotic

Numerator Statement: Use of aspirin or another antithrombotic.

Electronic Specification:

Documentation of use of aspirin or another antithrombotic during the measurement year. Refer to Table IVD-D to identify the code for prescribed oral anti-platelet therapy. Refer to Table IVD-E to identify medications for oral anti-platelet therapy.

Medical Record Specification:

Documentation of use of aspirin or another antithrombotic during the measurement year. At a minimum, documentation in the medical record must include a note indicating the date on which aspirin or another antithrombotic was prescribed or documentation of prescription from another treating physician.

Denominator Statement: Patients 18 years or older as of December 31 of the measurement year discharged alive for AMI, CABG, or PCI on or between January 1 and November 1 of the year prior to the measurement year or who had a diagnosis of IVD during both the measurement year and the year prior to the measurement year.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary Level of Analysis: Clinicians: Individual; Clinicians: Group

: Clinicians: Group Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical

Record NA ; <u>retooled eMeasure</u> Measure Steward: NCQA

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Performance gap demonstrated. The 25th percentile has not broken 90%.
- Cost-effective.

2. Scientific Acceptability of Measure Properties: C-2; P-14; M-4; N-1

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

Clearly specified with no significant exclusions.

0068 Ischemic vascular disease (IVD): use of aspirin or another antithrombotic

- Sufficient supplemental reliability and validity documentation was provided.
- Title and description do not match numerator.
- According to the measure developer, exclusions for clinical reasons thought to have been less than 5% aren't listed as an
 exclusion.

3. Usability: C-12; P-7; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

• Overlap with other measures using aspirin or other antithrombotics.

4. Feasibility: C-13; P-7; M-1; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Data will be generated as a byproduct of the care process during healthcare delivery as well as electronically.
- Important to note this measure has been retooled for meaningful use.

Does the Measure Meet Criteria for Endorsement: Y-20; N-1; A-0

Rationale:

- Important, effective care process.
- Gap in care—further opportunity for improvement.

If Applicable, Conditions/Questions for Developer:

- Title and description do not match numerator—developer clarified the description as above.
- Possible unintended consequences due to lack of exclusions

Developer response:

- While some exclusions may be coded and included in administrative data and are relatively easily accessible for chart review, a recent paper by Kmetik et al., indicates that most exclusions are relative. Many of the relative contraindications appear to be either minor in nature, or can be overcome by use of different medications. In terms of exceptions (patients removed from the denominator by the clinician at the time of service), the same research showed that the rates of physician added exceptions were quite low, inconsistent in rate, and many had to come from extensive manual chart review even from an EMR.
 - Codes (like CPT-II codes) that might be used to indicate exceptions are not widely used, and at the present time
 cannot be easily audited for accuracy.
- In addition, the measure allows for physician discretion in prescribing alternative oral anti-platelet therapies when aspirin is contraindicated.
- The performance goal is not 100%.
 - Kmetik KS, O'Toole MF, Bossley H, Brutico CA, Fischer G, Grund SL, Gulotta BM, Hennessey M, Kahn S, Murphy KM, Pacheco T, Pawlson LG, Schaeffer J, Schwamberger PA, Scholle SH, Wozniak G. Exceptions to outpatient quality measures for coronary artery disease in electronic health records. Ann Intern Med. 2011 Feb 15;154(4):227-34.
- Harmonization with 0076 and 0067:
- Developer response: NCQA is open to harmonizing this and other measures with other developers' measures and while in some other areas, PCPI and NCQA measures have been harmonized, no direct harmonization has been performed for CV measures at this time. NCQA and AMA PCPI-ACC_AHA have initiated discussions regarding harmonizing elements within this measure where there is potential for harmonization. Harmonization efforts will continue in areas of exclusions and whether it is possible (and/or alternative strategies) to harmonize denominator conditions (IVD vs. CAD) and the potential risks and benefits to populations being measured. There remain significant differences in the respective measures related to complexity, feasibility, standardization, and medication prescribing. As previously noted, the process for harmonization for most specifications must be carried out in a careful and deliberate manner since changes in specifications can affect both trendability of results as well as affect completeness, accuracy and reliability of data collection.

Evaluation of Competing and Related Measures

- 0076 Optimal vascular care (MNCM)
- 0068 IVD: Use of aspirin or antithrombotics (NCQA)
- 0067 CAD: Anti-platelet therapy (PCPI)

0068 Ischemic vascular disease (IVD): use of aspirin or another antithrombotic

Several Committee members suggested that the composite measure 0076 would be sufficient to address the use of anti-thrombotics along with other important aspects of care. The Committee was divided and did not reach consensus on whether to recommend the composite only.

In comparing measures 0068 and 0067, some Committee members questioned whether these are really competing measures because they have different data collection methods, applicable settings, and exclusions and cover different patients. Additionally:

- IVD is a broader denominator that includes coronary artery disease (CAD), cerebrovascular disease (CVD) and peripheral vascular disease (PAD).
- The evidence for aspirin use is very strong for CAD and CVD, less so for PAD although the guidelines do recommend aspirin in PAD.
- 0067 allows for exclusions, such as warfarin use.

Vote to recommend for endorsement: Yes - 11, No -4

RECOMMENDATION: MAINTAIN ENDORSEMENT

445

0067 Chronic stable coronary artery disease: Antiplatelet therapy

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period who were prescribed aspirin or clopidogrel.

Numerator Statement: Patients who were prescribed aspirin or clopidogrel* within a 12-month period.

*Prescribed may include prescription given to the patient for aspirin or clopidogrel at one or more visits in the measurement period OR patient already taking aspirin or clopidogrel as documented in current medication list.

Denominator Statement: All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period.

Exclusions: Documentation of medical reason(s) for not prescribing aspirin or clopidogrel (e.g., allergy, intolerant, receiving other thienopyridine therapy, bleeding coagulation disorders, receiving warfarin therapy, other medical reasons).

Documentation of patient reason(s) for not prescribing aspirin or clopidogrel (e.g., patient declined, other patient reasons).

Documentation of system reason(s) for not prescribing aspirin or clopidogrel (e.g., lack of drug availability, other reasons attributable to the healthcare system).

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Individual; Clinicians: Group Type of Measure: Process

Data Source: Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical Record; Registry data. This measure, in its previous specifications, is currently being used in the ACCF PINNACLE registry for the outpatient office setting. Retooled eMeasure

Measure Steward: AMA PCPI

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale: 1 a.

- Secondary prevention of coronary artery disease is a high impact aspect of healhcare.
- Quality gap has been extablished.
- This measured process leads to improved health outcomes.

2. Scientific Acceptability of Measure Properties: C-16; P-5; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Well-specified measure.
- Important to monitor the "other" exclusion option to prevent increasing percentages over time that may be misleading.

3. Usability: C-16; P-5; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing

24

0067 Chronic stable coronary artery disease: Antiplatelet therapy

measures)

Rationale:

- Meaningful and easily understandable to providers and consumers.
- Not used yet in public reporting initatives. AHA Get With The Guidelines uses this metric.
- Harmonization will need to be addressed.

4. Feasibility: C-19; P-2; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Data elements are readily available and retreiveable.
- Exlcusions are available with routine evaluation of the data that exist.
- Retooled eMeasure.

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale:

- High impact aspect of healthcare.
- Aspirin as part of a secondary prevention plan is a very important and proven intervention.
- Easy to understand and use this metric.

If Applicable, Conditions/Questions for Developer:

Harmonization with measures 0076 and 0068:

Developer Response: Upon original development of the measure set in 2003 and as part of the 2009 update, patients with chronic stable coronary artery disease were identified as the denominator for the measure set to be consistent with ACC/AHA clinical practice guidelines for patients with chronic stable angina which served as the primary evidence base to support measure development. The specific ICD-9 codes selected for CAD encompass all of the relevant codes in the 410-414 series, as well as procedure codes for patients who have undergone coronary bypass surgery or percutaneous coronary intervention. The 410-414 series of codes have been previously identified by other sources, including the American Heart Association as part of their yearly statistical reports, as representative of patients with coronary heart disease. The measure is limited to the only antiplatelet agents (i.e., aspirin and clopidogrel) recommended by the guideline, as follows: Aspirin should be started at 75 to 162 mg per day and continued indefinitely in all patients unless contraindicated (Class I Recommendation, Level A Evidence). Clopidogrel [is recommended] when aspirin is absolutely contraindicated (Class Ila Recommendation; Level of Evidence B). This represents an update to the previous version of the measure that allowed for aspirin, clopidogrel or a combination of aspirin and extended release dipyridamole and is consistent with changes to the evidence. The Work Group also included denominator exceptions for the measure so that physicians can exclude patients for whom aspirin or clopidogrel is not appropriate. If the patient has been prescribed another type of antithrombotic for valid reasons, the medical reason exception might apply.

Evaluation of Competing and Related Measures

- 0076 Optimal vascular care (MNCM)
- 0068 IVD: Use of aspirin or antithrombotics (NCQA)
- 0067 CAD: Anti-platelet therapy (PCPI)

Several Committee members suggested that the composite measure 0076 would be sufficient to address the use of anti-thrombotics along with other important aspects of care. The Committee was divided and did not reach consensus on whether to recommend the composite only.

In comparing measures 0068 and 0067, some Committee members questioned whether these are really competing measures because they have different data collection methods, applicable settings, and exclusions and cover different patients.

- IVD is a broader denominator that includes coronary artery disease (CAD), cerebrovascular disease (CVD) and peripheral vascular disease (PAD).
- The evidence for aspirin use is very strong for CAD and CVD, less so for PAD though the guidelines do recommend aspirin in PAD.
- 0067 allows for exclusions, such as warfarin use.

0067 Chronic stable coronary artery disease: Antiplatelet therapy

Vote to recommend for endorsement: Yes - 12, No -3

RECOMMENDATION: MAINTAIN ENDORSEMENT

446

0075 IVD: Complete lipid profile and LDL control <100

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The percentage of patients 18 years of age and older who were discharged alive for acute myocardial infarction (AMI), coronary artery bypass graft (CABG), or percutaneous coronary interventions (PCI) from January 1–November 1 of the year prior to the measurement year, or who had a diagnosis of ischemic vascular disease (IVD) during the measurement year and the year prior to measurement year, who had each of the following during the measurement year.

- · Complete lipid profile
- LDL-C control <100 mg/dL

Numerator Statement: A complete lipid profile performed during the measurement year. A LDL-C control result of <100mg/dL using the most recent LDL-C screening test during the measurement year.

Denominator Statement: Patients 18 years of age an older as of December 31 of the measurement year who were discharged alive for AMI, CABG, or PCI on or between January 1 and November 1 of the year prior to the measurement year or who had a diagnosis of IVD during both the measurement year and the year prior to the measurement year.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Individual; Clinicians: Group

Type of Measure: Outcome

Data Source: Paper medical record/flowsheet; Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical

Record; Lab data NA; retooled eMeasure

Measure Steward: NCQA

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

Evidence-based, intermediate outcome.

2. Scientific Acceptability of Measure Properties: C-15; P-6; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Reliability testing is in process and currently not available.
- Clarifications needed in the specifications for the target population's age: 18 years and older or 18 years to 75 years.

3. Usability: C-20; P-0; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Already in use as part of HEDIS measures and will need to be harmonized with other lipid measures.
- Data are generated as a byproduct of care processes during delivery and are available as electronic data.

4. Feasibility: C-20; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Measure has been retooled for EHR meaningful use.

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale:

- LDL <100 in IVD is an accepted standard backed by evidence.
- There is a gap in performance.

0075 IVD: Complete lipid profile and LDL control <100

The measurement is being done, it is feasible, and improvement would likely lead to health benefits.

If Applicable, Conditions/Questions for Developer:

What about intolerance to statins?

Response: While some exclusions to statins are coded and included in administrative data and are relatively easily accessible for chart review, a recent paper by Kmetik et al., indicates that most exclusions are relative so that the majority of patients who have "contraindications" to statins are actually on statins. Many of the relative contraindications (muscle cramping, Gl disturbance, etc.) appear to be either minor in nature, or can be overcome by use of different medications. In terms of exceptions (patients removed from the denominator by the clinician at the time of service), the same research showed that the rates of physician added exceptions were quite low, inconsistent in rate, and many had to come from extensive manual chart review even from an EMR.

In addition this measure is focused on reducing cholesterol, but is not prescriptive about the use of a statin. There are other mechanisms by which cholesterol reduction can be achieved (i.e., modifications in diet, exercise, etc.)

Evaluation of Competing and Related Measures

- 0076 Optimal vascular care (MNCM)
- 0075 IVD: Complete lipid profile and LDL control <100 (NCQA)
- 0074 Chronic stable coronary artery disease: Lipid control (PCPI)

Several Committee members suggested that the composite measure 0076 would be sufficient to address lipid lowering along with other important aspects of care. The Committee was divided and did not reach consensus on whether to recommend the composite only. In comparing measures 0075 and 0074, , some Committee members questioned whether these are really competing measures because they have different data collection methods, applicable settings, and exclusions and cover different patients.

Vote to recommend for endorsement: Yes - 9, No -6

RECOMMENDATION: MAINTAIN ENDORSEMENT

447

0074 Chronic stable coronary artery disease: Lipid control

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Endorsed measure 0074 was originally CAD: Drug therapy for lowering LDL-cholesterol *Percentage of patients with CAD who were prescribed a lipid – lowering therapy (based on current ACC/AHA guidelines)*. Original version is a <u>retooled eMeasure</u>.

Description: Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period who have a LDL-C result <100 mg/dL OR patients who have a LDL-C result ≥100 mg/dL and have a documented plan of care to achieve LDL-C <100mg/dL, including at a minimum the prescription of a statin.

Numerator Statement: Patients who have a LDL-C result <100 mg/dL OR

Patients who have a LDL-C result ≥100 mg/dL and have a documented plan of care* to achieve LDL-C <100 mg/dL, including at a minimum the prescription** of a statin within a 12-month period.

Definitions:

*Documented plan of care may also include: documentation of discussion of lifestyle modifications (diet, exercise); scheduled reassessment of LDL-C.

**Prescribed may include prescription given to the patient for a statin at one or more visits in the measurement period OR patient already taking a statin as documented in current medication list.

Numerator Instructions:

The first numerator option can be reported for patients who have a documented LDL-C < 100 mg/dL at any time during the measurement period.

Denominator Statement: All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period.

Exclusions: Documentation of medical reason(s) for not prescribing a statin (e.g., allergy, intolerance to statin medication(s), other medical reasons).

Documentation of patient reason(s) for not prescribing a statin (e.g., patient declined, other patient reasons).

Documentation of system reason(s) for not prescribing a statin (e.g., financial reasons, other system reasons).

Adjustment/Stratification: No risk adjustment necessary

0074 Chronic stable coronary artery disease: Lipid control

Level of Analysis: Clinicians: Individual; Clinicians: Group

Type of Measure: Process

Data Source: Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical Record; Registry data. This measure, in its previous specifications, is currently being used in the ACCF PINNACLE registry for the outpatient office setting.

Measure Steward: AMA PCPI

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Considerable evidence in terms of opportunity for improvement and impact.
- Performance gaps demonstrated across insured populations and across provider.
- A measure based on clinical guidelines.

2. Scientific Acceptability of Measure Properties: C-9; P-8; M-4; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

• Concerns regarding patient preference type or patient refusal type of exclusion; however, in general, exceptions are used rarely.

3. Usability: C-6; P-11; M-4; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Deomonstrated through multiple quality improvement programs.
- Not in use for public reporting at this time, but will be in the future.
- Additive values need to be addressed, and measure will need to be harmonized with other lipid measures.

4. Feasibility: C-8; P-11; M-1; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

• Data can be extracted electronically.

Does the Measure Meet Criteria for Endorsement: Y-17; N-4; A-0

Rationale:

- Opportunity for improvement.
- Evidence-based, outcome measure.

If Applicable, Conditions/Questions for Developer:

• How are patients who have not had an LDL test performed counted in the measure?

Response: All patients aged 18 years and older with a diagnosis of coronary artery disease must have an LDL-C recorded in order to satisfy the measure. The measure specifications will be clarified that patients who have not had an LDL test performed would not meet the measure.

Evaluation of Competing and Related Measures

- 0076 Optimal vascular care (MNCM)
- 0075 IVD: Complete lipid profile and LDL control <100 (NCQA)
- 0074 Chronic stable coronary artery disease: Lipid control (PCPI)

Several Committee members suggested that the composite measure 0076 would be sufficient to address lipid lowering along with other important aspects of care. The Committee was divided and did not reach consensus on whether to recommend the composite only. In comparing measures 0075 and 0074, , some Committee members questioned whether these are really competing measures because they have different data collection methods, applicable settings, and exclusions and cover different patients.

Vote to recommend for endorsement: Yes – 14, No -1

RECOMMENDATION: MAINTAIN ENDORSEMENT

0066 Chronic stable coronary artery disease: ACE inhibitor or ARB therapy—diabetes or left ventricular systolic dysfunction (LVEF <40%)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period who also have diabetes or a current or prior LVEF <40% who were prescribed ACE inhibitor or ARB therapy.

Numerator Statement: Patients who were prescribed ACE inhibitor or ARB therapy.*

*Prescribed may include prescription given to the patient for ACE inhibitor or ARB therapy at one or more visits in the measurement period OR patient already taking ACE inhibitor or ARB therapy as documented in current medication list.

Denominator Statement: All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period who also have diabetes or a current or prior LVEF <40%.

Exclusions: Documentation of medical reason(s) for not prescribing ACE inhibitor or ARB therapy (e.g., allergy, intolerant, pregnancy, renal failure due to ACE inhibitor, diseases of the aortic or mitral valve, other medical reasons).

Documentation of patient reason(s) for not prescribing ACE inhibitor or ARB therapy (e.g., patient declined, other patient reasons.) Documentation of system reason(s) for not prescribing ACE inhibitor or ARB therapy (e.g., lack of drug availability, other reasons attributable to the healthcare system).

Adjustment/Stratification: No risk adjustment necessary Level of Analysis: Clinicians: Individual; Clinicians: Group

Type of Measure: Process

Data Source: Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical Record; Registry data. This measure, in its previous specifications, is currently being used in the ACCF PINNACLE registry for the outpatient office setting. Retooled eMeasure

Measure Steward: AMA PCPI

1. Importance to Measure and Report: Y-18; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

Very high impact and strong evidence for this measure.

2. Scientific Acceptability of Measure Properties: C-12; P-8; M-1; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Why not include patients with coronary artery disease and hypertension, and patients with coronary artery disease and chronic kidney disease?
- "Most recent LVEF" would be better than "prior LVEF," particularly in recovery from STEMI.
- This is not a patient adherence measure but a provider adherence measure.
- A single point estimate is not ideal to measure use.

3. Usability: C-12; P-9; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Not yet publicly reported; however, it does have a signifigant amount of value if approved as it relates to clinical care.
- This measure will need to be harmonized with hospital measures.

4. Feasibility: C-13; P-8; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Data for this measure are easily extractable.
- Concerns about relative contraindications; however, including an explicit list of contraindications increases abstraction burden and raises clinical acceptability issues.

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale: An important clinical measure; however, a more stringent numerator criteria (i.e., must have X number of refills within defined time frame) would make it a stronger measure.

0066 Chronic stable coronary artery disease: ACE inhibitor or ARB therapy—diabetes or left ventricular systolic dysfunction (LVEF <40%)

If Applicable, Conditions/Questions for Developer:

• Why are patients with CAD + hypertension or CAD + chronic kidney disease not included—these are also indications for ACEI/ARB use?

Response: Whereas the guidelines on which these measures are based list CAD with heart failure or diabetes as specific indications for ACEI, they do not explicitly recommend ARB for patients with HTN or CKD. Because this measure combines ACEI and ARB therapy, including HTN or CKD in the denominator would be problematic with respect to the underlying guideline support for the measure.

RECOMMENDATION: MAINTAIN ENDORSEMENT

449

0071 Acute myocardial infarction (AMI): Persistence of beta-blocker treatment after a heart attack

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The percentage of patients age 18 years and older during the measurement year who were hospitalized and discharged alive July 1 of the year prior to the measurement year through June 30 of the measurement year with a diagnosis of acute myocardial infarction (AMI) and who received persistent beta-blocker treatment for six months after discharge.

Numerator Statement: A 180-day course of treatment with beta-blockers post discharge.

Denominator Statement: Patients 18 years and older as of December 31 of the measurement year discharged alive from an acute inpatient setting with an AMI from July 1 of the year prior to the measurement year through June 30 of the measurement year.

Exclusions: Exclude patients who are identified as having a contraindication to beta-blocker therapy or previous adverse reaction to beta-blocker therapy. Also exclude from the denominator hospitalizations in which the patient was transferred directly to a nonacute care facility for any diagnosis.

Adjustment/Stratification: No risk adjustment necessary NA None

Level of Analysis: Clinicians: Individual; Clinicians: Group; Health Plan Type of Measure: Process

Data Source: Paper medical record/flow-sheet; Electronic administrative data/claims; Pharmacy data; Electronic clinical data; Electronic Health/Medical Record NA

Measure Steward: National Committee for Quality Assurance | 1100 13th Street NW, Suite 1000 | Washington | District Of Columbia | 20005

STEERING COMMITTEE EVALUATION:

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- There is a significant performance gap for persistence of beta-blockers after acute myocardial infarction.
- High-level evidence for use of beta blockers for 1 year after AMI (Level A).

2. Scientific Acceptability of Measure Properties: C-8; P-11; M-2; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Very specific exclusion criteria. Discussion regarding whether the exclusion criteria are too strict.
- HEDIS health plan and clinician-level measure.

3. Usability: C-12; P-0; M-2; N-1

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- The measure is currently in use and publicly reported.
- No known issues on implementation.

4. Feasibility: C4; P-11; M-5; N-1

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

• The data are generated as a byproduct of care proceses during care delivery

0071 Acute myocardial infarction (AMI): Persistence of beta-blocker treatment after a heart attack

- The data elements are all collected electronically, but feasibility for a physician with paper records is questionable.
- Mainly based on pharmacy claims; regarding claims that aren't adjudicated or patients without insurance.

Does the Measure Meet Criteria for Endorsement? Y-13; N-8; A-0

Rationale: Adherence is a better measure than a single point in time assessment. Beta blocker use in the 6-12 months after AMI is strongly evidence-based.

If applicable, Conditions/Questions for Developer:

Clarify age specifications – response: "The measure looks at patients 18 years and older".

Evaluation of Competing and Related Measures:

- 0070 CAD: Beta blocker—prior MI (AMA PCPI)
- 0072 CAD: Beta-blocker treatment after a heart attack (NCQA) —retired by developer in favor of 0071
- 0160 Beta blocker prescribed at discharge [for AMI] (CMS)

The Committee agreed that a measure of adherence to beta blockers after AMI is superior to measuring a single point in time and selected this measure, 0071, as "best-in-class for outpatient measures of beta blocker use. The related hospital measure, 0160, has very high current performance and is recommended for reserve endorsement.

RECOMMENDATION: MAINTAIN ENDORSEMENT

450 451

Measures Not Recommended for Endorsement:

1486 Chronic stable coronary artery disease: Blood pressure control

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period with a blood pressure <140/90 mm Hg OR patients with a blood pressure =140/90 mm Hg and prescribed 2 or more antihypertensive medications during the most recent office visit

Numerator Statement: Patients with a blood pressure <140/90 mm Hg* OR

Patients with a blood pressure =140/90 mm Hg and prescribed** 2 or more anti-hypertensive medications during the most recent office visit

*BP value used for measure calculation:

- •Must be specified in medical record if >1 value (systolic/diastolic) recorded, and
- •Must be value upon which treatment decision was based, and
- •May be obtained by measurement during office visit or review of a home blood pressure log, OR of a 24-hour ambulatory blood pressure monitor, but the value on which the treatment decision is being made and which might represent the average of more than 1 reading must be documented as such in the medical record

**Prescribed may include prescriptions given to the patient for two or more anti-hypertensive medications at most recent office visit OR patient already taking 2 or more anti-hypertensive medications as documented in current medication list. (Each anti-hypertensive component in a combination medication should be counted individually.)

Instructions:

All patients aged 18 years and older with a diagnosis of coronary artery disease must have a measurement of blood pressure recorded in order to satisfy the measure.

Report number of patients for 1st numerator component (outcome) AND

Report number of patients for 2nd numerator component (process) AND

Report total number of patients for all numerator components

Denominator Statement: All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period

Exclusions: Documentation of medical reason(s) for not prescribing two or more antihypertensive medications (e.g., allergy, intolerant, postural hypotension, other medical reasons)

Documentation of patient reason(s) for not prescribing two or more anti-hypertensive medications (e.g., patient declined, other patient reasons)

Documentation of system reason(s) for not prescribing two or more antihypertensive medications (e.g., financial reasons, other reasons attributable to the healthcare delivery system)

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Individual; Clinicians: Group Type of Measure: Process

1486 Chronic stable coronary artery disease: Blood pressure control

Data Source: Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical Record; Registry data Measure Steward: American Medical Association (AMA) PCI

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- The Committee agreed that blood pressure control in this population is extremely important.
- The outcome target is consistent with guidelines, although there is no upper age limit in this measure. The Committee expressed concerns regarding appropriate treatment targets in the elderly.
- The Committee questioned the scientific evidence supporting use of only two drugs. Many Committee members did not agree that two drugs were adequate attempts at BP control in some patients.

2. Scientific Acceptability of Measure Properties: C-2; P-4; M-11 N-4

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Errors in measure submission form were addressed: developers confirmed that the numerator includes patients with BP
 ≥140/90
- Testing has not been completed. No data were provided.

3. Usability: <u>C-2; P-5; M-12; N-2</u>

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Clear need for harmonization.
- Developer stated the measure will be revised to reflect guidelines changes or updates as needed.

4. Feasibility: C-11; P-9; M-0; N-1

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

• Measure includes exceptions that address end stage renal disease and elderly patients.

Does the Measure Meet Criteria for Endorsement: Y-8; N-12; A-0

Rationale: Testing not completed.

RECOMMENDATION: NOT RECOMMENDED FOR ENDORSEMENT

452

0070 Chronic stable coronary artery disease: Beta-blocker therapy--Prior myocardial infarction (MI) or left ventricular systolic dysfunction (LVEF <40%)

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period who also have prior MI or a current or prior LVEF <40% who were prescribed beta-blocker therapy

Numerator Statement: Patients who were prescribed* beta-blocker therapy**

*Prescribed may include prescription given to the patient for beta-blocker therapy at one or more visits in the measurement period OR patient already taking beta-blocker therapy as documented in current medication list

** Beta-blocker therapy:

- •For patients with prior MI, no recommendations or evidence cited in current chronic stable angina guidelines for preferential use of specific agents
- •For patients with prior LVEF <40%, beta-blocker therapy should include bisoprolol, carvedilol, or sustained release metoprolol succinate

Denominator Statement: All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period who also have prior MI or a current or prior LVEF <40%

Exclusions: Documentation of medical reason(s) for not prescribing beta-blocker therapy (eg, allergy, intolerant, bradycardia, AV block

0070 Chronic stable coronary artery disease: Beta-blocker therapy--Prior myocardial infarction (MI) or left ventricular systolic dysfunction (LVEF <40%)

without permanent pacemaker, arrhythmia, hypotension, asthma, other medical reasons)

Documentation of patient reason(s) for not prescribing beta-blocker therapy (eq. patient declined, other patient reasons)

Documentation of system reason(s) for not prescribing beta-blocker therapy (eg, other reasons attributable to the health care system

Adjustment/Stratification: No risk adjustment necessary Level of Analysis: Clinicians: Individual; Clinicians: Group

Type of Measure: Process

Data Source: Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical Record; Registry data This measure, in its previous specifications, is currently being used in the ACCF PINNACLE registry for the outpatient office setting. Retooled eMeasure

Measure Steward: American Medical Association | 515 N. State St. | Chicago | Illinois | 60654

STEERING COMMITTEE EVALUATION:

1. Importance to Measure and Report: Y-17; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Cohort studies have demonstrated significant gaps in care regarding the measure.
- The measure takes into account specific beta blockers mentioned in the guidelines for patients with left ventricular systolic dysfuntion. However, data are lacking on beta blocker therapy with normal left ventricular function, more than three years after a myocardial infarction.

2. Scientific Acceptability of Measure Properties: C-4; P-9; M-2; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Measure can be modified to reflect any changes in the guideline recommendations.
- Exclusions include system reasons for not prescribing the beta blocker therapy. Examples provided: insurance, medication availability, and the availability of local cardiac rehabilitation programs.

3. Usability: C-9; P-10; M-2; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- The measure is already in use but is not in any public reporting initiative.
- Useful measure if it can be revised as needed to be consistent with guidelines.

4. Feasibility: C-9; P-8; M-2; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- Data are generated as part of the care process and are sometimes available from the EHR.
- Sixty-four percent of the submissions were rejected due to an inaccurate diagnoses code. This was an implementation issue that has been addressed.

Does the Measure Meet Criteria for Endorsement? Y-17; N-4; A-0

Rationale: The measure reports performance that has a strong positive impact on lowering mortality among patients with chronic CAD and LVEF <40%. It is in use and feasibility has been documented. Abstraction of the paper record is prone to error, however.

If applicable, Conditions/Questions for Developer:

What is the evidence for beta blocker use beyond 3 years?

Response: The newly released AHA guidelines for the prevention of cardiovascular disease in women do note that "Beta-blockers should be used for up to 12 mo (Class I; Level of Evidence A) or up to 3 y (Class I; Level of Evidence B) in all women after MI or ACS with normal left ventricular function unless contraindicated." As a result of this change to the evidence base, the Work Group will be consulted and any necessary modifications will be made to the measure.

Evaluation of Competing and Related measures:

0070 Chronic stable coronary artery disease: Beta-blocker therapy--Prior myocardial infarction (MI) or left ventricular systolic dysfunction (LVEF <40%)

- 0071 AMI: Persistence of beta blocker therapy (NCQA)
- 0072 CAD: Beta-blocker treatment after a heart attack (NCQA) —retired by developer in favor of 0071
- 0160 Beta blocker prescribed at discharge [for AMI] (CMS)

The Committee agreed that a measure of adherence to beta blockers after AMI is superior to measuring a single point in time and selected measure, 0071, as "best-in-class for outpatient measures of beta blocker use. Measure 0160 is recommended for reserve endorsement.

RECOMMENDATION: REMOVE ENDORSEMENT

453

0065 Chronic stable coronary artery disease: symptom and activity assessment

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period for whom there is documented results of an evaluation of level of activity AND an evaluation of presence or absence of anginal symptoms in the medical record.

Numerator Statement: Patients for whom there are documented results of an evaluation of level of activity AND an evaluation of presence or absence of anginal symptoms* in the medical record.

*Evaluation of level of activity and evaluation of presence or absence of anginal symptoms should include:

- •Documentation of Canadian Cardiovascular Society (CCS) Angina Class OR
- •Completion of a disease-specific questionnaire (eg, Seattle Angina Questionnaire or other validated questionnaire) to quantify angina and level of activity.

Numerator Definition:

Canadian Cardiovascular Society (CCS) Angina Classification

Class 0: Asymptomatic

Class 1: Angina with strenuous exercise

Class 2: Angina with moderate exertion

Class 3: Angina with mild exertion

- 1. Walking 1-2 level blocks at normal pace
- 2. Climbing 1 flight of stairs at normal pace

Class 4: Angina at any level of physical exertion

Denominator Statement: All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12-month period.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Individual; Clinicians: Group

Type of Measure: Process

Data Source: Electronic administrative data/claims; Electronic clinical data; Electronic Health/Medical Record; Registry data. This measure, in its previous specifications, is currently being used in the ACCF PINNACLE registry for the outpatient office setting.

Measure Steward: AMA

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-8; N-13

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Measure introduced as a means to ensure there was documentation of the symptom burden and the activity that precipitated those symptoms. Not an outcome measure.
- Evidence lacking; no documentation of gap.
- Testing data not provided.

Does the Measure Meet Criteria for Endorsement: No --Did not pass Importance to Measure and Report.

The developers submitted a <u>letter to the Steering Committee</u> disagreeing with the Committee's evaluation and requested a

0065 Chronic stable coronary artery disease: symptom and activity assessment

reconsideration of the measure evaluation citing the following:

- "a notable gap in patient-centric measures that would focus attention on patient-reported outcomes, including their symptoms, function and health-related quality of life"; and
- symptoms are an outcome and there are racial disparities in symptom management; they want to lay a foundation for future measures of efficacy and appropriateness.

The Steering Committee agreed that the measure, as specified, is a process measure that is not linked to an intermediate or ultimate outcome. The measure was introduced as a means to ensure documentation of the patient burden and the activity that precipitated those symptoms, and the Committee additionally noted:

- There is no reliability or validity data that say the results distinguish quality at the physician level.
- Evidence is lacking. What are the data/evidence that doing an assessment alone is related to patient satisfaction, better outcomes, more or less angioplasty, or less MIs?
- What is the gap? General perception that clinicians are not doing this well. PINNACLE data = 85.5%.
- Testing data not provided.

Steering Committee re-vote on Importance: Yes – 4, No -11

RECOMMENDATION: REMOVE ENDORSMENT

454

455

456

CORONARY ARTERY DISEASE—ACUTE PHASE: ACUTE MYOCARDIAL

INFARCTION AND PERCUTANEOUS CORONARY INTERVENTION

457 458 459

Recommended for endorsement:

0289 Median time to ECG

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Median time from emergency department arrival to ECG (performed in the ED prior to transfer) for acute myocardial infarction (AMI) or chest pain patients (with probable cardiac chest pain).

Numerator Statement: Continuous Variable Statement:

Time (in minutes) from emergency department arrival to ECG (performed in the ED prior to transfer) for acute myocardial infarction (AMI) or chest pain patients (with Probable Cardiac Chest Pain).

Included Populations:

- ICD-9-CM Principal or other diagnosis code for AMI as defined in Appendix A1, OP Table 6.1, or an ICD-9-CM Principal or other diagnosis code for angina, acute coronary syndrome, or chest pain as defined in Appendix A1, OP Table 6.1a, and
- E/M Code for emergency department encounter as defined in Appendix A1, OP Table 1.0a, and
- Patients receiving an ECG as defined in the Appendix A1, and
- Patients discharged/transferred to a short term general hospital for inpatient care, to a Federal healthcare facility, or to a critical access hospital.

Denominator Statement: Continuous Variable Statement:

Time (in minutes) from emergency department arrival to ECG (performed in the ED prior to transfer) for acute myocardial infarction (AMI) or chest pain patients (with probable cardiac chest pain)

Exclusions: Patients less than 18 years of age.

Adjustment/Stratification: No risk adjustment necessary Level of Analysis: Facility/Agency; Population: national

Type of Measure: Process

Data Source: Paper medical record/flow-sheet; Electronic administrative data/claims; Electronic Health/Medical Record N/A

Measure Steward: Centers for Medicare & Medicaid Services (CMS)

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-17; N-4

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

0289 Median time to ECG

Rationale:

- Large variation in performance in emergency departments.
- Questions raised regarding using the measure of median time as being useful and meaningful as an indicator of performance in an emergency department.
- What is the evidence for other conditions besides STEMI?
- Highest mismatched data element on measure was probable cardiac chest pain. Physician educational sessions and a quality assurance program have been implemented to help reduce error.

2. Scientific Acceptability of Measure Properties: C-7; P-10; M-4; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Steering Committee requested to have disparities information included.
- Concerns regarding reliability and validity. Time stamps on ECG machines are often inaccurate and are not as reliable as time stamps for arrival to ED or for administration of therapy.

3. Usability: C-7; P-12; M-2; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Currently being used in outpatient quality data programs.
- Patients not transferred are not included.

4. Feasibility: C-11; P-8; M-2; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Data are generated as a byproduct of care.

Does the Measure Meet Criteria for Endorsement: Y-17; N-2; A-0

Rationale:

• Important time marker in patients to be transferred. Not as important for patients who will get PCI or fibrinolytic therapy at the same location because this time is included in other measures.

If Applicable, Conditions/Questions for Developer:

What is the evidence for patients other than STEMI needing urgent evaluation?

Response: Current guidelines from the ACCF/AHA for STEMI note that ECG should be completed within 10 minutes for patients with persistent chest pain. You cannot diagnosis a STEMI until the ECG is completed.

Where is Appendix A, OP Table 1.1 referred to in the submission?

Response: Appendix A 1.1 (Acute Myocardial Infarction Diagnosis Codes) is found within the previously submitted documents. The table includes codes:

410.00 Anterolateral wall, acute myocardial infarction—episode of care unspecified

410.01 Anterolateral wall, acute myocardial infarction—initial episode

410.10 Other anterior wall, acute myocardial infarction—episode of care unspecified

410.11 Other anterior wall, acute myocardial infarction—initial episode

410.20 Inferolateral wall, acute myocardial infarction—episode of care unspecified

410.21 Inferolateral wall, acute myocardial infarction-initial episode

410.30 Inferoposterior wall, acute myocardial infarction—episode of care unspecified

410.31 Inferoposterior wall, acute myocardial infarction—initial episode

410.40 Other inferior wall, acute myocardial infarction—episode of care unspecified

410.41 Other inferior wall, acute myocardial infarction—initial episode

410.50 Other lateral wall, acute myocardial infarction—episode of care unspecified

410.51 Other lateral wall, acute myocardial infarction—initial episode

0289 Median time to ECG

410.60 True posterior wall, acute myocardial infarction—episode of care unspecified

410.61 True posterior wall, acute myocardial infarction—initial episode

410.70 Subendocardial, acute myocardial infarction—episode of care unspecified

410.71 Subendocardial, acute myocardial infarction—initial episode

410.80 Other specified sites, acute myocardial infarction—episode of care unspecified

410.81 Other specified sites, acute myocardial infarction—initial episode

410.90 Unspecified site, acute myocardial infarction—episode of care unspecified

410.91 Unspecified site, acute myocardial infarction—initial episode

Please provide data on disparities.

Response: The developer provided detailed tables depicting disparities data for the most recent performance data available.

RECOMMENDATION: MAINTAIN ENDORSEMENT

460

0286 Aspirin at arrival

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of emergency department acute myocardial infarction (AMI) patients or chest pain patients (with probable cardiac chest pain) without aspirin contraindications who received aspirin within 24 hours before ED arrival or prior to transfer.

Numerator Statement: Emergency department AMI or chest pain patients (with probable cardiac chest pain) who received aspirin within 24 hours before ED arrival or prior to transfer.

Denominator Statement: Emergency department AMI or chest pain patients (with probable cardiac chest pain) without aspirin contraindications.

Exclusions: Excluded Populations:

• Patients less than 18 years of age.

• Patients with a documented reason for no aspirin on arrival.

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic administrative data/claims; Electronic Health/Medical Record N/A

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-18; N-3

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- 25% are below 94% -indicates there may be more room for improvement here.
- No clear evidence to say patients outside of those having a myocardial infarction will benefit.

2. Scientific Acceptability of Measure Properties: C-7; P-11; M-3; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- No data provided for disparities.
- Validity is questionable-- about 20% of those patients who were initially identified as meeting criteria were then found to be invalid.

3. Usability: C-14; P-4; M-1; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- In use
- Very similar to measure 0132 for patient not being transferred reported on Hospital Compare

4. Feasibility: C-16; P-4; M-0; N-0

0286 Aspirin at arrival

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions— no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Data elements are easily generated from electronic or chart review.

Does the Measure Meet Criteria for Endorsement: Y-19; N-1; A-0

Rationale: Essentially the same measure as 0132, but applies to patients being transferred.

If Applicable, Conditions/Questions for Developer:

The title and description do not accurately describe what is being measured. Significant explanation from the developer was needed for the Committee to understand the intent of the measure. Using the same name for measures 0132 and 0286 is confusing to audiences, and some may assume they are redundant or competing measures.

Response: This measure includes both AMI and chest pain patients with probable cardiac chest pain. The population is emergency department patients who are transferred out to another facility and subsequently are not captured through measure 0132. This population differs from 0132 as patients with suspected cardiac chest pain are also included in the measure.

Provide data on disparities.

Response: Data tables on disparities provided to the Committee.

RECOMMENDATION: MAINTAIN ENDORSEMENT

461

462

0288 Fibrinolytic therapy received within 30 minutes of ED arrival

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Emergency department acute myocardial infarction (AMI) patients receiving fibrinolytic therapy during the ED stay and having a time from ED arrival to fibrinolysis of 30 minutes or less.

Numerator Statement: Emergency department AMI patients whose time from ED arrival to fibrinolysis is 30 minutes or less.

Denominator Statement: Emergency department AMI patients with ST-segment elevation or LBBB on ECG who received fibrinolytic therapy.

Exclusions: Excluded Populations:

- Patients <18 years of age.
- Patients who did not receive fibrinolytic administration within 30 minutes AND had a reason for delay in fibrinolytic therapy as defined in the Data Dictionary.

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national Type of Measure: Process

Data Source: Paper medical record/flowsheet: Electronic administrative data/claims: Electronic Health/Medical Record. See specifications at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=1196289981244.

Measure Steward: CMS21244-1850 0287 Median time to fibrinolysis

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Median time from emergency department arrival to administration of fibrinolytic therapy in ED patients with ST-segment elevation or left bundle branch block (LBBB) on the electrocardiogram (ECG) performed closest to ED arrival and prior to transfer.

Numerator Statement: Continuous Variable Statement:

Time (in minutes) from emergency department arrival to administration of fibrinolytic therapy in AMI patients with ST-segment elevation or LBBB on the ECG performed closest to ED arrival and prior to transfer.

Denominator Statement: Continuous Variable Statement:

Time (in minutes) from emergency department arrival to administration of fibrinolytic therapy in AMI patients with ST-segment elevation or LBBB on the ECG performed closest to ED arrival and prior to transfer.

Exclusions:

- Patients <18 years of age.
- Patients who did not receive fibrinolytic administration within 30 minutes and had a reason for delay in fibrinolytic therapy.

0288 Fibrinolytic therapy received within 30 minutes of ED arrival

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic administrative data/claims; Electronic Health/Medical Record. See specifications at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=1196289981244.

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

Significant disparities differences noted.

2. Scientific Acceptability of Measure Properties: C-19; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Patients who have long lengths of stay, >120 days, are excluded from this measure. These patients are a small proportion of the patients.
- This is a medium-to-large-hospital measure. Only those with more than 25 AMI cases per year are eligible (even if the number who receive fibrinolytics is small).

3. Usability: C-19; P-2; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

Important and meaningful for public reporting.

4. Feasibility: C-20; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Data can be collected either from electronic health records or chart review.
- Good information provided on susceptibility to inaccuracies, errors, or unintended consequences.
- Developers included a nice discussion of suceptibility to inaccuracies.

Does the Measure Meet Criteria for Endorsement: Y-20; N-0; A-0

Rationale:

- Same measure as 164 but different reporting mechanism for patients being transferred.
- Steering Committee duplicated voting on this measure with measure 164.
- 287 uses the same data as 288 but is presented in a different way. Justification for both is that median times may be more actionable in terms of quality improvement, and proportion facilitates comparisons among sites.
- Evaluation of 287 and 288 is the same (also for 164)

If Applicable, Conditions/Questions for Developer:

• The Committee concluded that 288 and 287 are the same measure with different representation of the results rather than competing measures and should be listed under the same NQF number.

Response: Measures are the same specifications, except 0288 and 0287 capture patients who are seen in the emergency department and are subsequently transferred out to another facility and thus are not captured by measure 0164.

RECOMMENDATION: MAINTAIN ENDORSEMENT as a single measure that includes specifications for the two methods of reporting the same data

463

0290 Median time to transfer to another facility for acute coronary intervention

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Median time from emergency department arrival to time of transfer to another facility for acute coronary intervention.

0290 Median time to transfer to another facility for acute coronary intervention

Numerator Statement: Continuous Variable Statement:

Time (in minutes) from emergency department arrival to transfer to another facility for acute coronary intervention. Included Populations:

- ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 6.1, and
- E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0a, and
- Patients discharged/transferred to a short-term general hospital for inpatient care, to a Federal healthcare facility, or to a Critical Access Hospital, and
- Patients not receiving Fibrinolytic Administration as defined in the Data Dictionary, and
- Patients with Transfer for Acute Coronary Intervention as defined in the Data Dictionary.

Denominator Statement: Time (in minutes) from emergency department arrival to transfer to another facility for acute coronary intervention.

Exclusions:

- Patients <18 years of age.
- Patients receiving fibrinolytic administration as defined in the Data Dictionary.

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national; Can be measured at all levels
Type of Measure: Process Data Source: Paper medical record/flowsheet; Electronic administrative data/claims; Electronic Health/Medical Record N/A

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

Measure supports national efforts on making the transfers more efficiently.

2. Scientific Acceptability of Measure Properties: C-13: P-8; M-0, N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Strictly defined population of patients with AMI/STEMI/LBBB who are specifically transferred for acute coronary intervention.
- Reliability of arrival time documentation considered. Data shows there was 20% error rate in arrival time when it was audited.
- Disparities are not defined but can be captured and calculated. Committee recommended the disparities element be included.

3. Usability: C-13; P-8;, M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

Measure is currently in use, reported, and harmonized.

4. Feasibility: C-0; P-21; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Abstractor collects data and needs to have a clear understaing of data definitions to accurately provide a data report.
- E-specifications not developed yet; funding is pending.
- Susceptibility to error not provided.

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale:

- Addresses timeliness of transfer for intervention.
- In use and harmonized with other measures.

If Applicable, Conditions/Questions for Developer:

The measure needs a better title and description of what is being measured. Response:

Measure Name: Median time to transfer to another facility for acute coronary intervention.

0290 Median time to transfer to another facility for acute coronary intervention

Description: Median time from emergency department arrival to time of transfer to another facility for acute coronary intervention.

RECOMMENDATION: MAINTAIN ENDORSEMENT

464

465

0132 Aspirin at arrival for acute myocardial infarction (AMI)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of acute myocardial infarction (AMI) patients who received aspirin within 24 hours before or after hospital arrival.

Numerator Statement: AMI patients who received aspirin within 24 hours before or after hospital arrival.

Denominator Statement: AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91).

Exclusions:

- •<18 years of age
- •Patients who have a length of stay >120 days.
- •Patients enrolled in clinical trials.
- •Discharged to another hospital on day of or day after arrival.
- •Discharged on day of arrival.
- •Expired on day of or day after arrival.
- •Left against medical advice on day of or day after arrival.
- •Patients with comfort measures only documented on day of or day after arrival.
- •Patients with a documented reason for no aspirin on arrival.

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national; Program: QIO Type of Measure: Process

Data Source: Paper medical record/flow-\sheet; Electronic Health/Medical CMS Abstraction & Reporting Tool (CART). Vendor tools also

available. Retooled eMeasure
Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Performance rates for this measure are very high, and there is not much variability but high impact.
- Early aspirin use has same effectiveness as reperfusion.

2. Scientific Acceptability of Measure Properties: C-19; P-2; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

Well-specified and good reliability and validity data provided.

3. Usability: C-18; P-2; M-1; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Existing measure that is meaningful and useful for public reporting.
- Measure is not harmonized with ambulatory CAD but concentrated on in-patient care of AMI.

4. Feasibility: C-19; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

NQF REVIEW DRAFT—DO NOT CITE, QUOTE, REPRODUCE, OR CIRCULATE

0132 Aspirin at arrival for acute myocardial infarction (AMI)

- Data are readily available and generated in care.
- No additional data sources are required for exclusions.

Does the Measure Meet Criteria for Endorsement: Y-18; N-1; A-0

Rationale:

- Little performance gap, but a large impact.
- Important process of care
- In use; data readily available.

If Applicable, Conditions/Questions for Developer:

• Does taking a daily low-dose aspirin 8 hours before the ED/hospital arrival for AMI count in the numerator?

Response: Yes, patients with documentation in the record of receiving aspirin (any dosage) within 24 hours prior to arrival are included in the numerator.

• What is the aspirin dose and timeframe required to meet the measure?

Response: Aspirin (any dosage) within 24 hours prior to arrival or 24 hours after arrival.

RECOMMENDATION: MAINTAIN ENDORSEMENT

466

467

0163 Primary PCI received within 90 minutes of hospital arrival

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of acute myocardial infarction (AMI) patients with ST-segment elevation or LBBB on the ECG closest to arrival time receiving primary percutaneous coronary intervention (PCI) during the hospital stay with a time from hospital arrival to PCI of 90 minutes or less.

Numerator Statement: AMI patients whose time from hospital arrival to primary percutaneous coronary intervention (PCI) is 90 minutes or less.

Denominator Statement: Principal diagnosis of AMI (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91); and PCI procedure (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal or other procedure code for PCI: 00.66); and ST-segment elevation or LBBB on the ECG performed closest to hospital arrival; and PCI performed within 24 hours after hospital arrival.

Exclusions:

- •<18 years of age.
- •Patients who have a length of stay >120 days.
- ·Patients enrolled in clinical trials.
- •Patients received as a transfer from an inpatient or outpatient department of another hospital.
- •Patients received as a transfer from the emergency/observation department of another hospital.
- •Patients received as a transfer from an ambulatory surgery center.
- •Patient administered fibrinolytic agent prior to PCI.
- •PCI described as non-primary by physician, advanced practice nurse, or physician assistant.
- •Patients who did not receive PCI within 90 minutes and had a reason for delay documented by a physician, advanced practice nurse, or physician assistant (e.g., social, religious, initial concern or refusal, cardiopulmonary arrest, balloon pump insertion, respiratory failure requiring intubation).

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national; Program: QIO

Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic Health/Medical Record CMS Abstraction & Reporting Tool (CART). Vendor

tools also available. Retooled eMeasure

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

0163 Primary PCI received within 90 minutes of hospital arrival

Rationale:

Good evidence and data that early PCI is very important.

2. Scientific Acceptability of Measure Properties: C-19; P-2; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- CDAC comparison to hospital data demonstrates reasonable reliability and validity.
- More data needed on disparities; 7% difference in rates for Caucasians going for PCI in a timely fashion, compared to African Americans.
- Measure excludes very unstable patients and patients transferred from another facility.

3. Usability: C-21; P-0; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

Information produced is meaningful and understandable. Has been used in different registries in the past.

4. Feasibility: C-21; P-0; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions— no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Data elements are easily obtainable through routine care processes.

Does the Measure Meet Criteria for Endorsement: Y-16; N-0; A-0

Rationale:

- Good evidence base.
- Reported on Hospital Compare

If Applicable, Conditions/Questions for Developer:

• How often is the exclusion for "system reason for delay" used? Given the potential for gaming, is this being monitored? Response: Current overall trends in measure numerator and denominator counts do not suggest gaming. There is no increasing trend in the use of this reason data element. In our last analysis, Reason for Delay in PCI was occurring in only 0.9% of cases (1Q10). Nevertheless, yes, this is being monitored.

RECOMMENDATION: MAINTAIN ENDORSEMENT

468

469

0164 Fibrinolytic therapy received within 30 minutes of hospital arrival

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of acute myocardial infarction (AMI) patients with ST-segment elevation or LBBB on the ECG closest to arrival time receiving fibrinolytic therapy during the hospital stay and having a time from hospital arrival to fibrinolysis of 30 minutes or less.

Numerator Statement: AMI patients whose time from hospital arrival to fibrinolysis is 30 minutes or less.

Denominator Statement: Principal diagnosis of AMI (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91); and ST-segment elevation or LBBB on the ECG performed closest to hospital arrival; and fibrinolytic therapy within 6 hours after hospital arrival; and fibrinolytic therapy is primary reperfusion therapy. Exclusions:

- •<18 years of age.</p>
- •Patients who have a length of stay >120 days.
- •Patients enrolled in clinical trials.
- •Patients received as a transfer from an inpatient or outpatient department of another hospital.
- •Patients received as a transfer from the emergency/observation department of another hospital.
- •Patients received as a transfer from an ambulatory surgery center.
- •Patients who did not receive fibrinolytic therapy within 30 minutes and had a reason for delay documented by a physician,

0164 Fibrinolytic therapy received within 30 minutes of hospital arrival

advanced practice nurse, or physician assistant (e.g., social, religious, initial concern or refusal, cardiopulmonary arrest, balloon pump insertion, respiratory failure requiring intubation).

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national; Program: QIO Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic Health/Medical Record CMS Abstraction & Reporting Tool (CART). Vendor

tools also available. Retooled eMeasure

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Performance around 50%.
- The Committee noted signifigant disparities differences: lower for females and patients aged > 75 years.
- Same discussion as for measure 288.

2. Scientific Acceptability of Measure Properties: C-19; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Patients who have long lengths of stay, >120 days, are excluded from this measure. These patients are a small proportion of the patients.
- This is a medium-to-large-hospital measure. Only those with more than 25 AMI cases per year are eligible (even if the number who receive fibrinolytics is small).

3. Usability: C-19; P-2; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

• Important and meaningful for public reporting.

4. Feasibility: C-20; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Data can be collected either from electronic health records or chart review.
- Good information provided on susceptibility to inaccuracies, errors, or unintended consequences.
- Developers included a nice discussion of suceptibilty to inaccuracies.

Does the Measure Meet Criteria for Endorsement?: Y-20; N-0; A-0

Rationale:

- Disparities differences.
- Rates highly on all four criteria.

If Applicable, Conditions/Questions for Developer: See discussion of measure 0288

RECOMMENDATION: MAINTAIN ENDORSEMENT

470

471

0137 ACEI or ARB for left ventricular systolic dysfunction- acute myocardial infarction (AMI) patients

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

0137 ACEI or ARB for left ventricular systolic dysfunction- acute myocardial infarction (AMI) patients

Description: Percentage of acute myocardial infarction (AMI) patients with left ventricular systolic dysfunction (LVSD) who are prescribed an ACEI or ARB at hospital discharge. For purposes of this measure, LVSD is defined as chart documentation of a left ventricular ejection fraction (LVEF) less than 40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction.

Numerator Statement: AMI patients who are prescribed an ACEI or ARB at hospital discharge.

Denominator Statement: AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91); with chart documentation of a left ventricular ejection fraction (LVEF) <40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction.

Exclusions:

- •<18 years of age.
- •Patients who have a length of stay >120 days.
- •Discharged to another hospital.
- Expired.
- •Left against medical advice.
- •Discharged to home for hospice care.
- •Discharged to a healthcare facility for hospice care.
- •Patients with comfort measures only documented.
- •Patients enrolled in clinical trials.
- •Patients with a documented reason for no ACEI and no ARB at discharge.

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national; Program: QIO

Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic Health/Medical Record CMS Abstraction & Reporting Tool (CART). Vendor

tools also available. Retooled eMeasure

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- High-impact measure.
- Strong evidence with multiple randomized trials showing ACE inhibitors reduce morbidity and mortality in post MI patients with LVSD and ARBs are shown to be good alternative for patients unable to tolerate ACE inhibitors.
- Concern regarding assumptions made on samples and bias to better results with voluntarily reported data.

2. Scientific Acceptability of Measure Properties: C-18; P-3; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- This is a measure of inpatient performance and is not a subset of measure 0066, which is a measure of outpatient performance.
- Reliability has been tested and documented to be adequate. Face validity is adequate.
- Almost 62% of exclusions were due to undocumented EF or description of LV dysfunction.
- Disparities can be identified but appear not to be present.

3. Usability: C-19; P-2; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

- Useful for public reporting and quality improvement.
- This is the only inpatient ACEI/ARB measure.
- 4. Feasibility: C-21; P-0; M-0; N-0

0137 ACEI or ARB for left ventricular systolic dysfunction- acute myocardial infarction (AMI) patients

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- The data are collected during the process of care.
- Abstraction can lead to errors of exclusion and inclusion, but efforts to limit these errors are continuous.
- The data collection system is already in use and does not impose an undue burden.

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale:

Strong evidence of benefit.

If Applicable, Conditions/Questions for Developer:

• There are a large number of excluded patients due to lack of assessment of LVEF. Is this a quality problem?

Response: Uncertain. The ACC/AHA STEMI/NSTEMI Performance Measure set includes an LVSF Evaluation specific to AMI patients. The Heart Care team has recommended addition of such a measure. Issue is currently under discussion at CMS.

RECOMMENDATION: MAINTAIN ENDORSEMENT

472

473

0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The measure estimates a hospital-level risk-standardized mortality rate (RSMR), defined as death from any cause within 30 days after the index admission date, for patients discharged from the hospital with a principal diagnosis of AMI.

Numerator Statement: This outcome measure does not have a traditional numerator and denominator like a core process measure (e.g., percentage of adult patients with diabetes aged 18-75 years receiving one or more hemoglobin A1c tests per year); thus, we are using this field to define the outcome.

The outcome for this measure is 30-day all-cause mortality. We define mortality as death from any cause within 30 days of the index admission date for patients discharged from the hospital with a principal diagnosis of AMI.

Denominator Statement: Note: This outcome measure does not have a traditional numerator and denominator like a core process measure; thus, we are using this field to define the patient cohort.

The cohort includes admissions for Medicare FFS beneficiaries age 65 years or older discharged from the hospital with a principal diagnosis of AMI (ICD-9-CM codes 410.xx except for 410.x2) and with a complete claims history for the 12 months prior to admission. Patients who are transferred from one acute care facility to another must have a principal discharge diagnosis of AMI at both hospitals. The initial hospital for a transferred patient is designated as the responsible institution for the episode.

If a patient has more than one AMI admission in a year, one hospitalization is randomly selected for inclusion in the measure. Exclusions: The measures exclude admissions for patients:

- Who were discharged on the day of admission or the following day and did not die or get transferred (because it is less likely they had a significant AMI).
- Who were transferred from another acute care hospital (because the death is attributed to the hospital where the patient was initially admitted).
- With inconsistent or unknown mortality status or other unreliable data (e.g., date of death precedes admission date).
- Enrolled in the Medicare Hospice program any time in the 12 months prior to the index hospitalization including the first day of the index admission (since it is likely these patients are continuing to seek comfort measures only).
- Who were discharged alive and against medical advice (AMA) (because providers did not have the opportunity to deliver full care and prepare the patient for discharge).
- Who were not the first hospitalization in the 30 days prior to a patient's death. We use this criteria to prevent attribution of a death to two admissions.

Adjustment/Stratification: Risk adjustment devised specifically for this measure/condition. Our approach to risk adjustment was tailored to and appropriate for a publicly reported outcome measure, as articulated in the American Heart Association (AHA) Scientific Statement, "Standards for Statistical Models Used for Public Reporting of Health Outcomes" (Krumholz et al., 2006).

The measure employs a hierarchical logistic regression model (a form of hierarchical generalized linear model [HGLM]) to create a

0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization

hospital level 30-day RSMR. This approach to modeling appropriately accounts for the structure of the data (patients clustered within hospitals), the underlying risk due to patients' comorbidities, and sample size at a given hospital when estimating hospital mortality rates. In brief, the approach simultaneously models two levels (patient and hospital) to account for the variance in patient outcomes within and between hospitals (Normand et al., 2007). At the patient level, each model adjusts the log-odds of mortality within 30 days of admission for age, sex, selected clinical covariates, and a hospital-specific intercept. The second level models the hospital-specific intercepts as arising from a normal distribution. The hospital intercept, or hospital specific effect, represents the hospital contribution to the risk of mortality, after accounting for patient risk and sample size, and can be inferred as a measure of quality. The hospital-specific intercepts are given a distribution in order to account for the clustering (non-independence) of patients within the same hospital. If there were no differences among hospitals, then after adjusting for patient risk, the hospital intercepts should be identical across all hospitals. Candidate and Final Risk-adjustment Variables: Candidate variables were patient-level risk-adjustors that are expected to be predictive of mortality, based on empirical analysis, prior literature, and clinical judgment, including demographic factors (age, sex) and indicators of comorbidity and disease severity. For each patient, covariates were obtained from Medicare claims extending 12 months prior to and including the index admission. The model adjusted for case differences based on the clinical status of the patient at the time of admission. We used condition categories (CCs), which are clinically meaningful groupings of more than 15,000 ICD-9-CM diagnosis codes. We did not risk adjust for CCs that were possible adverse events of care and that were only recorded in the index admission. In addition, only comorbidities that conveyed information about the patient at that time or in the 12 months prior, and not complications that arose during the course of the hospitalization were included in the risk adjustment.

The final set of risk-adjustment variables are:

Demographic

- Age-65 (years above 65, continuous)
- Male
- Cardiovascular
- History of PTCA
- History of CABG
- Congestive heart failure
- · History of AMI
- Unstable angina
- Anterior myocardial infarction
- Other location of myocardial infarction
- Chronic atherosclerosis
- Cardio-respiratory failure and shock
- Valvular and rheumatic heart disease

Comorbidity

- Hypertension
- Stroke
- · Cerebrovascular disease
- Renal failure
- · Chronic Obstructive Pulmonary Disease
- Pneumonia
- Diabetes and DM complications
- Protein-calorie malnutrition
- · Dementia and senility
- · Hemiplegia, paraplegia, paralysis, functional disability
- · Peripheral vascular disease
- Metastatic cancer, acute leukemia, and other severe cancers
- · Trauma in the last year
- · Major psychiatric disorders
- Chronic liver disease

References:

Krumholz HM, Brindis RG, Brush JE, et al., Standards for statistical models used for public reporting of health outcomes: an American Heart Association Scientific Statement From the Quality of Care and Outcomes Research Interdisciplinary Writing Group: cosponsored

0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization

by the Council on Epidemiology and Prevention and the Stroke Council Endorsed by the American College of Cardiology Foundation, *Circulation*, 2006;113:456-462.

Normand S-LT, Shahian DM, Statistical and clinical aspects of hospital outcomes profilin, *Stat Sci*, 2007;22(2):206-226. Results of this measure will not be stratified.

Level of Analysis: Facility/Agency Type of Measure: Outcome

Data Source: Electronic administrative data/claims. Two data sources were used to create the measure:

- 1. Medicare Part A Inpatient and Outpatient and Part B outpatient claims: This database contains claims data for fee-for-service inpatient and outpatient services including: Medicare inpatient hospital care, outpatient hospital services, skilled nursing facility care, some home health agency services, and hospice care, as well as inpatient and outpatient claims for the 12 months prior to an index admission.

 2. Medicare Enrollment Database (EDB): This database contains Medicare beneficiary demographic, benefit/coverage, and vital status
- information. This dataset was used to obtain information on several inclusion/exclusion indicators such as Medicare status on admission as well as vital status. These data have previously been shown to accurately reflect patient vital status (Fleming, Fisher, et al., 1992). The measure was originally developed with claims data from 1998. The models have been maintained and re-evaluated each year since public reporting of the measures began in 2007.

Fleming C., Fisher ES, Chang CH, et al., Studying outcomes and hospital utilization in the elderly: the advantages of a merged data base for Medicare and Veterans Affairs Hospitals, *Medical Care*, 1992;30(5):377-391.

Measure Steward: CMS

STEERING COMMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- This is an important indicator, as mortality rates after MI are high...
- There is wide variation in performance among hospitals, and this variation persists after adjustment for patient-level characteristics.

2. Scientific Acceptability of Measure Properties: C-19; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- The measure is precise.
- Reliability demonstrated in split-half analysis. Validity demonstrated by chart-based audit.
- Fully risk adjusted with hierarchical general linear modeling.
- Analysis indicates that disparities are small at the hospital level.
- Limited to patients great than 65 years.

3. Usability: C-18; P-2; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- The measure is publicly reported.
- The statistical adjustment method is the same one used for heart failure and pneumonia.
- AHRQ reports in-hospital mortality, but 30-day mortality is independent of length of stay and cannot be influenced by care
 decisions like early discharge.

NOTE: Developer indicates it is working on expanding the age range to include all patients in the near future.

4. Feasibility: <u>C-20; P-0; M-0; N-0</u>

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

- Data are byproduct of routine medical record coding.
- Data are available electronically, and no additional sources are required.

0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization

Measure is already in use.

Does the Measure Meet Criteria for Endorsement: Y-18; N-0; A-0

Rationale:

- Risk-adjusted outcome measure.
- Well developed and tested.
- In use for public reporting.
- Complete measure information in submission, including disparities data.

If Applicable, Conditions/Questions for Developer:

• Developer indicated it is working on expanding the measure to apply to all patients, not just those over 65 years. On June 3, 2011 the developer forwarded testing results for the AMI 30 day mortality applied to all payer data. The Committee will review these results in the coming months and perform a full evaluation as an addendum.

RECOMMENDATION: MAINTAIN ENDORSEMENT

On June 3, 2011 NQF and the Steering Committee received initial results of testing this measure on all payer data. The Committee will further evaluate the testing results as an addendum to this recommendation.

474

475

0355 Bilateral cardiac catheterization rate (IQI 25)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percent of discharges with heart catheterizations in any procedure field with simultaneous right and left heart (bilateral) catheterizations.

Numerator Statement: Discharges with ICD-9-CM procedure code for right and left heart catheterization in any procedure code field.

Denominator Statement: Discharges with ICD-9-CM procedure code for heart catheterizations in any procedure code field.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary. None Observed (raw) rates may be stratified by gender, age groups, race/ethnicity categories, and payer categories.

Risk adjustment of the data is recommended using age and sex. Reliability adjustment is also recommended.

Level of Analysis: Facility/Agency Type of Measure: Outcome

Data Source: Electronic administrative data/claims; Hospital administrative discharge data. See data requirements in the AHRQ QI

Windows Application Documentation: http://www.gualityindicators.ahrg.gov/software.htm

Measure Steward: AHRQ

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-18; N-3

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Recently modified to add the list of procedure indications. Implemented in Version 4.0 of IQI software. An indicator of overuse or unnecessary procedure or a component of a procedure performed without appropriate indications.
- Downward trend over past 10 years resulted from changes in the specifications.

2. Scientific Acceptability of Measure Properties: C-10; P-9; M-2; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

- Looks at heart catherizations in any procedure field but only to include cases with coronary disease.
- Long list of exclusions including diagnoses that would lead to an indication for right heart catherization.
- Reliability and validity testing have been done using large databases.
- Disparaties across payers probably reflect difference across ages.
- There is a 1.3% difference in the rate of inappropriate right heart catherizations between the 5th and 95th percentile.
- Steering Committee interested in seeing more recent regional variation data...

0355 Bilateral cardiac catheterization rate (IQI 25)

3. Usability: C-15; P-5; M-0; N-1

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Measures in use across multiple states and national reporting agencies.
- No harmonization issues are apparent.

4. Feasibility: C-17; P-4; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale

Data are collected from coding; easily obtainable from electronic record sounces.

Does the Measure Meet Criteria for Endorsement: <u>Y-17; N-3; A-0</u> Rationale:

- An indicator of overuse; looking at appropriateness.
- Hospital-level measure.

If Applicable, Conditions/Questions for Developer:

RECOMMENDATION: MAINTAIN ENDORSEMENT

476 477

0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Patients undergoing PCI who receive prescriptions for all medications (aspirin, P2Y12 and satins) for which they are eligible for at discharge.

Numerator Statement: Patients who receive all medications for which they are eligible.

Aspirin prescribed at discharge (if eligible for aspirin as described in denominator)

AND

P2Y12 agent (clopidogrel, prasurgel, or ticlopidine) prescribed at discharge (if eligible for P2Y12 as described in denominator) AND

Statin prescribed at discharge (if eligible for statin as described in denominator).

Denominator Statement: All patients surviving hospitalization who are eligible to receive any one of the three medication classes:

- Eligible for aspirin (ASA): Patients undergoing PCI who do not have contraindication to aspirin documented OR
- 2. Eligibility for P2Y12 agent (clopidogrel, prasurgel, or ticlopidine): Patients undergoing PCI with stenting who do not have a contraindication to P2Y12 agent documented

OR

Eligibility for statin therapy: Patients undergoing PCI who do not have a contraindication to stain therapy.

Exclusions: Discharge statue of expired; not eligible for aspirin, P2Y12, or statin (contraindicated or blinded to all 3 medications).

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility

Type of Measure: Composite with component measures combined at patient level.

Data Source: Registry Data http://www.ncdr.com/WebNCDR/ELEMENTS.ASPX

Measure Steward: American College of Cardiology Foundation, 2400 N. Street NW, Washington, DC 20037

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

- Developed as a result of the Phase I in-person meeting
- Performance gap higher with composite
- High impact and solid evidence

0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge

2. Scientific Acceptability of Measure Properties: C-16; P-4; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

Includes FDA approved drugs

3. Usability: C-19; P-1; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Used in cath labs already
- Adds value to existing measures as a composite.

4. Feasibility: C-19; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

Burden for public reporting purposes as a hybrid measure if only 50 percent of physicians' offices use electronic health

Does the Measure Meet Criteria for Endorsement: Yes -18; No-1

Rationale:

Exclusions possible if LDL is low

If Applicable, Conditions/Questions for Developer:

RECOMMENDATION: Recommend for endorsement

478

479

0133 PCI mortality (risk-adjusted)©

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Risk-adjusted PCI mortality rate.

Numerator Statement: Patients 18 years of age and older with a PCI procedure performed during admission who expired.

Denominator Statement: Patients 18 years of age and older with a PCI procedure performed during admission.

Exclusions: 1. NCDR Registry patients who did not have a PCI (Patient admissions with a diagnostic cath only during that admission);

- 2. Data submissions that do not pass the data quality and completeness reports.
- 3. Procedure variables for subsequent PCIs during the same admission (if the patient had more than one PCI procedure during that
- 4. Patient admissions with PCI who transferred to another facility on discharge.
- 5. Patient admissions with PCI who have more than two variables in the risk model that are missing.

Adjustment/Stratification: Risk-adjustment devised specifically for this measure/condition. Risk adjustment methodology is a logistic

Weights were assigned to risk factors or variables reflecting the strength of their association to PCI in-hospital mortality. Each patient in a facilities submission is given a risk score to predict risk of in hospital mortality and accurately report risk-adjusted mortality rates during hospitalization.

Data from 181,775 procedures performed from January 2004 to March 2006 were used to develop risk models based on pre-procedural and/or angiographic factors using logistic regression.

The most noteworthy risk factors or variables in the model include:

1. ST-segment elevation MI defined as a patient who had a STEMI on admission, with an onset within 24 hours, or the procedure indication was primary, rescue, or facilitated PCI.

0133 PCI mortality (risk-adjusted)©

- 2. Discharge status (alive or expired). The interaction between this variable with other variables were key in the analysis.
- 3. The glomerular filtration rate (GFR) variable is calculated using abbreviated MDRD formula [GFR = $186 \times ?(last creatinine)-1.154 \times (age)-0.203 \times (gender factor) \times (race factor)$ where (gender factor) = 1 for male and 0.742 for female, (race factor) = 1.21 for black and 1 for others].
- 4. The body mass index (BMI) (kg/m²) is calculated from height (cm) and weight (kg): BMI = weight × 10000 / (height) 2.

All Risk Adjustment Variables

STEMI patients:

Age (for age ≤70, for age >70) Cardiogenic Shock at Admission

Previous History—CHF

Peripheral Vascular Disease

Chronic Lung Disease

GFR (for STEMI, for non-STEMI)

NYHA Class IV (for STEMI, for non-STEMI)

PCI Status (for STEMI, for non STEMI)

- Urgent
- Emergency
- Salvage

Previous Vascular Disease

Cerebrovascular Disease

Previous PCI

PreOp IABP

Ejection Fraction Percentage

Coronary Lesion ≥50%: Subacute

Thrombosis? Yes vs. No

Highest Risk Pre-Procedure TIMI Flow = None vs. Yes

1.19 1.02 1.38 4.84

Diabetes/Control (Non-Insulin Diabetes vs. No Diabetes; Insulin Diabetes vs. No Diabetes)

Highest Risk Lesion: SCAI Lesion Class (II or III vs. I; IV vs. I)

BMI [kg/m²] (for STEMI, for Non-STEMI)

Highest Risk Lesion - Segment Category (for STEMI, for non STEMI)

-pRCA/mLAD/pCIRC

-pLAD

-Left Main N/A

Level of Analysis: Facility/Agency Type of Measure: Outcome

Data Source: Registry data National Cardiovascular Data Registry (NCDR) CathPCI Registry®

Measure Steward: ACC

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Outcome measure; is a very frequently performed procedure that can have major impact on patients' lives.
- Expensive procedure so information and knowledge about how centers are performing and where improvements can be made are very important.
- There is a gap in terms of mortality after PCI among different hospitals, and database allows hospitals to compare themselves against each other and against a national baseline.
- Goal is to have a composite measure.

2. Scientific Acceptability of Measure Properties: C-13; P-7; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

0133 PCI mortality (risk-adjusted)©

- Concerns included: data submissions that don't pass a data quality and completeness assessment are excluded; the fact that
 excluding reports because of completeness might bias the mortality to be lower than it actually it is; how to handle patients
 taken back for a second procedure as a result of a poorly performed first procedure.
- Transfers excluded; can lower mortality by transferring to another facility; however, this includes only about 0.7%.

3. Usability: C-8; P-12; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Has been in use by many hospitals.
- Outpatient sites are not captured in registry.

4. Feasibility: C-21; P-0; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Data are available and retrievable.

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale:

- Includes all PCIs performed (30% with AMI; 70% "elective")—data from NCDR registry.
- In-patient mortality—outpatient sites not captured in the registry.

Implementation Comment Received:

• Concern over the definition of PCI status of "salvage" and how it reflects in the PCI RAM model which does not accurately reflect each facilities patient population.

Developer Response to Implementation Comment:

• The definition of a PCI salvage procedure: The definition of salvage in the v3 dataset was harmonized with the definition in the Society for Thoracic Surgery dataset. It was revised in the v4 dataset in 2009 because the committees that develop, review and approve the data elements felt the previous definition was inadequately precise for use for a non-surgical procedure. After implementation of the more focused definition, there was a slightly lower aggregate rate of salvage cases in the registry (0.4% with the v3 dataset compared with 0.3% with the v4 dataset).

	v3 dataset (2005-July 2009)	v4 dataset (July 2009-present)
Definition of PCI Salvage	The patient is undergoing CPR en route to the Cardiac Cath Lab or prior to procedure.	The procedure is a last resort. The patient is in cardiogenic shock at the start of the procedure. Within the last ten minutes prior to the start of the procedure the patient has also received chest compressions for a total of at least sixty seconds or has been on unanticipated extracorporeal circulatory support (e.g. extracorporeal membrane oxygenation, cardiopulmonary support)

Currently, the CathPCI Registry Steering Committee is finalizing a dataset update for 2012. They have proposed several additional variables to further refine case severity to capture cases that may fall between the emergency and salvage status. This includes additional elements describing <u>cardiac arrest</u>, as well as <u>neurologic status</u> of the patient at the start of the procedure.

PCI RAM model revision – Duke Clinical Research Institute, with the oversight of a workgroup of physicians, revised the PCI RAM model using v4 data. The model was approved by the NCDR committees for inclusion in the CathPCI 2011 q2 report. The revised model includes new elements not available in v3 dataset (such as cardiac arrest). The model also included a new 6-level variable matrix, combining PCI status and presence of shock. The model provides predicted mortality ranging from 0.2% for a patient undergoing elective PCI with no shock, to 71% for a patient with shock undergoing salvage PCI. The model accurately predicts mortality, as evidenced by a C-index of 0.934.

The next step for this workgroup is to apply and study the model performance in subsets of the PCI population, such those at particularly high risk for death, or in groups of hospitals, such as STEMI referral centers.

Competing and related measures:

0133 PCI mortality (risk-adjusted)©

- 535: 30-day all-cause risk-standardized mortality rate of percutaneous coronary intervention (PCI) for patients without ST segment elevation myocardial infarction (STEMI) and without cardiogenic shock (CMS)
- 536:30-day all-cause risk-standardized mortality rate of Percutaneous Coronary Intervention (PCI) for patients with ST segment elevation myocardial (STEMI) or cardiogenic shock (CMS)

The variables in all three measures are harmonized in that they use the same clinical registry data elements and definitions (derived from the NCDR CathPCI Registry). Related measures, not competing.

RECOMMENDATION: MAINTAIN ENDORSEMENT

480

Measures recommended for endorsement and placement in reserve status:

481 482

0160 Beta-blocker prescribed at discharge for AMI

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of acute myocardial infarction (AMI) patients who are prescribed a beta-blocker at hospital discharge. Numerator Statement: AMI patients who are prescribed a beta-blocker at hospital discharge.

Denominator Statement: AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91).

Exclusions: Exclusions

- •<18 years of age.
- •Patients who have a length of stay >120 days.
- •Patients enrolled in clinical trials.
- Discharged to another hospital.
- Expired.
- ·Left against medical advice.
- •Discharged to home for hospice care.
- •Discharged to a healthcare facility for hospice care.
- •Patients with comfort measures only documented
- •Patients with a documented reason for no beta blocker at discharge.

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national; Program: QIO Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic Health/Medical Record CMS Abstraction & Reporting Tool (CART). Vendor

tools also available. Retooled eMeasure.

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-0; N-21

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Important measure in terms of reducing morbidity and mortality; ongoing use is designed to ensure high performance.
- Very high performance, concern about not being enough room for improvement to justify the effort.

Steering Committee asked about a special category for good, important measures that seem to be "topped out". In May 2011, the NQF Board approved a policy for a special category "reserve measures".

Committee re-voted on Importance except for 1b, opportunity for improvement:

Y-21; N-0

2. Scientific Acceptability of Measure Properties: C-14; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

0160 Beta-blocker prescribed at discharge for AMI

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3. Usability: C-11; P-4; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

• Unless there is another way to get at the question of disparities identified by the TAP analyses, reserve status appears to be the most cost effective option for this measure.

4. Feasibility: C-14; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Widely accepted and in use by CMS

Does the Measure Meet Criteria for Endorsement: Y-15; N-0; A-0 Rationale: Meets all criteria except for opportunity for improvement

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: MAINTAIN ENDORSEMENT AND PLACEMENT IN RESERVE STATUS

Additional recommendation: The Steering Committee also recommends the measure be recalculated again in 3-5 years to monitor performance.

483

0142 Aspirin prescribed at discharge for AMI

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of acute myocardial infarction (AMI) patients who are prescribed aspirin at hospital discharge Numerator Statement: AMI patients who are prescribed aspirin at hospital discharge.

Denominator Statement: AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91).

Exclusions:

- •<18 years of age.
- •Patients who have a length of stay >120 days.
- •Patients enrolled in clinical trials.
- •Discharged to another hospital.
- Expired.
- ·Left against medical advice.
- •Discharged to home for hospice care.
- •Discharged to a healthcare facility for hospice care.
- •Patients with comfort measures only documented.
- Patients with a documented reason for no aspirin at discharge.

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency; Population: national; Program: QIO Type of Measure: Process

Data Source: Paper medical record/flowsheet; Electronic Health/Medical Record CMS Abstraction & Reporting Tool (CART). Vendor

tools also available. Retooled eMeasure

Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-4; N-17

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

• Very important and high impact; however, room for improvement when 98.5% of performance rates are documented is extremely small.

0142 Aspirin prescribed at discharge for AMI

Suggest an all-or-none composite for the AMI discharge medication measures.

The Steering Committee asked about a special category for good, important measures that seem to be "topped out". In May 2011, the NQF Board approved a policy for a special category "reserve measures."

Committee re-voted on Importance except for 1b, opportunity for improvement:

Y-21; N-0

2. Scientific Acceptability of Measure Properties: C-14; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

3. Usability: C-11; P-3; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

 Unless there is another way to get at the question of disparities identified by the TAP analyses, reserve status appears to be the most cost effective option for this measure.

4. Feasibility: C-12; P-3; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Widely accepted and in use by CMS

Does the Measure Meet Criteria for Endorsement: Y-15; N-0; A-0 Rationale: Meets all criteria except for opportunity for improvement

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: MAINTAIN ENDORSEMENT AND PLACEMENT IN RESERVE STATUS

Additional recommendation: The Steering Committee also recommends the measure be recalculated again in 3-5 years to monitor performance.

484

485 **Measures not recommended:**

960 Composite measure of hospital quality for acute myocardial infarction (AMI)

For More Information: Complete Measure Submission: Meeting/Call Proceedings

Description: A composite measure of in-hospital process and outcome of care for acute myocardial infarction (AMI) patients. Components of the Composite: Hospital process-of-care indicators

- 1. Percent of AMI patients given aspirin on arrival (NQF #0132; Endorsed May 9, 2007)
- 2. Percent of AMI patients given aspirin at discharge (NQF #0142; Endorsed May 9, 2007)
- 3. Percent of AMI patients given ACE inhibitor or ARB for LVSD (NQF #0137; Endorsed May 9, 2007)
- 4. Percent of AMI patients given smoking cessation advice/counseling (NQF #0027; Endorsed May 1, 2006)
- 5. Percent of AMI patients given beta blocker at discharge (NQF #0160; Endorsed May 9, 2007)
- Percent of AMI patients given fibrinolytic medication within 30 min. of arrival (NQF #0164; Endorsed May 9, 2007)
- 7. Percent of AMI patients given PCI within 90 min. of arrival (NQF #0163; Endorsed May 9, 2007)

Hospital outcome-of-care indicators

- AMI 30-day risk-standardized mortality (NQF #0230; Endorsed May 9, 2007)
- 2. AMI 30-day risk-standardized readmission (NQF #0505; Endorsed Oct. 28, 2008)

Numerator Statement: The sum of all successes for acute myocardial infarction process-of-care indicators, weighted by one-half the reciprocal of the share of opportunities represented by acute myocardial infarction process-of-care indicators in total opportunities, plus the sum of all successes for acute myocardial infarction outcome-of-care indicators, weighted by one-half the reciprocal of the share of opportunities represented by acute myocardial infarction outcome-of-care indicators in total opportunities.

Denominator Statement: The total number of opportunities for success on all acute myocardial infarction indicators used in the composite.

Exclusions: Hospitals missing three or more acute myocardial infarction process-of-care indicators and one or more outcome-of-care indicator were excluded.

Adjustment/Stratification: None

Level of Analysis: Facility Type of Measure: Composite

Data Source: Paper medical record/flowsheet; Electronic Health/Medical Record CMS Abstraction & Reporting Tool (CART). Vendor

tools also available. Measure Steward: CMS

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Yes-21; No-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

Composite measure of NQF endorsed measures for AMI.

2. Scientific Acceptability of Measure Properties: C-0; P-9; M-7; N-5

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- A lot of imputation of values due to missing data.
- Narrow range of results: 25th percentile = 83.1%; 75th percentile = 84.9%.
- Includes smoking cessation measure that has been determined to be invalid.

3. Usability: C-1; P-9; M-8; N-3

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Narrow range of results limts usefulness.
- Providers will find it hard to understand.

4. Feasibility: C-7; P-10; M-1; N-2

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Uses existing data from component measures.

Does the Measure Meet Criteria for Endorsement: Y-7; N-14; A-0 Rationale:

- Includes invalid smoking measure no longer endorsed by NQF.
- Limited variation in results.
- Question handling of large amount of missing data by imputation of national means.
- Complicated composite methodology—harder to understand compared to an "all or none."

If Applicable, Conditions/Questions for Developer:

RECOMMENDATION: DO NOT RECOMMEND

486

0282 Angina without procedure (PQI 13)

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: All non-maternal discharges of age 18 years and older with ICD-9-CM principal diagnosis code for angina.

Numerator Statement: All discharges of age 18 years and older with ICD-9-CM principal diagnosis code for angina.

Denominator Statement: Population in Metro area or county, age 18 years and older.

Exclusions: None

Adjustment/Stratification: Risk adjustment method widely or commercially available. The predicted value for each case is computed using standard logistic regression and covariates for gender and age (in 5-year age groups). The reference population used in the regression is the universe of discharges for states that participate in the HCUP State Inpatient Databases (SID) for the year 2007, a database consisting of approximately 35 million discharges from 43 states. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., county or state). The risk-adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate. Observed rates may be stratified by age and sex.

Data Source: Electronic administrative data/claims; Hospital administrative discharge data. See data requirements in the AHRQ QI

Windows Application Documentation: http://www.qualityindicators.ahrq.gov/software.htm

Measure Steward: AHRQ

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-0; N-21

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Coding of angina has demonstrated high variability and therefore reliability concerns. Changes in coding practices lead to significant changes in results.
- Should all admissions get a procedure? Seems to encourage procedures—wrong incentive.
- Developer states: "This indicator has unclear construct validity, because it has not been validated except as part of a set of indicators."
- There is wide variation in hospitalization rates by zip code.
- This is a community/population/geographic measure, not a hospital-level measure.

Does the Measure Meet Criteria for Endorsement: No --Did not pass Importance to Measure and Report. Rationale:

If Applicable, Conditions/Questions for Developer:

RECOMMENDATION: REMOVE ENDORSEMENT

487

1495 P2Y12 Inhibitor at discharge for patients with percutaneous coronary intervention (PCI) (with stents)

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of adult patients (age 18 or older) who undergo a percutaneous coronary intervention (PCI) (without a documented contraindication) with a stent implanted that had a P2Y12 inhibitor prescribed at discharge.

Numerator Statement: Count of patients with a PCI procedure with a P2Y12 inhibitor (Clopidogrel, Prasugrel, or Ticlopidine) prescribed at discharge.

Denominator Statement: Count of patients with a PCI procedure with a stent implanted.

Exclusions:

1495 P2Y12 Inhibitor at discharge for patients with percutaneous coronary intervention (PCI) (with stents)

-P2Y12 coded as contraindicated or blinded.

-Discharge status of expired.

-Discharge location of "other acute care hospital," "hospice," or "against medical advice."

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency Type of Measure: Process

Data Source: Registry data National Cardiovascular Data Registry (NCDR®) CathPCI Registry®

Measure Steward: ACC

STEERING COMMITTEE EVALAUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- This is based off a guideline that is the most widely recognized professional guideline in the United States for cardiovascular medicine in the area of PCI care.
- The value of the measure is high, but the performance gap is small and may represent reporting issues rather than true performance given the small gap of 7%.
- When the performance gap gets low, why not eliminate most exclusions? A key factor in terms of exclusions is they are the same as CMS inpatient measures as a means to reduce provider burden.

2. Scientific Acceptability of Measure Properties: C-19; P-2; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

3. Usability: C-17; P-4; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Harmonized to the extent possible with existing CMS measure and are specified identically.
- Is being used everywhere the NCDR is.
- Harmonization suggested with measure 558 and combined with 1493.

4. Feasibility: <u>C-17; P-4; M-0; N-0</u>

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Getting the outcome of transfers should not be too difficult.

Does the Measure Meet Criteria for Endorsement: Y-21: N-0: A-0

Rationale: Steering Committee would like to see this measure as a composite score with measure 1493 and 1498.

If Applicable, Conditions/Questions for Developer:

Have you considered an all or none composite for the PCI medication measures (1495, 1493, 1498)?

Response: Developer submitted a new composite measure 0964

Composite measure versus composite measure plus individual component measures:

The Committee vote to recommend only the composite and not the individual measures: Y- 11, N-8

RECOMMENDATION: DO NOT RECOMMEND AS AN INDIVIDUAL MEASURE

488

1493 Aspirin at discharge for patients with percutaneous coronary intervention (PCI)

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of adult patients (age 18 or older) who undergo a percutaneous coronary intervention (PCI) and are prescribed aspirin at discharge.

Numerator Statement: Count of patients with a PCI procedure with aspirin prescribed at discharge.

Denominator Statement: Count of patients with a PCI procedure.

1493 Aspirin at discharge for patients with percutaneous coronary intervention (PCI)

Exclusions:

-Aspirin coded as contraindicated or blinded.

-Discharge status of deceased.

-Discharge location of "other acute care hospital," "hospice," or "against medical advice."

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Facility/Agency Type of Measure: Process

Data Source: Registry data National Cardiovascular Data Registry (NCDR®) CathPCI Registry®

Measure Steward: ACC

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale: The Steering Committee agreed to duplicate voting on this measure to be the same as measure 1495. Unanimous

agreement to recommend that developer to combine 1495 and 1493.

If Applicable, Conditions/Questions for Developer:

Have you considered an all or none composite for the PCI medication measures (1495, 1493, 1498)?

Response: Developer submitted a new composite measure 0964

Composite measure versus composite measure plus individual component measures:

The Committee vote to recommend only the composite and not the individual measures: Y- 11, N-8

RECOMMENDATION: DO NOT RECOMMEND AS AN INDIVIDUAL MEASURE

489

1498 Statins at discharge for patients with percutaneous coronary intervention (PCI)

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of adult patients (age 18 or older) who undergo a percutaneous coronary intervention (PCI) and are prescribed a statin at discharge.

Numerator Statement: Count of patients with a PCI procedure with statin prescribed at discharge.

Denominator Statement: Count of patients with a PCI procedure.

Exclusions:

-Discharge status of deceased.

-Discharge location of "other acute care hospital," "hospice," or "against medical advice."

-Statins coded as contraindicated or blinded.

Adjustment/Stratification: N/A

Level of Analysis: Facility/Agency Type of Measure: Process

Data Source: Registry data National Cardiovascular Data Registry (NCDR®) CathPCI Registry®

Measure Steward: ACC

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-21; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Measure will encourage improvement in the rates of statin prescribing, which reduces the risk of coronary events and coronary artery disease following PCI.
- There is a performance gap. Prescribing rate from the 5th to the 98th percentile was from 72% to 98%.
- Stratified analysis indicated the lower SES hospitals did as well as or better than others.

2. Scientific Acceptability of Measure Properties: C-18; P-3; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Content validity tested by review by an expert consensus panel.
- Measure describes appropriate exclusions as well as option for contraindications.
- Consistent results reported for derivation cohort and testing cohort.

3. Usability: C-20; P-1; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing

1498 Statins at discharge for patients with percutaneous coronary intervention (PCI)

measures)

Rationale:

• This voluntarily reported measure is currently in use. Participating institutions receive an outcomes report each quarter with their individual results.

4. Feasibility: C-20; P-0; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Electronic sources are used.
- Reasonable information was provided about their efforts to reduce inaccuracies and follow-up on the process.

Does the Measure Meet Criteria for Endorsement: Y-21; N-0; A-0

Rationale:

If Applicable, Conditions/Questions for Developer:

Have you considered an all or none composite for the PCI medication measures (1495, 1493, 1498)?

Response: Developer submitted a new composite measure 0964

Composite measure versus composite measure plus individual component measures:

The Committee vote to recommend only the composite and not the individual measures: Y-11, N-8

RECOMMENDATION: DO NOT RECOMMEND AS AN INDIVIDUAL MEASURE

490 491

CARDIAC REHABILITATION

Measures not recommended:

492 493

1496 Cardiac rehabilitation/secondary prevention (CR) program structure-based measurement set to set safety standards for CR programming

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Cardiac rehabilitation/secondary prevention (CR) program measurement set to assess the presence of four safety standards.

Numerator Statement: The cardiac rehabilitation/secondary prevention (CR) program has policies in place that demonstrate all of the below:

- 1. A physician-director is responsible for the oversight of CR program policies and procedures and ensures that policies and procedures are consistent with evidence-based guidelines, safety standards, and regulatory standards. This includes appropriate policies and procedures for the provision of alternative CR program services, such as home-based CR.
- 2. An emergency response team is immediately available to respond to medical emergencies. (See numerator details for care setting details).
- 3. All professional staff have successfully completed the national Cognitive and Skills examination in accordance with the AHA curriculum for BLS with at least one staff member present who has completed the National Cognitive and Skills examination in accordance with the AHA curriculum for ACLS and has met state and hospital or facility medical-legal requirements for defibrillation and other related practices.
- 4. Functional emergency resuscitation equipment and supplies for handling cardiovascular emergencies are immediately available in the exercise area.

Denominator Statement: All CR programs.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Group; Facility/Agency; Integrated delivery system; Other Interdisciplinary teams of cardiac rehabilitation/secondary prevention professionals providing CR services.

Type of Measure: Structure/management

Data Source: Paper medical record/flowsheet; Organizational policies and procedures; Program policies and procedures and documentation of compliance using departmental records. This can be submitted electronically.

Measure Steward: American Association of Cardiovascular and Pulmonary Rehabilitation/American College of Cardiology

1496 Cardiac rehabilitation/secondary prevention (CR) program structure-based measurement set to set safety standards for CR programming

Foundation/American Heart Association (AACVPR/ACCF/AHA)

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-1

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Cardiac rehabilitation is an important and effective care process.
- Steering Committee questioned the evidence for the criteria.
- Only looks at 40% of programs that are certified.

2. Scientific Acceptability of Measure Properties: C-3; P-11; M-3; N-4

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- The program initially had to deny two-thirds of applications for remediation efforts, whereas more recently, all but two met criteria for safety.
- Measure is dependent on AACVPR certification, but can a program be just as compliant without being certified?
- Stewards state they are not aware of alternative data sources and note controversy regarding the applicability of the requirement for resuscitation equipment and supplies be available in the testing area when the testing area is in the home or other alternative settings.

3. Usability: <u>C-2; P-12; M-4; N-3</u>

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Currently in use for those programs that are currently certified.
- No data available for programs using the measure but are not certified. About 60% of the programs are not certified.
- NQF criteria does not require widespread national testing.

4. Feasibility: C-2; P-7; M-8; N-3

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

Feasible if certified; not that feasible if not certified.

Does the Measure Meet Criteria for Endorsement: Y-6; N-15; A-0

Rationale:

- Linkage to being certified in order to meet the measure.
- Absence of non-certification data.
- Structural measure:
- Unclear relationship to outcomes

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Not recommended for endorsement

494

1494 Cardiac rehabilitation/secondary prevention (CR) program measurement set related to monitoring response to therapy and documenting program effectiveness

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Cardiac rehabilitation/secondary prevention (CR) program measurement set to assess the presence of a written policy in place that demonstrates program effectiveness.

Numerator Statement: The cardiac rehabilitation/secondary prevention (CR) program is monitoring a response to therapy, and the program effectiveness has a written policy in place to capture all four of the elements below:

1. Document the percentage of patients for whom the CR program has received a formal referral request who actually enroll in

1494 Cardiac rehabilitation/secondary prevention (CR) program measurement set related to monitoring response to therapy and documenting program effectiveness

the program.

- 2. Document for each patient a standardized plan to assess completion of the prescribed course of CR as defined on entrance to the program.
- 3. Document for each patient a standardized plan to assess outcome measurements at the initiation and again at the completion of CR, including at least one outcome measure for the core program components as outlined in the Proposed AACVPR/ACCF/AHA Performance Measure: Individualized Assessment and Evaluation of Modifiable Cardiovascular Risk Factors, Development of Individualized Interventions, and Communication With Other Health Care Providers.
- 4. Describe the program's methodology to document program effectiveness and initiate quality improvement strategies.

Denominator Statement: All CR programs.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Group; Facility/Agency; Integrated delivery system; Program: Other

Type of Measure: Structure/management

Data Source: Paper medical record/flow-sheet; Organizational policies and procedures; Program policies and procedures and documentation of compliance using departmental records.

In addition, a National Outcomes Data Registry is being established by AACVPR to use in future to collect and analyze this data.

Measure Steward: AASVPR/ACCF/AHA

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-1

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Similar construct and comments as measure 1496.
- 55% patients are referred, but only 19% actually enroll.
- Not known if there is a gap in performance because no data are available beyond the remediation efforts of the overall certification.
- Structural measure

2. Scientific Acceptability of Measure Properties: C-3; P-15; M-3; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Disparities information included: less prescribed for elderly, women, and minorities.
- Results of reliability testing show good agreement (kappa inter-rater reliability testing), and Delphi-like peer review was used for validity testing.
- Four components in the numerator; three patient level and one system level.
- Impact of CR is four times the impact of timely PCI.
- No exclusions and no known disparities.

3. Usability: C-7; P-8; M-6; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Currently in use and publicly reported on several websites.
- Harmonized with other cardiac rehabilitation measures being reviewed.
- Stimulates quality improvement strategies for cardiac rehabilitation professionals, if they are certified.

4. Feasibility: C-1; P-12; M-4; N-4

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

• If the patient fails to complete the program it may affect the program's ability to capture the individual outcomes and accurately

1494 Cardiac rehabilitation/secondary prevention (CR) program measurement set related to monitoring response to therapy and documenting program effectiveness

reflect the program effectiveness.

Feasible and relatively low cost, although dependent on the AASCPR.

Does the Measure Meet Criteria for Endorsement: Y-3; N-17; A-0

Rationale:

- Similar to 1496. Standard is measured through certification; however, 60% cardiac programs do not participate in the certification program.
- Structural measure
- Unclear relationship to outcomes

If Applicable, Conditions/Questions for Developer:

RECOMMENDATION: Do not recommend for endorsement

495

1497 Cardiac rehabilitation/secondary prevention (CR) program measurement set to assess risk for adverse cardiovascular events

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Cardiac rehabilitation/secondary prevention (CR) program measurement set to assess the presence of two assessments of risk for adverse cardiovascular events.

Numerator Statement: The cardiac rehabilitation/secondary prevention (CR) program performs assessments of risk for adverse cardiovascular events:

- 1. Documentation, at program entry, that each patient undergoes an assessment of clinical status (e.g., symptoms, medical history) in order to identify high-risk conditions for adverse cardiovascular events.
- 2. A policy to provide recurrent assessments for each patient during the time of participation in the CR program in order to identify any changes in clinical status that increase the patient's risk of adverse cardiovascular events.

Denominator Statement: All CR Programs.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Group; Facility/Agency; Integrated delivery system

Type of Measure: Structure/management

Data Source: Organizational policies and procedures program policies and procedures and documentation of compliance using

departmental records. This can be submitted electronically.

Measure Steward: AAVCPR/ACCF/AHA

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-2

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Much of the discussion from the previous two measures, 1496 and 1494, applies here.
- The measure submitters use program certification data to indicate a gap. Information submitted is unclear whether failure to obtain certification is directly related to the lack of the policies and behaviors included in the measure or for other reasons.

2. Scientific Acceptability of Measure Properties: C-1; P-13; M-6; N-1

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Much of the discussion from the previous two measures, 1496 and 1494, applies here.
- Stewards state that there is no standardized risk assessment method in use. This is a concern for a performance measure.
- The measure did not meet criteria for endorsement because there is no "one best or standard" method of screening.
- Reliability testing minimally addressed this specific measure.
- Evidence for scoring seems to be on the composite of all CR measures taken together, but not individually.

3. Usability: <u>C-2; P-10; M-7; N-1</u>

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing

1497 Cardiac rehabilitation/secondary prevention (CR) program measurement set to assess risk for adverse cardiovascular events

measures)

Rationale:

Much of the discussion from the previous two measures, 1496 and 1494, applies here.

4. Feasibility: C-0; P-11; M-8; N-1

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- Much of the discussion from the previous two measures, 1496 and 1494, applies here.
- Electronic sources were not addressed.
- Review is audit of policies, not an audit of actual use in patients.

Does the Measure Meet Criteria for Endorsement: Y-2; N19; A-0

Rationale:

- The Steering Committee encouraged the measure developers to rework this measure in to one that would be much more
 usable.
- The Steering Committee believed it was important to note that its vote against the measures should <u>not</u> be interpreted as a rejection of the importance of, and the need for, a standard in America for cardiac rehabilitation programs.

If Applicable, Conditions/Questions for Developer:

RECOMMENDATION: Not recommended for endorsement

496

497

960 Cardiac rehabiltation composite

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: This measure evaluates whether a cardiac rehabilitation/secondary prevention program has processes in place for individualized assessment and evaluation of modifiable cardiovascular risk factors, development of individualized interventions, and communication with other health care providers.

Numerator Statement: The cardiac rehabilitation/secondary prevention (CR) program has all 11 processes in place for an individualized assessment and evaluation of modifiable cardiovascular risk factors, development of individualized interventions, and communication with other healthcare providers.

Denominator Statement: All CR Programs.

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinicians: Group; Facility/Agency; Integrated delivery system

Type of Measure: Structure/management

Data Source: Organizational policies and procedures program policies and procedures and documentation of compliance using

departmental records. This can be submitted electronically.

Measure Steward: AACVPR/ACCF/AHA

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-2

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- The discussion from the previous three measures applies here.
- The measure submitters use program certification data to indicate a gap. Information submitted is unclear whether failure to obtain certification is directly related to the lack of the policies and behaviors included in the measure or for other reasons.

2. Scientific Acceptability of Measure Properties: C-1; P-13; M-6; N-1

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2a. Comparability; 2h. Disparities)

960 Cardiac rehabiltation composite

Rationale:

- Much of the discussion from the previous two measures, 1496 and 1494, applies here.
- Stewards state that there is no standardized risk assessment method in use. This is a concern for a performance measure.
- The measure did not meet criteria for endorsement because there is no "one best or standard" method of screening.
- Evidence for scoring seems to be on the composite of all CR measures taken together, but not individually.

3. Usability: C-2; P-10; M-7; N-1

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

• See discussion of component measures.

4. Feasibility: C-0; P-11; M-8; N-1

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions—no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified; 4e. Data collection strategy can be implemented)

Rationale:

- The discussion from the previous three measures applies here.
- Electronic sources were not addressed.
- Review is audit of policies, not an audit of actual use in patients.

Does the Measure Meet Criteria for Endorsement?: Y-2; N19; A-0

Rationale:

- The Steering Committee encouraged the measure developers to rework this measure into one that would be much more
 usable.
- The Steering Committee believed it was important to note that its vote against the measures should <u>not</u> be interpreted as a rejection of the importance of, and the need for, a standard in America for cardiac rehabilitation programs.
- Specific issues:
 - The absence of noncertified validity and reliability data.
 - The linkage of these measures to certification.
 - The absence of outcomes or favorable outcomes related to certification.
 - The need for patient-level measures.

If Applicable, Conditions/Questions for Developer:

RECOMMENDATION: Do not recommend for endorsement

498

499

500

ATRIAL FIBRILLATION

Recommended for endorsement:

1524 Assessment of thromboembolic risk factors (CHADS 2)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

1524 Assessment of thromboembolic risk factors (CHADS 2)

Description: : Patients with nonvalvular atrial fibrillation or atrial flutter in whom assessment of thromboembolic risk factors using the CHADS2 risk criteria has been documented

Numerator Statement: Patients with nonvalvular atrial fibrillation or atrial flutter in whom assessment of all of the specified thromeboembolic risk factors is documented

For patients with nonvalvular atrial fibrillation or atrial flutter, assessment of thromboembolic risk should include the following factors: Electronic Specifications:

- Risk factors:
- Prior stroke or transient ischemic attack--> High risk
- Age = 75 years--> Moderate risk
- Hypertension--> Moderate risk
- Diabetes mellitus--> Moderate risk
- Heart failure or impaired LV systolic function--> Moderate risk

Denominator Statement: All patients 18 years of age or older with nonvalvular atrial fibrillation or atrial flutter other than those specifically excluded

Exclusions:

- Patients with mitral stenosis or prosthetic heart valves
- Patients with transient or reversible causes of atrial fibrillation (e.g., pneumonia or hyperthyroidism)
- Postoperative patients
- Patients who are pregnant
- Medical reason(s) documented by a physician, nurse practitioner, or physician assistant for not assessing risk factors.
 Examples of medical reasons for not assessing risk factors include but are not limited to the following:
 - Allergy to warfarin
 - Risk of bleeding

Adjustment/Stratification: No risk adjustment necessary None

Level of Analysis: Clinician: Individual Type of Measure: Process

Data Source: Electronic Clinical Data, Paper medical record/flow-sheet, Registry data

Measure Steward: American College of Cardiology (ACC) Foundation/American Heart Association (AHA)/American Medical

Association's Physician Consortium for Performance Improvement, 2400 N. Street NW, Washington, DC, 20037

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-18; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Hospital admissions for atrial fibrillation have increased 66% in the past decade.
- Approximately 60,000 strokes each year are preventable with appropriate risk stratification and anticoagulation with warfarin.
- Strong evidence base.
- Vague title. Steering Committee recommended changing the title to be more specific.

2. Scientific Acceptability of Measure Properties: C-12; P-6; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Uses CHAD2 score, which are in AHA/ACC Guidelines.
- Rigorously tested. Reliable and valid.
- Requires good documentation; may underestimate. More documentation needed if warfarin is not recommended.
- Testing of measure used Pinnacle registry data.

3. Usability: C-13; P-7; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

- Promotes better physician documentation.
- Requires good documentation or results will underestimate performance.

1524 Assessment of thromboembolic risk factors (CHADS 2)

4. Feasibility: C-7; P-12; M-0; N-1

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

 All of the data elements are available through a paper source, electronic health record (EHR) or electronic medical record (EMR). No exclusions..

Does the Measure Meet Criteria for Endorsement?:: Y-17; N-3; A-0 Rationale:

- Documentation that calculating a CHADS2 score improves the match of anticoagulation with the risk factors for stroke.
 However, the target of the measure, prevention of stroke due to atrial fibrillation is important, and the measure appears feasible.
- The most frequent reason for low scoring is failure of the physician to document the CHADS2 score.

If applicable, Conditions/Questions for Developer: Specifically mention the CHADS2 criteria in the measure specification. Title is vaque.

Developer Response: The developer revised the specifications to include the CHADS2. The developer changed the title to "Assessment of Thromboembolic Risk Factors (CHADS2)".

RECOMMENDATION: Recommend for endorsement

501

1525 Chronic anticoagulation therapy

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Prescription of warfarin or another anticoagulant drug that is FDA approved for the prevention of thromboembolism for all patients with nonvalvular atrial fibrillation or atrial flutter at high risk for thromboembolism, according to CHADS2 risk stratification.

Numerator Statement: All patients with nonvalvular atrial fibrillation or atrial flutter at high risk of thromboembolism (i.e., those with any high-risk factor or more than 1 moderate-risk factor) who are prescribed warfarin OR another anticoagulant drug that is FDA approved for the prevention of thromboembolism.

Denominator Statement: Patients with nonvalvular AF or atrial flutter for whom assessment of the specified thromboembolic risk factors documented one or more high-risk factor or more than one moderate-risk factor

Exclusions:

- Patients with valvular AF, specifically those with prosthetic heart valves or mitral stenosis
- Patients at low risk for thromboembolism (i.e., those with none of the risk factors listed above)
- Patients with only one moderate risk factor
- Postoperative patients
- Patients with transient or reversible causes of AF (e.g., pneumonia or hyperthyroidism)
- Patients who are pregnant
- Medical reason(s) documented by a physician, nurse practitioner, or physician assistant for not prescribing warfarin. Examples of medical reasons for not prescribing warfarin include, but are not limited to:
 - Allergy
 - Risk of bleeding
- Documentation of patient reason(s) for not prescribing warfarin or another anticoagulant drug that is FDA approved for the prevention of thromboembolism (e.g., economic, social, and/or religious impediments, noncompliance or patient refusal)

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinician: Individual Type of Measure: Process

Data Source: Electronic Clinical Data, Paper medical record/flow-sheet, Registry data

Measure Steward: American College of Cardiology Foundation/ American Heart Association/American Medical Association's Physician Consortium for Performance Improvement, 2400 N. Street NW, Washington, DC 20037

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

1525 Chronic anticoagulation therapy

Rationale:

- Importance demonstrated by decrease in stroke by 66% for patients with atrial fibrillation treated with warfarin.
- 45-55% of candidates for anticoagulation do not receive risk assessment or treatment.
- Race and gender data disparities are evident.
- Class I Level A evidence. CHADS2 score has been validated.
- 2. Scientific Acceptability of Measure Properties: C-1; P-4; M-10; N-5 (As submitted)

If conditions are met: C-3; P-13; M-3; N-1

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Steering Committee discussed including the newer anticoagulants or other FDA-approved drugs besides warfarin.
- Measure doesn't specify CHADS2. Should be consistent with measure 1524.
- Second vote with conditions set by Steering Committee (as submitted in addition to the following): 1) Include CHADS2 in specifications. 2) Numerator to include "other FDA-approved drugs".; and 3) Exclusions include patient or physician preference reason for alternative treatment.

3. Usability: C-13; P-7; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

Not used in public reporting yet but will be eligible for use in PQRS in 2012.

4. Feasibility: C-14; P-5; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- Recognizes the need for better documentation to achieve more accurate assessment of physician performance.
- Data are generated through the usual care processes. Electronic sources are available.

Does the Measure Meet Criteria for Endorsement?: (as submitted) Y-7; N-12

With Conditions: 1) Include all Food and Drug Administration (FDA)-approved drugs for this condition, i.e., dabigatran; and 2) Specify CHADS2 risk assessment: Y-16; N-3

Rationale:

- Important process of care—high morbidity.
- Developer complied with conditions.
- Evidence-based action based on standardized risk assessment.

If applicable, Conditions/Questions for Developer:

- What about newer anticoagulants besides warfarin?
- Why not use CHADS2 scoring for consistency?

Developer Response:

- Developer revised the measure to include "all FDA approved drugs for this condition."
- Developer revised the measure to specify CHADS2 scoring.

RECOMMENDATION: Recommend for endorsement

502

Not recommended:

1505 Adult patient(s) with atrial fibrillation taking amiodarone that had serum ALT or AST test in last 12 reported months

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: This measure identifies adults with atrial fibrillation, 18 years of age or older, taking amiodarone that had at least one serum ALT or AST test in last 12 months of the report period.

1505 Adult patient(s) with atrial fibrillation taking amiodarone that had serum ALT or AST test in last 12 reported months

Numerator Statement: Patients who are diagnosed with atrial fibrillation and who are treated with amiodarone, who have had a serum AST/ALT test during the following time period: last 12 months of the report period through 90 days after the end of the report period Denominator Statement: All patients 18 years of age or older who have a diagnosis of atrial fibrillation and who are actively being treated with amiodarone.

Exclusions: Criteria for inclusion in the denominator are as follows:

- 1. All male and female patients who are 18 years or older at the end of the report period
- 2. Patient must have been continuously enrolled in medical benefits throughout the 12 months prior to the end of the report period AND pharmacy benefit plan for 6 months prior to the end of the report period. The standard EBM Connect® enrollment break logic allows unlimited breaks in coverage of no more than 45 days and no breaks greater than 45 days.
- The patient is listed in the Disease Registry Input File for this condition OR

Patient fulfills both criteria A and B:

- A. During the 24 months prior to the end of the report period, the patient has two or more of the following services or events, at least 14 days apart, with a diagnosis of atrial fibrillation (code set DX0014):
 - Professional Encounter (code set PR0107, RV0107)
 - Professional Supervision (code set PR0108)
 - Facility Event—Confinement/Admission (i.e., hospitalization)
 - Facility Event—Emergency Room
 - Facility Event—Outpatient Surgery

AND

- B. During the 12 months prior to the end of the report period, the patient has one or more of the following services or events, with a diagnosis of atrial fibrillation (code set DX0014):
 - Professional Encounter (code set PR0107, RV0107)
 - Professional Supervision (code set PR0108)
 - Facility Event—Confinement/Admission (i.e., hospitalization)
 - Facility Event—Emergency Room
 - Facility Event—Outpatient Surgery
- 4. The patient must have filled a prescription for amiodarone (code set RX-9) during the following time period: last 120 days of the report period through 90 days after the end of the report period AND the duration of treatment was greater than 90 days.

Code Set Code Set Description Diagnosis Code

DX0014 Atrial Fibrillation 427.3 DX0014 Atrial Fibrillation 427.31 DX0014 Atrial Fibrillation 427.32

Code Set Code Set Description Procedure Code

PR0107 Professional encounter 99201-99205, 99211-99223 (except 99216), 99231-99245 (except 99237, 99240), 99251-99255, 99261-99263, 99271-99275, 99281-99285, 99301-99313, 99315, 99316, 99318, 99341-99350 (except 99346), 99381-99387, 99391-99397, 99401-99404, 99411-412, 99420, 99429, S0270-S0273

Code Set Code Set Description Procedure Code

PR0108 Professional supervision 99321-99328, 99331-99337, 99339-99340, 99371-99380 (except 99376), 99441-99444, G0179-G0182

Code Set Code Set Description Revenue Code

RV0107 Professional encounter 0510-0517, 0519-0526, 0528-0529, 0981, 0983

Rx code set Rx code set description ndc Amiodarone

Adjustment/Stratification: Does not apply; No risk adjustment necessary

Level of Analysis: Type of Measure: Process

Data Source: A 15 million patient population sample was chosen to analyze the potential patient safety gap in care. The sample was derived from more than 60 million patients based on criteria including national geographic representation, commercial health coverage, and patient age less than 65.

Measure Steward: Ingenix

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-1; N-17

1505 Adult patient(s) with atrial fibrillation taking amiodarone that had serum ALT or AST test in last 12 reported months

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Too narrow; there are other toxicities for this drug. Why choose this one?
- Why not the multitude of tests for potential issues with many drugs?
- This drug warrants a composite of multiple side effects monitoring.
- Low numbers of incidence; measure overload.

Does the Measure Meet Criteria for Endorsement?: No

Rationale: This measure did not pass Importance to Measure and Report.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Do not recommend

504

505

506

IMPLANTABLE CARDIOVERTER DEFIBRILLATORS (ICD)

Recommended for endorsement:

1522 ACE/ARB therapy at discharge for ICD implant patients with LVSD

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of ICD implant patients with a diagnosis of LVSD who are prescribed ACE-I or ARB therapy at discharge

Numerator Statement: Count of patients with ACE-I or ARB therapy prescribed at discharge

Denominator Statement: Count of patients with an ICD implant with moderate or severe LVSD (LVEF<40%) without contraindication to

ACE inhibitors and ARBs

Exclusions:

- Patients who expired prior to discharge
- Patients with ACE-I and ARB therapy contraindicated or blinded

Adjustment/Stratification: N/A Level of Analysis: Facility/Agency Type of Measure: Process Data Source: Registry data

Measure Steward: American College of Cardiology Foundation, 2400 N Street NW, Washington, DC 20037

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Patient group of high morbidity and mortality.
- Still a performance gap, although narrowing with the implementation of current guality improvement programs.
- Strong outcome evidence in terms of efficacy.

2. Scientific Acceptability of Measure Properties: C-18; P-2; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Reliability and validity of the measure are strong.
- Indication for ICD is based on maximum medical therapy.

3. Usability: C-19; P-0; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

Adds value to exisiting measures.

1522 ACE/ARB therapy at discharge for ICD implant patients with LVSD

Useful for public reporting.

4. Feasibility: C-20; P-0; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

Easily obtained from the electronic source/registry.

Does the Measure Meet Criteria for Endorsement?: Y-19; N-0; A-0

Rationale:

- Recommend an all-or-none composite for medications.
- Recommend as a stand-alone as well as part of composite 0965.
- Recommend amending the wording to clarify inclusion and include a broader scope of patients (biventricular without ICD).

If applicable, Conditions/Questions for Developer: Is ICD being used here as a generic or a specific term?

Developer Response: This applies to patients receiving any rhythm management device.

Steering Committee Follow-up: Why not include biventricular device without ICD?

Developer Follow-up: Could clarify to include patients who get biventricular device without ICD.

RECOMMENDATION: Recommend for endorsement as an individual measure as well as a component of the

composite 965

507 508

1528 Beta blocker at discharge for ICD implant patients with a previous MI

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of ICD implant patients with a diagnosis of previous myocardial infarction (MI) who are prescribed a beta

blocker at discharge

Numerator Statement: Count of patients discharged on beta-blocker therapy

Denominator Statement: Count of patients with an ICD implant without contraindication to beta-blockers

Exclusions:

-Patients who expired

-Beta-blocker therapy contraindicated or blinded.

Contraindicated supporting definition:

Medication was not prescribed because of a contraindication.

Contraindications must be documented explicitly by the physician, or clearly evidenced within the medical record

Blinded supporting definition:

Patient was in research study or clinical trial and administration of this specific medication is unknown

Adjustment/Stratification: N/A

Level of Analysis: Facility/Agency Type of Measure: Process

Data Source: Registry data

Measure Steward: American College of Cardiology Foundation (ACCF), 2400 N Street NW, Washington, DC 20037

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- High impact and large population at risk.
- There is a relatively small but significant "performance" gap with median performance of around 87-90%, quartile 1 at 83%, and quartile 3 at 96%.

2. Scientific Acceptability of Measure Properties: C-19; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

1528 Beta blocker at discharge for ICD implant patients with a previous MI

- Aligned with specifications from other CMS measures for ICD.
- Well-defined measure with well-accepted, well-documented exclusions.
- Good face validity and supported by evidence-based guidelines.
- Data analysis shows that this measure discerns differences in performance—mostly from ICD registry of 144,000 patient records in 1,305 hospitals from 2008-2009.
- No disparities have been reported.

3. Usability: C-20; P-0; M-1; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

Measure is meaningful, understandable, and easy to use in different formats.

4. Feasibility: C-19; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

 NCDR electronic database is well tested and takes many steps to minimize inaacuracies, including thorough training of data abstractors, certification process of hospital EMR or NCDR's web-based tool, frequent edit checks, frequent validity checks, and an onsite audit program.

Does the Measure Meet Criteria for Endorsement?: <u>Y-20; N-0; A-0</u> Rationale:

• Recommend as a stand-alone as well as part of composite 965.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Recommend for endorsement as an individual measure as well as a component of the composite 965

509

1529 Beta blocker at discharge for ICD implant patients with LVSD

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of ICD implant patients with a diagnosis of LVSD who are prescribed beta blocker therapy on discharge

Numerator Statement: Count of patients with beta blocker therapy prescribed on discharge

Denominator Statement: Count of patients with an ICD implant with LVSD without contraindication to beta blockers

Exclusions: Procedure type=initial generator implant=yes or generator change=yes

Most recent LVEF<40%

Adjustment/Stratification: N/A Discharge status=deceased

Beta blocker (any)=contraindicated or blinded

Contraindicated supporting definition:

Medication was not prescribed because of a contraindication.

Contraindications must be documented explicitly by the physician, or clearly evidenced within the medical record

Blinded supporting definition:

Patient was in research study or clinical trial and administration of this specific medication is unknown

Level of Analysis: Affects large numbers; Frequently performed procedure; Leading cause of morbidity/mortality; High resource use;

Severity of illness

Type of Measure: Process

Data Source: N/A

Measure Steward: American College of Cardiology Foundation, 2400 N Street NW, Washington, DC 20037

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

1529 Beta blocker at discharge for ICD implant patients with LVSD

High-risk population and impact gap.

2. Scientific Acceptability of Measure Properties: C-20; P-0; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

Tested for reliability and validity.

3. Usability: <u>C-18; P-0; M-0; N-0</u>

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Information produced is meaningful and easy to understand.
- Data are currently being used in registries.

4. Feasibility: C-19; P-0; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

Does the Measure Meet Criteria for Endorsement?: Y-20; N-0; A-0

Rationale:

- Recommended as a stand-alone as well as part of composite 0965.
- Patients not captured in beta blocker after AMI measure (1528) because ICD is the primary diagnosis.
- Evaluation the same as 1528.
- Recommended an all-inclusive measure for beta blockers.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Recommend for endorsement as an individual measure as well as a component of the composite 965

510

511

0965 Patients with an ICD implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of patients with an ICD implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge (all-or-none composite measure of two medication classes)

Numerator Statement: Patients who receive all medications for which they are eligible.

- 1. ACE/ARB prescribed at discharge (if eligible for ACE/ARB as described in denominator) AND
- 2. Beta blockers prescribed at discharge (if eligible for beta blockers as described in denominator)

Denominator Statement: All patients with an ICD implant surviving hospitalization who are eligible to receive any one of the two medication classes:

- 1. Eligiblility for ACE/ARB: Patients who have an ejection fraction (EF) of <40% AND do not have a documented contraindication to ACE/ARB documented OR
- 2. Eligibility for beta blockers: Patients who do not have a documented contraindication to beta blocker therapy and have either:
- a. EF of <40% OR
- b. A previous myocardial infarction (MI)

Exclusions: Discharge status of expired; not eligible for either ACE/ARB or beta blockers

Adjustment/Stratification: N/A

Level of Analysis: Hospital (inpatient and outpatient)

Type of Measure: Process

Data Source: N/A

0965 Patients with an ICD implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge

Measure Steward: American College of Cardiology Foundation, 2400 N Street NW, Washington, DC 20037

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- High-risk population and impact gap.
- Composite combines three medication measures.

2. Scientific Acceptability of Measure Properties: C-20; P-0; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaninaful differences; 2a. Comparability; 2h. Disparities)

Rationale:

Tested for reliability and validity.

3. Usability: C-18; P-0; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Information produced is meaningful and easy to understand.
- Data are currently being used in registries.

4. Feasibility: C-19; P-0; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale: Uses same data as the individual measures.

Does the Measure Meet Criteria for Endorsement?: Y-20; N-0; A-0

Rationale:

All or none composite.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Recommend for endorsement

512

513 Not recommended:

1530 Prophylactic antibiotics prior to ICD (lead or implant) procedure

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Proportion of patients that receive an ICD implant or lead procedure that receive antibiotics within 1 hour (if fluoroquinolone or vancomycin, 2 hours) prior to procedure

Numerator Statement: Count of patients that receive antibiotics prior to the ICD implant or leads procedure

Denominator Statement: Count of patients with an ICD implant or lead procedure

Exclusions: Count of patients with arrival/discharge dates from data submissions that pass NCDR data inclusion thresholds Adjustment/Stratification: N/A Prophylactic antibiotics within 1 hour of procedure start time=No—not given, medical reason documented, including:

- Patients with a documented contraindication to receiving prophylactic antibiotics prior to the ICD implant
- Patients receiving continuous antibiotics >24 hours prior to the implant

Level of Analysis: Affects large numbers; Frequently performed procedure; Leading cause of morbidity/mortality; High resource use; Severity of illness

Type of Measure: Process

Data Source: N/A

Measure Steward: American College of Cardiology Foundation (ACCF), 2400 N Street NW, Washington, DC, 20037

STEERING COMMITTEE EVALUATION

1530 Prophylactic antibiotics prior to ICD (lead or implant) procedure

1. Importance to Measure and Report: Y-3; N-17

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Should be incorporated into SCIP measure
- High current performance—median is 100%
- Little gap— criteria 1b not met.

Does the Measure Meet Criteria for Endorsement?: Not recommended

Rationale: Did not meet criteria for Importance to Measure and Report

- Unclear at this point if there is a performance gap.
- No data on reliability of measure or disparities.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Do not recommend

514

515

516

HEART FAILURE

Recommended for endorsement:

0079 Heart failure: Left ventricular ejection fraction assessment (outpatient setting)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of heart failure for whom the quantitative or qualitative results of a recent or prior (any time in the past) LVEF assessment is documented within a 12-month period

Numerator Statement: Patients for whom the quantitative or qualitative results of a recent or prior (any time in the past) LVEF assessment is documented* within a 12-month period

*Documentation must include documentation in a progress note of the results of an LVEF assessment, regardless of when the evaluation of ejection fraction was performed.

Qualitative results correspond to numeric equivalents as follows:

- Hyperdynamic: corresponds to LVEF greater than 70%
- Normal: corresponds to LVEF 50% to 70% (midpoint 60%)
- Mild dysfunction: corresponds to LVEF 40% to 49% (midpoint 45%)
- Moderate dysfunction: corresponds to LVEF 30% to 39% (midpoint 35%)
- Severe dysfunction: corresponds to LVEF less than 30%

Denominator Statement: All patients aged 18 years and older with a diagnosis of heart failure

Exclusions: None

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinician: Individual Type of Measure: Process

Data Source: Electronic administrative data/claims, Electronic Clinical Data, Paper medical record/flow-sheet, Registry data

Measure Steward: American Medical Association, 515 N State St., Chicago, IL 60654

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-1

(1a. Impact: 1b. Performance gap: 1c. Outcome or Evidence)

Rationale:

- Heart failure is a common, high-mortality condition that comprises two entities—systolic and diastolic heart failure. The ejection fraction needs to be known in order to differentiate the two conditions.
- Evidence is Level C, Class I recommendation.
- Important measure and is used to base other measures.
- Will this be interpreted as needing a new test every 12 months even though the specification requires that the test results, even if done in the past, be in the current documentation?
- 2. Scientific Acceptability of Measure Properties: C-12; P-6; M-1; N-0

0079 Heart failure: Left ventricular ejection fraction assessment (outpatient setting)

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Well-defined and has been shown to be reliable and valid.
- There are no exclusions.
- Risk adjustment is not necessary.
- Disparities have not been identified.

3. Usability: C-12; P-6; M-2; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- The measure is meaningful, understandable, and provides distinct value.
- Selection codes are harmonized with measure 0135.
- Some concern with promoting overuse of LVSD testing by misinterpreting the measure.

4. Feasibility: C-7; P-11; M-1; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- Data can be collected with paper or electronic medical record, claims, or registry data.
- Concern that the measure may drive overuse.

Does the Measure Meet Criteria for Endorsement?: Y-18; N-1; A-0

Rationale:

- Basis of other treatments.
- Well-defined; demonstrated to be reliable and valid.

If applicable, Conditions/Questions for Developer: The Steering Committee suggested changing title and description to more accurately reflect what is measured.

RECOMMENDATION: Recommend for endorsement

517

518

0081 Heart Failure: Angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) therapy for left ventricular systolic dysfunction

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40% who were prescribed ACE inhibitor or ARB therapy either within a 12-month period when seen in the outpatient setting or at hospital discharge

Numerator Statement: Patients who were prescribed* ACE inhibitor or ARB therapy either within a 12-month period when seen in the outpatient setting or at hospital discharge

*Prescribed may include prescription given to the patient for ACE inhibitor or ARB therapy at one or more visits in the measurement period OR patient already taking ACE inhibitor or ARB therapy as documented in current medication list.

Denominator Statement: All patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40%. LVEF < 40% corresponds to qualitative documentation of moderate dysfunction or severe dysfunction.

Exclusions:

- Documentation of medical reason(s) for not prescribing ACE inhibitor or ARB therapy; Append modifier to CPT II code 4009F-1P
- Documentation of patient reason(s) for not prescribing ACE inhibitor or ARB; Append modifier to CPT II code 4009F-2P
- Documentation of system reason(s) for not prescribing ACE inhibitor or ARB; Append modifier to CPT II code 4009F-3P

Adjustment/Stratification: No risk adjustment necessary

0081 Heart Failure: Angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) therapy for left ventricular systolic dysfunction

Level of Analysis: Clinician: Individual Type of Measure: Process

Data Source: Electronic administrative data/claims, Electronic Clinical Data, Paper medical record/flow-sheet, Registry data; Retooled eMeasure

Measure Steward: American Medical Association, 515 N State St., Chicago, IL 60654

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-18; N-1

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- The topic of measurement (ACE/ARB for HF with low EF) is of high impact, there are definite quality problems, and there is RCT evidence that prescribing ACE/ARB improves outcomes.
- Signifigant performance gap in the outpatient setting and strong outcome in evidence.

2. Scientific Acceptability of Measure Properties: C-19; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Very well specified.
- Reliability and validity are both extensively discussed in the PCPI review.
- Exlcusions justified and consistent with other ACE and ARB measures.

3. Usability: C-13; P-7; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- The information produced by the measure is meaningful and useful.
- It is harmonized with measure 0162.

4. Feasibility: <u>C-16; P-3; M-0; N-0</u>

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- The data elements for the measure are routinely generated from phamacy claims.
- The data tend to be accurate, and being in use already, feasibility has been documented.

Does the Measure Meet Criteria for Endorsement?: Y-19; N-0; A-0

Rationale:

- ACE/ARB for HF with low EF in the ambulatory setting offers important therapeutic benefits.
- Significant disparities and variations in care exist.
- The measure is already used successfully.

If applicable, Conditions/Questions for Developer: Please explain why you're requesting endorsement of this measure at an individual clinician level of measurement to avoid duplication (measure 0162).

Response: The intent is to really enhance care on the outpatient side, looking at individual clinicians on the outpatient performance.

RECOMMENDATION: Recommend for endorsement

519

0083 Heart failure: Beta-blocker therapy for left ventricular systolic dysfunction

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40% who were prescribed beta-blocker therapy either within a 12-month period when seen in the outpatient setting or at hospital discharge Numerator Statement: Patients who were prescribed* beta-blocker therapy** either within a 12-month period when seen in the outpatient setting or at hospital discharge

*Prescribed may include prescription given to the patient for beta-blocker therapy at one or more visits in the measurement period OR

0083 Heart failure: Beta-blocker therapy for left ventricular systolic dysfunction

patient already taking beta-blocker therapy as documented in current medication list.

**Beta-blocker therapy should include bisoprolol, carvedilol, or sustained release metoprolol succinate.

Denominator Statement: All patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40%.

LVEF < 40% corresponds to qualitative documentation of moderate dysfunction or severe dysfunction.

Exclusions:

- Documentation of medical reason(s) for not prescribing beta-blocker therapy
- Documentation of patient reason(s) for not prescribing beta-blocker therapy
- Documentation of system reason(s) for not prescribing beta-blocker therapy

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinician: Individual Type of Measure: Process

Data Source: Electronic administrative data/claims, Electronic Clinical Data, Paper medical record/flow-sheet, Registry data; Retooled eMeasure

Measure Steward: American Medical Association, 515 N State St., Chicago, IL 60654

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- High impact; heart failure is prevalent and associated with high mortality rates.
- Beta blockers have been shown to reduce mortality, but wide variability still exists.

2. Scientific Acceptability of Measure Properties: C-18; P-0; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- The measure is well-defined and precise.
- Certain beta-blocker drugs, based on the evidence, are specified.
- Reliability was tested on a previous measure that is related.
- The measure is valid and exclusions are identified.
- Disparities in care have not yet been identified.

3. Usability: C-18; P-2; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Information provided by the measure is meaningful.
- Information about harmonization is not provided.
- The measure is already being used successfully

4. Feasibility: C-19; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale: The data are routinely generated from pharmacy records. Exclusions do not require additional data sources. Reasonable accuracy has been demonstrated, and data collection is feasible.

Does the Measure Meet Criteria for Endorsement?: Y-17; N-0; A-0

Rationale:

- The prescription of beta blockers for heart failure has been shown to improve outcomes.
- Prescription rates do vary.
- The measure is already being used successfully.

If applicable, Conditions/Questions for Developer: Exclusions indicate there may be systemic or organizational reasons for excluding someone. What might the reasons be?

Response: We have to talk about patient reasons for exclusion as well as system reasons. System reasons could be high cost or other reasons related to resources. Patient would be excluded because of valid reasons if why they haven't received a beta blocker is

0083 Heart failure: Beta-blocker therapy for left ventricular systolic dysfunction

indicated somewhere in the record.

RECOMMENDATION: Recommend for endorsement

520

0162 ACEI or ARB for left ventricular systolic dysfunction—heart failure (HF) patients

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of heart failure (HF) patients with left ventricular systolic dysfunction (LVSD) who are prescribed an ACEI or ARB at hospital discharge. For purposes of this measure, LVSD is defined as chart documentation of a left ventricular ejection fraction (LVEF) less than 40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction.

Numerator Statement: HF patients who are prescribed an ACEI or ARB at hospital discharge

Denominator Statement: HF patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of HF: 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, 428.0, 428.1, 428.20, 428.21, 428.22, 428.23, 428.30, 428.31, 428.32, 428.33, 428.40, 428.41, 428.42, 428.43, 428.9); with chart documentation of a left ventricular ejection fraction (LVEF) < 40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction

Exclusions:

- Patients who had a left ventricular assistive device (LVAD) or heart transplant procedure during hospital stay (ICD-9-CM procedure code of LVAD or Heart Transplant: 33.6, 37.51, 37.52, 37.53, 37.54, 37.60, 37.62, 37.63, 37.65, 37.66, 37.68)
- <18 years of age
- Patients who have a length of stay greater than 120 days
- Discharged to another hospital
- Expired
- Left against medical advice
- Discharged to home for hospice care
- Discharged to a health care facility for hospice care
- Patients enrolled in clinical trials
- Patients with comfort measures only documented
- Patients with a documented reason for no ACEI and no ARB at discharge

Adjustment/Stratification: No risk adjustment necessary N/A

Level of Analysis: Facility/Agency, Population: National Type of Measure: Process

Data Source: Paper medical record/flow-sheet

Measure Steward: Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-18; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Multiple large multicenter clinical trials demonstrate importance of use of ACE/ARBs for patients with reduced LV function, with significant impact on long-term outcome.
- National Performance is 94%; lower in Native Americans.

2. Scientific Acceptability of Measure Properties: C-11; P-7; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

Patients with a missing LVSD value are excluded.

3. Usability: C-14; P-4; M-1; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

• Submission form included thorough discussion of harmonization.

0162 ACEI or ARB for left ventricular systolic dysfunction—heart failure (HF) patients

Currently in use/Hospital Compare.

4. Feasibility: C-13; P-5; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- Patients without LVEF documented are excluded—measure 0135 assesses measurement of LVSD and has high current performance.
- Data are easily obtainable.

Does the Measure Meet Criteria for Endorsement?: <u>Y-20; N-0; A-0</u> Rationale:

- Effective process of care that improves outcomes.
- Strong evidence base.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Recommend for endorsement

521

0358 Congestive heart failure (CHF) mortality rate (IQI 16)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Perecent of discharges with principal diagnosis code of CHF with in-hospital mortality

Numerator Statement: Number of deaths (DISP=20) among cases meeting the inclusion and exclusion rules for the denominator.

Denominator Statement: All discharges, age 18 years and older, with a principal diagnosis code of CHF.

Exclusions:

- missing discharge disposition (DISP=missing)
- transferring to another short-term hospital (DISP=2)
- MDC 14 (pregnancy, childbirth, and puerperium)

Adjustment/Stratification: Risk-adjustment method widely or commercially available. The predicted value for each case is computed using a hierarchical model (logistic regression with hospital random effect) and covariates for gender, age in years (in 5-year age groups), All Patient Refined-Diagnosis Related Group (APR-DRG), and APR-DRG risk-of-mortality subclass. The reference population used in the model is the universe of discharges for states that participate in the HCUP State Inpatient Databases (SID) for the year 2007 (updated annually), a database consisting of 43 states and approximately 30 million adult discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., hospital, state, and region). The risk-adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate.

Required data elements: Patient gender; age in years at admission; International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM) principal and secondary diagnosis codes. A limited license 3M APR-DRG grouper is included with the AHRQ QI Software. Gender, age (5-year age groups), race / ethnicity, primary payer, custom

Level of Analysis: Facility/Agency Type of Measure: Outcome

Data Source: Electronic administrative data/claims

Measure Steward: Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, MD 20850

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-12; N-7

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Heart failure is common and associated with high mortality rates.
- Committee recommended more recent evidence citations.

2. Scientific Acceptability of Measure Properties: C-1; P-14; M-3; N-1

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

Well-defined, valid and reliable.

0358 Congestive heart failure (CHF) mortality rate (IQI 16)

- Risk-adjustment algorithms are available and scoring and analysis allow for identification of disparities in outcome.
- No data element available that would allow exclusion for DNR.
- Detailed disparities information presented in measure submission.

3. Usability: C-8; P-7; M-3; N-1

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- The information provided by the measure is useful and meaningful.
- Many states already report the measure.
- If patient is admitted for palliative care, it is not captured as an acute admission.

4. Feasibility: C-15; P-5; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- The data are routinely generated.
- Exclusions do not require additional data.

Does the Measure Meet Criteria for Endorsement?: Y-13; N-7; A-0

Rationale:

- The measure has a long history of use since 2001.
- The outcome is important.
- The measure is meaningful, reliable, and valid.
- It can be calculated electronically.
- Disparities information presented.

If applicable, Conditions/Questions for Developer: The developer was asked to update the evidence information in the submission.

RECOMMENDATION: MAINTAIN ENDORSEMENT

522

0277 Congestive heart failure admission rate (PQI 8)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percent of county population with an admissions for CHF

Numerator Statement: All discharges of age 18 years and older with ICD-9-CM principal diagnosis code for CHF

Denominator Statement: Population in Metro Area or county, age 18 years and older

Exclusions: None

Adjustment/Stratification: Risk-adjustment method widely or commercially available. The predicted value for each case is computed using a logistic regression model and covariates for gender and age in years (in 5-year age groups). The reference population used in the model is the universe of discharges for states that participate in the HCUP State Inpatient Databases (SID) for the year 2007 (updated annually), a database consisting of 43 states and approximately 30 million adult discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., county, state, and region). The risk-adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate. Observed rates may be stratified by gender, age (5-year age groups), race/ethnicity.

Level of Analysis: Population: Counties or cities Type of Measure: Outcome

Data Source: Electronic administrative data/claims

Measure Steward: Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, MD 20850

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-15; N-5

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Performance gaps by age, gender, and income.
- No benchmark for the indicator.

0277 Congestive heart failure admission rate (PQI 8)

- Some concern that use of the measure may create perverse incentives to improve performance by reducing admissions without improving quality of care.
- Some concern about interpretation of "preventable".
- An "ambulatory care sensitive measure".

2. Scientific Acceptability of Measure Properties: C-5; P-15; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Precisely defined.
- Very strong disparities
- Risk adjusted by age and gender only.
- Committee would like to see stratification for race/disparities
- Does not include emergency department (ED) admission, only hospital admission.

3. Usability: C-2; P-18; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Committee would like to see stratification for race.
- Developer notes that county and state health departments have used this as a tool to allocate resources toward primary care workforce development in communities that are felt to have a disproportionate burden of avoidable hospitalizations.

4. Feasibility: C-9; P-11; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

Does not include ED admissions data; only hospital admission data.

Does the Measure Meet Criteria for Endorsement?: Y-19; N-1

Rationale:

- Population health measures in use for more than 10 years.
- Gaps by age, gender, and income.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: Recommend for endorsement

523

0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The measure estimates a hospital-level risk-standardized mortality rate (RSMR), defined as death from any cause within 30 days after the index admission date, for patients discharged from the hospital with a principal diagnosis of HF.

Numerator Statement: This outcome measure does not have a traditional numerator and denominator like a core process measure (e.g., percentage of adult patients with diabetes aged 18-75 years receiving one or more hemoglobin A1c tests per year); thus, we are using this field to define the outcome.

The outcome for this measure is 30-day all-cause mortality. We define mortality as death from any cause within 30 days of the index admission date for patients discharged from the hospital with a principal diagnosis of HF.

Denominator Statement: Note: This outcome measure does not have a traditional numerator and denominator like a core process measure; thus, we are using this field to define the patient cohort and to define exclusions to the patient cohort.

The cohort includes admissions for Medicare FFS beneficiaries age 65 years or older discharged from the hospital with a principal diagnosis of HF (ICD-9-CM codes 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, and 428.xx) and with a complete claims history for the 12 months prior to admission. Patients who are transferred from one acute care facility to another must have a principal discharge diagnosis of HF at both hospitals. The initial hospital for a transferred patient is designated as the responsible institution for the episode.

0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization

If a patient has more than one HF admission in a year, one hospitalization is randomly selected for inclusion in the measure. Exclusions: The measures exclude admissions for patients:

- who were discharged on the day of admission or the following day and did not die or get transferred (because it is less likely they had a significant HF diagnosis);
- who were transferred from another acute care hospital (because the death is attributed to the hospital where the patient was initially admitted);
- with inconsistent or unknown mortality status or other unreliable data (e.g., date of death precedes admission date);
- enrolled in the Medicare Hospice program any time in the 12 months prior to the index hospitalization including the first day of the index admission (because it is likely these patients are continuing to seek comfort measures only);
- who were discharged alive and against medical advice (AMA) (because providers did not have the opportunity to deliver full care and prepare the patient for discharge);
- that were not the first hospitalization in the 30 days prior to a patient's death. We use this criteria to prevent attribution of a
 death to two admissions.

Adjustment/Stratification: Risk-adjustment devised specifically for this measure/condition. Our approach to risk adjustment was tailored to and appropriate for a publicly reported outcome measure, as articulated in the American Heart Association (AHA) Scientific Statement, "Standards for Statistical Models Used for Public Reporting of Health Outcomes" (Krumholz et al., 2006).

The measure employs a hierarchical logistic regression model (a form of hierarchical generalized linear model [HGLM]) to create a hospital level 30-day RSMR. This approach to modeling appropriately accounts for the structure of the data (patients clustered within hospitals), the underlying risk due to patients' comorbidities, and sample size at a given hospital when estimating hospital mortality rates. In brief, the approach simultaneously models two levels (patient and hospital) to account for the variance in patient outcomes within and between hospitals (Normand et al., 2007). At the patient level, each model adjusts the log-odds of mortality within 30-days of admission for age, sex, selected clinical covariates and a hospital-specific intercept. The second level models the hospital-specific intercepts as arising from a normal distribution. The hospital intercept, or hospital specific effect, represents the hospital contribution to the risk of mortality, after accounting for patient risk and sample size, and can be inferred as a measure of quality. The hospital-specific intercepts are given a distribution in order to account for the clustering (non-independence) of patients within the same hospital. If there were no differences among hospitals, then after adjusting for patient risk, the hospital intercepts should be identical across all hospitals. Candidate and Final Risk-adjustment Variables: Candidate variables were patient-level risk-adjustors that are expected to be predictive of mortality, based on empirical analysis, prior literature, and clinical judgment, including demographic factors (age, sex) and indicators of comorbidity and disease severity. For each patient, covariates were obtained from Medicare claims extending 12 months prior to and including the index admission. The model adjusted for case differences based on the clinical status of the patient at the time of admission. We used condition categories (CCs), which are clinically meaningful groupings of more than 15,000 ICD-9-CM diagnosis codes. We did not risk-adjust for CCs that were possible adverse events of care and that were only recorded in the index admission. In addition, only comorbidities that conveyed information about the patient at that time or in the 12-months prior, and not complications that arose during the course of the hospitalization were included in the risk-adjustment. The final set of risk-adjustment variables are: Demographic

- Age-65 (years above 65, continuous)
- Male

Cardiovascular

- History of PTCA
- History of CABG
- Congestive heart failure
- Acute myocardial infarction
- Unstable angina
- Chronic atherosclerosis
- Cardio-respiratory failure and shock
- Valvular and rheumatic heart disease

Comorbidity

- Hypertension
- Stroke
- Renal failure
- Pneumonia

0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization

- Diabetes and DM complications
- Protein-calorie malnutrition
- Dementia and senility
- Hemiplegia, paraplegia, paralysis, functional disability
- Peripheral vascular disease
- Metastatic cancer, acute leukemia, and other severe cancers
- Trauma in last year
- Major psych disorders
- Chronic liver disease

References:

Krumholz HM, Brindis RG, Brush JE, et al. 2006. Standards for Statistical Models Used for Public Reporting of Health Outcomes: An American Heart Association Scientific Statement From the Quality of Care and Outcomes Research Interdisciplinary Writing Group: Cosponsored by the Council on Epidemiology and Prevention and the Stroke Council Endorsed by the American College of Cardiology Foundation. Circulation 113: 456-462.

Normand S-LT, Shahian DM. 2007. Statistical and Clinical Aspects of Hospital Outcomes Profiling. Stat Sci 22 (2): 206-226. Results of this measure will not be stratified.

Level of Analysis: Facility/Agency Type of Measure: Outcome

Data Source: Electronic administrative data/claims

Measure Steward: Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-02-01, Baltimore, MD 21244-9045

STEERING COMMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Most common admission under Medicare; second most costly total bill.
- Outcome measure.
- Important outcome measure

2. Scientific Acceptability of Measure Properties: C-19; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Data were published in a manuscript last year, looking at long-term trends in cardiovascular quality and outcomes.
- Risk adjustment used is administrative data. Methodology was validated against clinical data.

3. Usability: C-17; P-2; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Measure is currently in use.
- Public may not view data on website as often as was hoped, but doctors and administrators are using the data for internal
 quality improvement.

4. Feasibility: C-19; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- Measure is in use and publicly reported.
- Uses administrative data.

Does the Measure Meet Criteria for Endorsement?: Y-17; N-1; A-0

Rationale:

• A detailed, comprehensive submission form demonstrates that the measure meets all the criteria.

0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization

- Published in the literature.
- In use and publicly reported.

If applicable, Conditions/Questions for Developer: Disparities in race and socioeconomic status have been reported at the patient level. Does CMS plan on stratifying the measure?

Response: Disparities at the hospital level haven't been seen in facilities with higher percentages of African-American patients.

RECOMMENDATION: MAINTAIN ENDORSEMENT

On June 3, 2011, NQF and the Steering Committee were advised that the developer will complete testing of this measure on all payer data. The Committee will evaluate possible revisions to the measure as an addendum.

524

0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure hospitalization

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The measure estimates a hospital 30-day risk-standardized readmission rate (RSRR), defined as readmission for any cause within 30 days after the date of discharge of the index admission for patients discharged from the hospital with a principal diagnosis of heart failure (HF).

Numerator Statement: This outcome measure does not have a traditional numerator and denominator like a core process measure (e.g., percentage of adult patients with diabetes aged 18-75 years receiving one or more hemoglobin A1c tests per year); thus, we are using this field to define the outcome.

The outcome for this measure is 30-day all-cause readmission. We define this as readmission for any cause within 30 days from the date of discharge of the index HF admission.

In addition, if a patient has one or more admissions within 30 days of discharge from the index admission, only one was counted as a readmission.

Denominator Statement: Note: This outcome measure does not have a traditional numerator and denominator like a core process measure; thus, we are using this field to define the patient cohort and to define exclusions to the patient cohort.

The cohort includes admissions for Medicare fee-for-service (FFS) beneficiaries age 65 years or older discharged from the hospital with a principal diagnosis of HF (ICD-9-CM codes 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, and 428.xx) and with a complete claims history for the 12 months prior to admission.

Exclusions: We excluded admissions for patients:

- with an in-hospital death (because they are not eligible for readmission);
- without at least 30 days post-discharge enrollment in Medicare FFS (because the 30-day readmission outcome cannot be assessed in this group);
- transferred to another acute care facility (When a patient is transferred from one acute care hospital to another, these multiple
 contiguous hospitalizations are considered one episode of care. Readmissions for transferred patients are attributed to the
 hospital that ultimately discharges the patient to a non-acute care setting.);
- discharged against medical advice (AMA) (because providers did not have the opportunity to deliver full care and prepare the
 patient for discharge);
- admitted with HF within 30 days of discharge from an index admission (Admissions within 30 days of discharge of an index admission will be considered readmissions. No admission is counted as a readmission and an index admission. The next eligible admission after the 30-day time period following an index admission will be considered another index admission.)

Adjustment/Stratification: Risk-adjustment devised specifically for this measure/condition. Our approach to risk adjustment was tailored to and appropriate for a publicly reported outcome measure, as articulated in the American Heart Association (AHA) Scientific Statement, "Standards for Statistical Models Used for Public Reporting of Health Outcomes" (Krumholz et al., 2006).

The measure employs a hierarchical logistic regression model (a form of hierarchical generalized linear model [HGLM]) to create a hospital level 30-day RSRR. This approach to modeling appropriately accounts for the structure of the data (patients clustered within hospitals), the underlying risk due to patients' comorbidities, and sample size at a given hospital when estimating hospital readmission rates. In brief, the approach simultaneously models two levels (patient and hospital) to account for the variance in patient outcomes within and between hospitals (Normand et al., 2007). At the patient level, each model adjusts the log-odds of readmission within 30-days of admission for age, sex, selected clinical covariates and a hospital-specific intercept. The second level models the hospital-specific intercepts as arising from a normal distribution. The hospital intercept, or hospital specific effect, represents the hospital contribution to the risk of readmission, after accounting for patient risk and sample size, and can be inferred as a measure of quality. The hospital-specific intercepts are given a distribution in order to account for the clustering (non-independence) of patients within the same hospital.

0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure hospitalization

If there were no differences among hospitals, then after adjusting for patient risk, the hospital intercepts should be identical across all hospitals.

Candidate and Final Risk-adjustment Variables: Candidate variables were patient-level risk-adjustors that are expected to be predictive of readmission, based on empirical analysis, prior literature, and clinical judgment, including demographic factors (age, sex) and indicators of comorbidity and disease severity. For each patient, covariates were obtained from Medicare claims extending 12 months prior to and including the index admission. The model adjusted for case differences based on the clinical status of the patient at the time of admission. We used condition categories (CCs), which are clinically meaningful groupings of more than 15,000 ICD-9-CM diagnosis codes. In addition, only comorbidities that conveyed information about the patient at that time or in the 12-months prior, and not complications that arose during the course of the hospitalization were included in the risk-adjustment. We did not risk-adjust for CCs that were possible adverse events of care and that were only recorded in the index admission,

The final set of risk-adjustment variables are:

Demographic

- Age-65 (years above 65, continuous)
- Male

Cardiovascular

- History of CABG
- Cardio-respiratory failure or shock
- Congestive heart failure
- Acute coronary syndrome
- Coronary atherosclerosis or angina
- Valvular or rheumatic heart disease
- Specified arrhythmias
- Other or unspecified heart disease
- Vascular or circulatory disease

Comorbidity

- Metastatic cancer or acute leukemia
- Cancer
- Diabetes or DM complications
- Protein-calorie malnutrition
- Disorders of fluid, electrolyte, acid-base
- Liver or biliary disease
- · Peptic ulcer, hemorrhage, other specified gastrointestinal disorders
- Other gastrointestinal disorders
- Severe hematological disorders
- Iron deficiency or other anemias and blood disease
- Dementia or other specified brain disorders
- Drug/alcohol abuse/dependence/psychosis
- Major psychiatric disorders
- Depression
- Other psychiatric disorders
- · Hemiplegia, paraplegia, paralysis, functional disability
- Stroke
- Chronic obstructive pulmonary disease
- Fibrosis of lung or other chronic lung disorders
- Asthma
- Pneumonia
- End stage renal disease or dialysis
- Renal failure
- Nephritis
- Other urinary tract disorders

0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure hospitalization

Decubitus ulcer or chronic skin ulcer

References:

Krumholz HM, Brindis RG, Brush JE, et al. 2006. Standards for Statistical Models Used for Public Reporting of Health Outcomes: An American Heart Association Scientific Statement From the Quality of Care and Outcomes Research Interdisciplinary Writing Group: Cosponsored by the Council on Epidemiology and Prevention and the Stroke Council Endorsed by the American College of Cardiology Foundation. Circulation 113: 456-462.

Normand S-LT, Shahian DM. 2007. Statistical and Clinical Aspects of Hospital Outcomes Profiling. Stat Sci 22 (2): 206-226. Results of this measure will not be stratified.

Level of Analysis: Facility/Agency

Type of Measure: Outcome Data Source: Electronic administrative data/claims

Measure Steward: Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-02-01, Baltimore, MD 21244-

9045

STEERING COMMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

Heart failure is the number one cause of hospitalization and readmission among Medicare members.

2. Scientific Acceptability of Measure Properties: C-18; P-1; M-0; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Very well specified.
- Disparities information should be publicly disclosed on Hospital Compare.
- Stratified analyses are done instead of controlling for socioeconomic status.

3. Usability: C-18; P-1; M-0; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Has been in use without any major issues for some time.
- Captures an important domain of quality that's not captured in the mortality measure or other measures reviewed.

4. Feasibility: C-18; P-1; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- Data generated during care process. Uses administrative data.
- Data could be obtained from electronic health records or paper.
- Isn't particularly susceptible to inaccuracies and is easily implemented.

Does the Measure Meet Criteria for Endorsement?: Y-20; N-0; A-0

Rationale:

- High readmission rates—20% within 30 days; 50% within 1 years
- Significant variation
- Addresses all criteria

If applicable, Conditions/Questions for Developer: Strongly recommend that disparities data be reported on Hospital Compare. Developer Response: Disparities surveillance is on-going and reported on another CMS website. Will consider recommendation to include in Hospital Compare.

RECOMMENDATION: MAINTAIN ENDORSEMENT

On June 3, 2011, NQF and the Steering Committee were advised that the developer will complete testing of this measure on all payer data. The Committee will evaluate possible revisions to the measure as an addendum.

Recommended for endorsement and placement in reserve status:

0135 Evaluation of left ventricular systolic function (LVS)

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of heart failure (HF) patients with documentation in the hospital record that left ventricular systolic (LVS) function was evaluated before arrival, during hospitalization, or is planned for after discharge.

Numerator Statement: HF patients with documentation in the hospital record that LVS function was evaluated before arrival, during hospitalization, or is planned for after discharge

Denominator Statement: HF patients (ICD-9-CM principal diagnosis of HF: 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, 428.0, 428.1, 428.20, 428.21, 428.22, 428.23, 428.30, 428.31, 428.32, 428.33, 428.40, 428.41, 428.42, 428.43, 428.9)

Exclusions: Exclusions:

- <18 years of age</p>
- Patients who have a length of stay greater than 120 days
- Discharged to another hospital
- Expired
- Left against medical advice
- Discharged to home for hospice care
- Discharged to a health care facility for hospice care
- Patients enrolled in clinical trials
- Patients with comfort measures only documented
- Reasons for no LVS function evaluation documented by a physician, advanced practice nurse, or physician assistant
- Patients who had a left ventricular assistive device (LVAD) or heart transplant procedure during hospital stay (ICD-9-CM procedure code of LVAD or Heart Transplant: 33.6, 37.51, 37.52, 37.53, 37.54, 37.60, 37.62, 37.63, 37.65, 37.66, 37.68)

Adjustment/Stratification: No risk adjustment necessary N/A

Level of Analysis: Facility/Agency, Population: National Type of Measure: Process

Data Source: Paper medical record/flow-sheet

Measure Steward: Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-15; N-3

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Important to document this measurement; impacts long-term outcome and therapy.
- Current performance is very high.
- Disparities evident among Native American population.
- No explicit guideline recommendation as to what an appropriate time interval is.

2. Scientific Acceptability of Measure Properties: C-7; P-6; M-5; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- Concern with misinterpretation of measure so that testing is done at every hospitalization which is not required by the
 measure.
- Data abstraction may be difficult. Documentation challenge if test wasn't done during that hospitalization period.

3. Usability: C-5; P-10; M-4; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

 May stimulate overuse of imaging because of misinterpretation of measures inclusions—test done before or after hospitalization is credited

4. Feasibility: C-5; P-8; M-6; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to

0135 Evaluation of left ventricular systolic function (LVS)

inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

- Unintended consequence may be to encourage overuse.
- Upcoding issues with heart failure diagnosis.
- Implemenation issues—difficult to find data in charts.

Does the Measure Meet Criteria for Endorsement?: Y-5; N-13; A-0

Rationale:

- Current high performance. Possibly candidate for "topped out" category.
- Concern that this measure is a starting point for therapy, and if eliminated could impact other measures.
- A composite format may better serve this measure.

If applicable, Conditions/Questions for Developer:

RECOMMENDATION: MAINTAIN ENDORSEMENT AND PLACEMENT IN RESERVE STATUS

Additional recommendation: The Steering Committee also recommended that the measure be recalculated again in 3-5 years to monitor performance.

527

528 Not recommended:

0077 Heart failure: Symptom and activity assessment

For More Information: Complete Measure Submission: Meeting/Call Proceedings

Description: Percentage of patient visits for those patients aged 18 years and older with a diagnosis of heart failure with quantitative results of an evaluation of both current level of activity and clinical symptoms documented

Numerator Statement: Patient visits with quantitative results of an evaluation of both current level of activity and clinical symptoms documented*

*Evaluation and quantitative results documented should include:

- documentation of New York Heart Association (NYHA) Class OR
- documentation of completion of a valid, reliable, disease-specific instrument (e.g., Kansas City Cardiomyopathy Questionnaire, Minnesota Living with Heart Failure Questionnaire, Chronic Heart Failure Questionnaire)

Denominator Statement: All patient visits for those patients aged 18 years and older with a diagnosis of heart failure

Exclusions: Documentation of medical reason(s) for not evaluating both current level of activity and clinical symptoms (eg, severe cognitive or functional impairment)

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Clinician: Individual Type of Measure: Process

Data Source: Electronic administrative data/claims, Electronic Clinical Data, Paper medical record/flow-sheet, Registry data

Measure Steward: American Medical Association, 515 N State St., Chicago, IL 60654

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-8; N-10

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Process measure based on a clinical guideline recommendation supported by Level C evidence (expert consensus).
- There is evidence to suggest that the variability in provider determination of NYHA class is considerable.
- Use of psychometrically standardized questionnaires is more defensible; however, there is no evidence of a link between performing and assessment and outcome..
- Unclear if there is a gap in documentation or a gap in clinically asking or assessing.
- Testing results not included with submission.

Steering Committee Recommendation for Endorsement: Not recommended.

Rationale: Does not meet the criterion for importance to measure.

- What is the evidence of realtionship to outcomes?
- Gap is likely a gap in documentation.

0077 Heart failure: Symptom and activity assessment

The developers submitted a <u>letter to the Steering Committee</u> disagreeing with the Committee's evaluation and requested a reconsideration of the measure evaluation citing the following:

- "a notable gap in patient-centric measures that would focus attention on patient-reported outcomes, including their symptoms, function and health-related quality of life"; and
- symptoms are an outcome and there are racial disparities in symptom management; they want to lay a foundation for future measures of efficacy and appropriateness.

The Steering Committee agreed that the measure, as specified, is a process measure that is not linked to an intermediate or ultimate outcome and additionally noted:

- There is no reliability or validity data that say the results distinguish quality at the physician level.
- Evidence is lacking. What is the data/evidence that just doing an assessment is related to patient satisfaction, better outcomes, more or less angioplasty, or less MIs?
- What is the gap? General perception that clinicians are not doing this well. PINNACLE data = 85.5%.
- Testing data not provided.

Steering Committee re-vote on Importance: Y-6, N-9

RECOMMENDATION: Not recommended

529

530

962 Composite measure of hospital quality for heart failure (HF)

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: A composite measure of in-hospital process- and outcome-of-care for Heart Failure (HF) patients.

Composite Numerator Statement: For the process-of-care domain, the numerator is equal to the weighted sum of four terms. Each term is equal to the ratio of the hospital's raw performance rate to the national performance rate for the indicator. The weight is equal to the total number of observations, that is, the number of patients 'at risk' for the indicator.

For the outcome-of-care domain, the numerator is equal to the weighted sum of two terms. Each term is equal to the ratio of the hospital's risk-standardized performance rate to the national performance rate for the indicator. The weight is equal to the total number of eligible discharges for the indicator.

Denominator Statement: For the process-of-care domain, the denominator is equal to the total number of observations for all HF process indicators. It is thus equal to the number of patients 'at risk' for the four process indicators.

For the outcome-of-care domain, the denominator is equal to the total number of observations for all HF outcome indicators. It is thus equal to the number of eligible discharges for the two outcome indicators.

Exclusions: The following two criteria were applied as exclusion restrictions:

- 1. Hospitals with less than five eligible patient cases for the process-of-care indicators and less than 25 eligible discharges for the outcome-of-care indicators.
- 2. Hospitals that were missing rates for one or more process-of-care and/or outcome-of-care indicators.

Adjustment/Stratification:

Level of Analysis: Hospital Type of Measure: Composite

Data Source: The composite is constructed from component measures posted on the Hospital Compare website.

Measure Steward: Centers for Medicare & Medicaid Services

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-8; N-10

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

• While a composite is desirable, the components are not the right ones.

Does the Measure Meet Criteria for Endorsement?: <u>Not recommended.</u> Rationale: <u>Does not meet Importance to Measure and Report criteria:</u>

962 Composite measure of hospital quality for heart failure (HF)

- Includes smoking measure no longer endorsed by NQF and discharge instructions measure that is not recommended to maintain endorsement.
- Does not address improtant aspects of care for HF: beta blocker use; better discharge measure; cardiac rehab.
- The process of care measures are on all patients; the outcome measures (mortality and readmissions) are Medicare only.
- Weighting should be by impact.

If applicable, Conditions/Questions for Developer: Why not create an all-or-none composite? What about other important aspects of care for HF patients such as beta blocker use, patient education and self management, functional status and symptom control or a valid smoking cessation measure?

Response: They were limited to the measures used on Hospital Compare.

RECOMMENDATION: Not recommended

531

533

532 **HYPERTENSION**

Recommended for endorsement:

0018 Controlling high blood pressure

For More Information: Detailed Measure Specifications; Complete Measure Submission; Meeting/Call Proceedings

Description: The percentage of members 18–85 years of age who had a diagnosis of hypertension (HTN) and whose blood pressure (BP) was adequately controlled (<140/90) during the measurement year. Use the Hybrid Method for this measure.

Numerator Statement: The number of members in the denominator whose most recent BP is adequately controlled during the measurement year. For a member's BP to be controlled, both the systolic and diastolic BP must be <140/90 (adequate control). To determine if a member's BP is adequately controlled, the organization must identify the representative BP.

Denominator Statement: Patients 18-85 with hypertension. A patient is considered hypertensive if there is at least one outpatient encounter with a diagnosis of HTN during the first 6 months of the measurement year.

Exclusions:

- Exclude from the eligible population all members with evidence of end-stage renal disease (ESRD) (Table CBP-C) on or prior to December 31 of the measurement year. Documentation in the medical record must include a dated note indicating evidence of ESRD. Documentation of dialysis or renal transplant also meets the criteria for evidence of ESRD.
- Exclude from the eligible population all members with a diagnosis of pregnancy (Table CBP-C) during the measurement year.
- Exclude from the eligible population all members who had an admission to a nonacute inpatient setting any time during the measurement year. Refer to Table FUH-B for codes to identify nonacute care.

Adjustment/Stratification: No risk adjustment necessary.

Level of Analysis: Clinician. Clinician: Group/Practice. Clinician: Individual. Health Plan

Type of Measure: Outcome

Data Source: Administrative claims, Electronic administrative data/claims, Electronic Clinical Data, Electronic Clinical Data: Electronic Health Record. Paper medical record/flow-sheet. Paper Records: retooled eMeasure

Measure Steward: National Committee for Quality Assurance, 1100 13th Street NW, Suite 1000, Washington, DC 20005

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-20; N-0

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Important intermediate outcome measure.
- Strong evidence for relationship to long-term outcomes.
- There is less precision in the evidence for BP targets for patients greater than 85 years.

2. Scientific Acceptability of Measure Properties: C-4; P-12; M-3; N-0

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

• The level of measurement or analysis should be clinician and health plan. Submission form indicates clinician only.

0018 Controlling high blood pressure

Intolerance of low BP not included.

3. Usability: C-12; P-6; M-1; N-0

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

- Value added is in exclusions specified in this measure.
- Measure is essentially the same as the PCPI measure (0013).

4. Feasibility: C-12; P-8; M-0; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

Measure has been retooled for EHRs.

Does the Measure Meet Criteria for Endorsement?: Y-19; N-0; A-0

Rationale: Clearer measurement defintion than comparable PCPI measure (0013).

If applicable, Conditions/Questions for Developer:

- 1. How is timeframe for control defined?
- 2. How was age 85 chosen?
- 3. Is white coat hypertension in the exclusions?
- 4. Why isn't home blood pressure monitoring included?

Developer Response:

- 1. From onset of diagnosis to the following 12 month period.
- 2. The age was chosen as a result of multiple comorbidities and functional status issues.
- 3. No. This is office-based and the last measurement recorded.
- 4. This measure hasn't been tested to incorporate home monitoring.

Steering Committee Follow-up:

4. As new JNC-8 guidelines are released, the inclusion of home monitoring is recommended, as well as age inclusions. Developer Follow-up:

4. May consider retesting of the measure.

RECOMMENDATION: MAINTAIN ENDORSEMENT

534

535 Not recommended:

0013 Hypertension: Blood pressure management

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Endorsed measure 0013 was originally Blood pressure measurement *Percentage of patient visits with blood pressure measurement recorded among all patient visits for patients aged* \geq 18 years with diagnosed hypertension. (Retooled eMeasure)

Endorsed measure 0017 was originally Hypertension plan of care Percentage of patient visits during which either systolic blood pressure \geq 140 mm Hg or diastolic blood pressure \geq 90 mm Hg, with documented plan of care for hypertension. The revised submission replaces both measures.

Description: Percentage of patients aged 18 years and older with a diagnosis of hypertension with a blood pressure <140/90 mm Hg OR patients with a blood pressure ≥ 140/90 mm Hg and prescribed two or more anti-hypertensive medications during the most recent office visit within a 12-month period

Numerator Statement: Patients with a blood pressure <140/90 mm Hg OR

Patients with a blood pressure ≥ 140/90 mm Hg and prescribed two or more anti-hypertensive medications during the most recent office visit within a 12-month period

Instructions:

- Report number of patients for 1st numerator component (outcome) AND
- Report number of patients for 2nd numerator component (process) AND

0013 Hypertension: Blood pressure management

Report total number of patients for all numerator components

Denominator Statement: All visits for patients aged 18 years and older with a diagnosis of hypertension Exclusions:

- Documentation of medical reason(s) for not prescribing two or more anti-hypertensive medications (e.g., allergy, intolerant, postural hypotension)
- Documentation of patient reason(s) for not prescribing two or more anti-hypertensive medications (e.g., patient declined)
- Documentation of system reason(s) for not prescribing two or more anti-hypertensive medications (e.g., financial reasons)

Adjustment/Stratification: No risk adjustment necessary

Level of Analysis: Type of Measure: Process

Data Source: Electronic administrative data/claims, Electronic Clinical Data, Paper medical record/flow-sheet, Registry data

Measure Steward: American Medical Association, 515 N State St., Chicago, IL 60654

This is an updated version of measure 0013 Blood pressure measurement combined with 0017 Plan of care.

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-19; N-1

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- This is a new measure combining intermediate outcome and plan of care.
- More evidence is needed to support that two or more anti-hypertensive medications is considered a positive outcome without some additional definition of the measure related to the extent of control achieved (e.g., reduction in BP by a certain % from baseline after medications prescribed).
- Concern that credit could be given for undertreatment.

2. Scientific Acceptability of Measure Properties: C-3; P-5; M-7; N-5

(2a. Precise specifications; 2b. Reliability testing; 2c. Validity testing; 2d. Exclusions justified; 2e. Risk adjustment/stratification; 2f. Meaningful differences; 2g. Comparability; 2h. Disparities)

Rationale:

- No current performance data. Reliability and validity are not known.
- Based on more than one BP measurement.
- BP values from home, office or 24-hour monitoring.
- Unintended consequence for the two medication threshold if patients should be on three.
- Concerns for patients that don't tolerate BP <140/90 versus undertreatment of patients who should be at target.

3. Usability: C-4; P-9; M-6; N-1

(3a. Meaningful/useful for public reporting and quality improvement; 3b. Harmonized; 3c. Distinctive or additive value to existing measures)

Rationale:

Title seems misleading because it captures patients who are not under control.

4. Feasibility: C-9; P-6; M-5; N-0

(4a. Clinical data generated during care process; 4b. Electronic sources; 4c. Exclusions – no additional data source; 4d. Susceptibility to inaccuracies/ unintended consequences identified 4e. Data collection strategy can be implemented)

Rationale:

Data are generated during care; collection easily implemented.

Does the Measure Meet Criteria for Endorsement?: Y-6; N-14; A-0

Rationale:

- Lack of evidence for two or more drugs component.
- Reliablity and validity not known.
- Some patients may need three+ drugs—measure gives credit for patients that may be undertreated.
- New measure—no current performance data.

If applicable, Conditions/Questions for Developer:

- 1. What is the added value of this measure on top of previous ones?
- 2. Title seems misleading—it is not just BP control.

0013 Hypertension: Blood pressure management

Developer Response:

- Addresses other issues: blood pressure >140/90; includes ambulatory, home, and office monitoring.
- 2. Developer changed the title to "BP management".

RECOMMENDATION: Not recommended

536

0276 Hypertension admission rate (PQI 7)

For More Information: Complete Measure Submission; Meeting/Call Proceedings

Description: Percentage of county population with an admission for hypertension.

Numerator Statement: All discharges of age 18 years and older with ICD-9-CM principal diagnosis code for hypertension (see below). Denominator Statement: Population in Metro Area or county, age 18 years and older.

Exclusions: None

Adjustment/Stratification: Risk-adjustment method widely or commercially available. The predicted value for each case is computed using a logistic regression model and covariates for gender and age in years (in 5-year age groups). The reference population used in the model is the universe of discharges for states that participate in the HCUP State Inpatient Databases (SID) for the year 2007 (updated annually), a database consisting of 43 states and approximately 30 million adult discharges. The expected rate is computed as the sum of the predicted value for each case divided by the number of cases for the unit of analysis of interest (i.e., county, state, and region). The risk adjusted rate is computed using indirect standardization as the observed rate divided by the expected rate, multiplied by the reference population rate. Observed rates may be stratified by gender, age (5-year age groups), race / ethnicity.

Level of Analysis: Population: Counties or cities Type of Measure: Outcome

Data Source: Electronic administrative data/claims

Measure Steward: Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, MD 20850

STEERING COMMITTEE EVALUATION

1. Importance to Measure and Report: Y-7; N-11

(1a. Impact; 1b. Performance gap; 1c. Outcome or Evidence)

Rationale:

- Submitted documentation states "Little evidence exists regarding the validity of this indicator," and "some of the variance in age-sex adjusted rates does not reflect true differences in area performance."
- Patients with uncontrolled blood pressure are admitted for many reasons (CHF, AMI, stroke). Only hypertension as primary diagnosis is captured.
- Could be missing an important population.

Does the Measure Meet Criteria for Endorsement?: Do not recommend.

Rationale: Did not pass Importance criteria

If applicable, Conditions/Questions for Developer: How is this data better than NHANES or BRFSS?

Response: Intended to describe population health; designed for use at the geographic area level.

RECOMMENDATION: REMOVE ENDORSEMENT

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RETIRED MEASURES

- The measure developers have indicated that they no longer maintain the following measures and
- request retirement from NQF's measure portfolio. The Committee agreed that better measures
- have replaced these in NQF's portfolio.

0072 CAD: beta-blocker treatment after a heart attack (NCQA)	DESCRIPTION: Percentage of patients who have a claim indicating beta blocker therapy or who received an ambulatory prescription for beta-blockers rendered within 7 days after discharge.
0161 AMI inpatient mortality (risk-	DESCRIPTION: Percentage of acute myocardial infarction
adjusted) (The Joint Commission)	(AMI) patients who expired during hospital stay.
0165 Percutaneous coronary	DESCRIPTION: Percentage of patient admissions for
intervention (PCI) volume (ACC)	percutaneous coronary intervention (PCI) procedure.
0082 Heart Failure (HF) : Patient	DESCRIPTION: Percentage of patients who were provided
education (AMA PCPI)	with patient education on disease management and health
	behavior changes during one or more visit(s).
0084 Heart Failure (HF) : Warfarin	DESCRIPTION: Percentage of patients with HF who also
therapy patients with atrial fibrillation	have paroxysmal or chronic atrial fibrillation who were
(AMA PCPI)	prescribed warfarin therapy.
0085 Heart Failure (HF): Weight	DESCRIPTION: Percentage of patient visits for patients with
measurement (AMA PCPI)	HF with weight measurement recorded.

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546 NOTES

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1. Lloyd-Jones D, Adams RJ, Brown TM, et al. <u>Heart Disease and Stroke Statistics—2010</u> <u>Update. A Report from the American Heart Association Statistics Committee and Stroke Statistics Subcommittee & Circulation</u>. 2010;121:e1-e170.

551	APPENDIX A—SPECIFICATIONS FOR THE NATIONAL VOLUNTARY CONSENSUS
552	STANDARDS: CARDIOVASCULAR ENDORSEMENT MAINTENANCE, 2010: A
553	CONSENSUS REPORT
554	
555	0018 Controlling High Blood Pressure98
556 557	0066 Chronic Stable Coronary Artery Disease: ACE Inhibitor or ARB TherapyDiabetes or Left Ventricular Systolic Dysfunction (LVEF <40%)
558	0067 Chronic Stable Coronary Artery Disease: Antiplatelet Therapy105
559	0068 Ischemic Vascular Disease (IVD): Use of Aspirin or another Antithrombotic107
560	0071 Acute Myocardial Infarction (AMI): Persistence of Beta-Blocker Treatment After a Heart
561	Attack
562	0073 IVD: Blood Pressure Management
563	0074 Chronic Stable Coronary Artery Disease: Lipid Control
564	0075 IVD: Complete Lipid Profile and LDL Control <100122
565	0076 Optimal Vascular Care126
566	0079 Heart Failure: Left Ventricular Ejection Fraction Assessment (Outpatient Setting)
567	0081 Heart Failure: Angiotensin-Converting Enzyme (ACE) Inhibitor or Angiotensin Receptor
568	Blocker (ARB) Therapy for Left Ventricular Systolic Dysfunction
569	0083 Heart Failure : Beta-blocker therapy for Left Ventricular Systolic Dysfunction134
570	0132 Aspirin at arrival for acute myocardial infarction (AMI)136
571	0133 PCI mortality (risk-adjusted)©
572	0135 Evaluation of Left ventricular systolic function (LVS)142
573	0137 ACEI or ARB for left ventricular systolic dysfunction- Acute Myocardial Infarction (AMI)
574	Patients
575	0142 Aspirin prescribed at discharge for AMI148
576	0160 Beta-blocker prescribed at discharge for AMI151

577	0162 ACEI or ARB for left ventricular systolic dysfunction - Heart Failure (HF) Patients	154
578	0163 Primary PCI received within 90 minutes of Hospital Arrival	158
579	0164 Fibrinolytic Therapy received within 30 minutes of hospital arrival	161
580	0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart fail	lure
581	(HF) hospitalization	164
582	0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute	
583	myocardial infarction (AMI) hospitalization	168
584	0277 Congestive Heart Failure Admission Rate (PQI 8)	171
585	0286 Aspirin at Arrival	186
586	0288 Fibrinolytic Therapy Received Within 30 Minutes of ED Arrival	188
587	0289 Median Time to ECG	190
588	0290 Median Time to Transfer to Another Facility for Acute Coronary Intervention	192
589	0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure	
590	hospitalization	194
591	0355 Bilateral Cardiac Catheterization Rate (IQI 25)	198
592	0358 Congestive Heart Failure (CHF) Mortality Rate (IQI 16)	209
593	1522 ACE/ARB Therapy at Discharge for ICD implant patients with LVSD	213
594	1524 Assessment of Thromboembolic Risk Factors (CHADS2)	215
595	1525 Chronic Anticoagulation Therapy	218
596	1528 Beta Blocker at Discharge for ICD implant patients with a previous MI	221
597	1529 Beta Blocker at Discharge for ICD implant patients with LVSD	223
598	0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge (ACCF)	
599	0965 Therapy with ACE/ARB and beta blocker at discharge following ICD implantation (AC	<u>(CF)</u>
600		

601

0018 Controlling high blood pressure

	0018 Controlling high blood pressure
Steward	National Committee for Quality Assurance, 1100 13th Street NW, Suite 1000, Washington, DC 20005
Description	The percentage of patients 18-85 years of age who had a diagnosis of hypertension (HTN) and whose blood pressure (BP) was adequately controlled (<140/90) during the measurement year.
Туре	Outcome
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Paper medical record/flow-sheet
Level	Clinicians: Group, Clinicians: Individual
Setting	All settings, Ambulatory Care: Amb Surgery Center, Ambulatory Care: Clinic, Ambulatory Care: Emergency Dept, Ambulatory Care: Hospital Outpatient, Ambulatory Care: Office
Numerator Statement	The number of patients in the denominator whose most recent, representative BP is adequately controlled during the measurement year. For a member's BP to be controlled, both the systolic and diastolic BP must be <140/90mm Hg.
Numerator Details	Time Window: The measurement year.
	The number of patients in the denominator whose most recent, representative BP is adequately controlled during the measurement year. For a member's BP to be controlled, both the systolic and diastolic BP must be <140/90 mm Hg. Follow these steps to identify the representative BP:
	• Identify the most recent blood pressure reading noted during the measurement year. The reading must occur after the date when the diagnosis of hypertension was made or confirmed. Do not include readings that meet the following criteria: taken during an acute inpatient stay or an ED visit, taken during an outpatient visit that was for the sole purpose of having a diagnostic test or surgical procedure performed (e.g., sigmoidoscopy, removal of a mole), taken the same day as a major diagnostic procedure (e.g., stress test, administration of IV contrast for a radiology procedure, endoscopy), reported by or taken by the patient, documentation of "VS within normal limits" or "vital signs normal".
	 Identify the lowest systolic and lowest diastolic reading from the most recent blood pressure notation in the medical record. If multiple readings were recorded for a single date, use the lowest systolic and the lowest diastolic reading on that date as the representative blood pressure. Results do not need to come from the same reading.
Denominator Statement	Patients 18-85 with hypertension. A patient is considered hypertensive if there is at least one outpatient encounter with a diagnosis of HTN during the first six months of the measurement year.
Denominator Categories	Female; Male 18-85 years
Denominator Details	Time Window: Age range verified as of December 31st of the measurement year, while the hypertensive diagnosis is verified in the first 6 months of the measurement year.
	Patients 18-85 as of December 31st of the measurement year who meet the following inclusion criteira:

Continuous enrollment using health plan data: Patients continuously enrolled during the measurement year with no more than one gap in enrollment of up to 45 days during the measurement year. Continuous enrollment using non-health plan data: any enrollment, claim or encounter transaction any time during the measurement year.	
Event/Diagnosis: Hypertensive: At least one outpatient encounter (Table CBP-B) with a diagnosis of hypertension (Table CBP-A) during the first six months of the measurement year.	
Table CBP-A: Codes to Identify Hypertension	
Description ICD-9-CM Diagnosis	
Hypertension 401	
Table CBP-B: Codes to Identify Outpatient Visits	
Description CPT	
Outpatient visits: 99201-99205, 99211-99215, 99241-99245, 99384-99387, 99394-99397	
The diagnosis of hypertension must be confirmed by chart review on or before June 30 of the measurement year finding notation of one of the following: HTN, High BP, Elevated BP, Borderline HTN, Intermittent HTN, History of HTN, Hypertensive vascular disease, Hyperpiesia, Hyperpiesis.	
Exclude from the eligible population all patients with evidence of end-stage renal disease (ESRD) (including dialysis or renal transplant), all patients who are pregnant, and all patients who had an admission to a nonacute inpatient setting on or prior to December 31 of the measurement year.	
Exclude from the eligible population all patients with evidence of end-stage renal disease (ESRD) on or prior to December 31 of the measurement year. Documentation in the medical record must include a dated note indicating evidence of ESRD. Documentation of dialysis or renal transplant also meets the criteria for evidence of ESRD.	
 Exclude from the eligible population all members with a diagnosis of pregnancy during the measurement year. Exclude from the eligible population all members who had an admission to a nonacute inpatient setting any time during the measurement year. Table CBP-C: Codes to Identify ESRD and Pregnancy Exclusions 	
Description: CPT HCPCS ICD-9-CM ICD-9-CM UB UB POS	
Diagnosis Procedure Revenue type of	
Bill	
Evidence 36145, 36800, G0257 585.5, 38.95 0367 72X 65	
of ESRD 36810, 36815, G0308-G0313 585.6, 39.27 080x	
36818, 36819, G0314-G0319 V42.0, 39.42 082x	
36820, 36821, G0322 V45.1 39.43 085x	
36831-36833, G0323 V56 39.53 088x	

	0018 Controlling high blood pressure
	50300, 50320,
	50340, 50360, G0327 54.98
	50365, 50370, G0392 55.6
	50380, 90920, G0393
	90921, 90924, S9339
	90925, 90935,
	90937, 90939,
	90940, 90945,
	90947, 90989,
	90993, 90997,
	90999, 99512
	Evidence of Pregnancy: ICD-9-CM Diagnosis: 630-679, V22, V23, V28
	Table FUH-B codes to identify non-acute inpatient exclusions:
	Hospice: UB Rev (0115, 0125, 0135, 0145, 0155, 0650, 0656, 0658, 0659), UB Type Bill (81x, 82x), POS (34)
	SNF: UB Rev (019x), UB Type Bill (21x, 22x, 28x), POS (31, 32)
	Hospital Transitional Care: UB Type Bill (18x)
	Rehabilitation: UB Rev (0118, 0128, 0138, 0148, 0158)
	Respite: UB Rev (0655)
	Intermediate Care Facility: POS (54)
	Residential Substance Abuse Treatment Facility: UB Rev (1002), POS (55)
	Psychiatric Residential Treatment Facility Center: HCPCS (T2048, H0017-19), UB Rev (1001), POS (56)
	Comprehensive Inpatient Rehabilitation Facility: POS (61)
Risk Adjustment	No risk adjustment necessary
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	

602

	0066 Chronic stable coronary artery disease: ACE Inhibitor or ARB Therapy—diabetes or left ventricular systolic dysfunction (LVEF <40%)
Steward	American Medical Association, 515 N. State St., Chicago, IL 60654
Description	Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period who also have diabetes or a current or prior LVEF <40% who were prescribed ACE inhibitor or ARB therapy.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Registry data URL www.pinnacleregistry.org Attachment PCPI_CAD-8_ACE-ARB Diabetes LVSD NQF 0066.pdf
Level	Clinicians: Group, Clinicians: Individual
Setting	Ambulatory Care: Clinic, Ambulatory Care: Hospital Outpatient, Ambulatory Care: Office, Assisted Living, Group homes, Home, Nursing home (NH)/Skilled Nursing Facility (SNF)
Numerator Statement	Patients who were prescribed ACE inhibitor or ARB therapy.* *Prescribed may include prescription given to the patient for ACE inhibitor or ARB therapy at one or more visits in the measurement period OR patient already taking ACE inhibitor or ARB therapy as documented in current medication list.
Numerator Details	Time Window: Once during measurement period.
	See attached for EHR Specifications.
	For Claims/Administrative: Report CPT II Code 4009F: Angiotensin converting enzyme (ACE) inhibitor or Angiotensin Receptor Blocker (ARB) therapy prescribed.
Denominator Statement	All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period who also have diabetes or a current or prior LVEF <40%.
Denominator Categories	Female; Male Aged 18 years and older
Denominator Details	Time Window: 12 consecutive months
	See attached for EHR Specifications.
	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, CPT).

0066 Chronic stable coronary artery disease: ACE Inhibitor or ARB Therapy—diabetes or left ventricular systolic dysfunction (LVEF <40%)
Documentation of medical reason(s) for not prescribing ACE inhibitor or ARB therapy (e.g., allergy, intolerant, pregnancy, renal failure due to ACE inhibitor, diseases of the aortic or mitral valve, other medical reasons).
Documentation of patient reason(s) for not prescribing ACE inhibitor or ARB therapy (e.g., patient declined, other patient reasons).
Documentation of system reason(s) for not prescribing ACE inhibitor or ARB therapy (e.g., lack of drug availability, other reasons attributable to the health care system).
See attached for EHR Specifications.
For Claims/Administrative:
Documentation of medical reason(s) for not prescribing ACE inhibitor or ARB therapy
 Append modifier to CPT II code 4009F-1P. Documentation of patient reason(s) for not prescribing ACE inhibitor or ARB therapy
 Append modifier to CPT II code 4009F-2P. Documentation of system reason(s) for not prescribing ACE inhibitor or ARB therapy
Append modifier to CPT II code 4009F-3P.
No risk adjustment necessary
Rate/proportion better quality = higher score
See attached for calculation algorithm.

603

	0067 Chronic stable coronary artery disease: Antiplatelet therapy
Steward	American Medical Association, 515 N. State St., Chicago, IL 60654
Description	Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period who were prescribed aspirin or clopidogrel.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Registry data URL www.pinnacleregistry.org Attachment PCPI_CAD-6_AntiplateletTherapy NQF 0067.pdf
Level	Clinicians: Group, Clinicians: Individual
Setting	Ambulatory Care: Clinic, Ambulatory Care: Hospital Outpatient, Ambulatory Care: Office, Assisted Living, Group homes, Home, Nursing home (NH)/Skilled Nursing Facility (SNF)
Numerator Statement	Patients who were prescribed aspirin or clopidogrel * within a 12 month period. *Prescribed may include prescription given to the patient for aspirin or clopidogrel at one or more visits in the measurement period OR patient already taking aspirin or clopidogrel as documented in current medication list.
Numerator Details	Time Window: Once during the measurement period.
	See attached for EHR Specifications.
	For Claims/Administrative: Report CPT II Code 4011F: Oral antiplatelet therapy prescribed
Denominator Statement	All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period.
Denominator Categories	Female; Male Aged 18 years and older
Denominator Details	Time Window: 12 consecutive months
	See attached for EHR Specifications.
	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, CPT).
Exclusions	Documentation of medical reason(s) for not prescribing aspirin or clopidogrel (e.g., allergy, intolerant, receiving other thienopyridine therapy, bleeding coagulation disorders, receiving warfarin therapy, other medical reasons).
	Documentation of patient reason(s) for not prescribing aspirin or clopidogrel (e.g., patient declined, other patient

	0067 Chronic stable coronary artery disease: Antiplatelet therapy
	reasons).
	Documentation of system reason(s) for not prescribing aspirin or clopidogrel (e.g., lack of drug availability, other reasons attributable to the health care system).
Exclusion	See attached for EHR Specifications.
Details	For Claims/Administrative:
	Documentation of medical reason(s) for not prescribing aspirin or clopidogrel
	 Append modifier to CPT II code 4011F-1P (in development). Documentation of patient reason(s) for not prescribing aspirin or clopidogrel
	 Append modifier to CPT II code 4011F-2P (in development). Documentation of system reason(s) for not prescribing aspirin or clopidogrel
	 Append modifier to CPT II code 4011F-3P (in development).
Risk Adjustment	No risk adjustment necessary
Stratification	
Type Score	Rate/proportion better quality = higher score
Algorithm	See attached for calculation algorithm.

	0068 Ischemic vascular disease (IVD): Use of aspirin or another antithrombotic
Steward	National Committee for Quality Assurance, 1100 13th Street NW, Suite 1000, Washington, DC 20005
Description	The percentage of patients 18 years and older with ischemic vascular disease who were discharged alive for acute myocardial infarction (AMI), coronary artery bypass graft (CABG) or percutaneous coronary interventions (PCI) from January 1-November 1 of the year prior to the measurement year, or who had a diagnosis of ischemic vascular disease (IVD) during the measurement year and the year prior to the measurement year and who had the following during the measurement year. • Use of aspirin or another antithrombotic
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Paper medical record/flow-sheet
Level	Clinicians: Group, Clinicians: Individual
Setting	All settings, Ambulatory Care: Clinic
Numerator Statement	Use of aspirin or another antithrombotic. Electronic specification: Documentation of use of aspirin or another antithrombotic during the measurement year. Refer to Table IVD-D to identify the code for prescribed oral anti-platelet therapy. Refer to Table IVD-E to identify medications for oral anti-platelet therapy. Medical Record Specification: Documentation of use of aspirin or another antithrombotic during the measurement year. At a minimum, documentation in the medical record must include a note indicating the date on which aspirin or another antithrombotic was prescribed or documentation of prescription from another treating physician.
Numerator Details	Time Window: 12 months
	Use of aspirin or another antithrombotic.
	Electronic specification:
	Documentation of use of aspirin or another antithrombotic during the measurement year. Refer to table IVD-D to identify the code for prescribed oral anti-platelet therapy. Refer to Table IVD-E to identify medications for oral anti-platelet therapy.
	Medical Record Specification:
	Documentation of use of aspirin or another antithrombotic during the measurement year. At a minimum,

	0068 Ischemic vascular disease (IVD): Use of aspirin or another antithrombotic			
	documentation in the medical record must include a note indicating the date on which aspirin or another antithrombotic was prescribed or documentation of prescription from another treating physician. Table IVD-D: Codes to Identify Prescribed Oral Anti-Platelet Therapy			
	Description CPT Category II ICD-9-CM Diagnosis			
	Oral anti-platelet therapy prescribed 4011F V58.63, V58.66			
	Table IVD-E: Oral Anti-Platelet Therapies			
	Description Prescription			
	Oral anti-platelet therapies • aspirin			
	• clopidogrel			
	aspirin-dipyridamole			
	• ticlopidine			
Denominator Statement	Patients 18 years or older as of December 31 of the measurement year discharged alive for AMI, CABG or PCI on or between January 1 and November 1 of the year prior to the measurement year or who had a diagnosis of IVD during both the measurement year and the year prior to the measurement year.			
Denominator Categories	Female; Male 18 years of age and older			
Denominator Details	Time Window: From January 1st of the year prior to the measurement year through December 31st of the measurement year.			
	Patients 18 years or older as of December 31 of the measurement year.			
	Patient inclusion criteria:			
	For physician assessment with generated from a health plan: continuous medical benefit enrollment for the measurement year, with no more than one gap in continuous enrollment of up to 45 days during the measurement year. To determine continuous enrollment for a Medicaid beneficiary for whom enrollment is verified monthly, there may not be more than a 1-month gap in coverage during each year of continuous enrollment. The patient must be enrolled as of December 31 of the measurement year.			
	For physician assessment from data that comes from a non-health plan: Any enrollment, claim or encounter transaction any time during the measurement year.			
	Event/diagnosis Event. Discharged alive for AMI, CABG or PCI on or between January 1 and November 1 of the year prior to the measurement year. Use the codes listed in Table IVD-A to identify AMI, PCI and CABG. AMI and CABG cases should be from inpatient claims only. All cases of PCI should be included, regardless of setting (e.g.,			

0000 150	Hennic V	asculai U	isease (IV	D). USE UI Z	ispiriii 0i	anome	er antithrombo	AIC	
inpatien	t, outpatie	ent, ED).							
_		• .	•						during both the same across both y
At least	st one ac	ute inpatie	nt visit (Ta		vith an IV	D diagno	(Table IVD-B), osis (Table IVI ocludes:		
 Angina Coron Cardio Occlus Athero Athero Arteria Athero Athero Note: Us 	ary ather ary artery ovascular sion or sto osclerosis osclerosis ic total od al embolisr	osclerosis / occlusion disease enosis of p s of renal a s of native ecclusion of sm and thr m. logs, patie	orecerebra ortery arteries of artery of t ombosis	the extremit he extremition	ies es		arotid and verte ominator, ther		teries) e medical record to
Table IV	D-A: Coo	des to Ider	ntify AMI, F	CI and CAB	G				
Descript	ion	CPT	HCPCS	ICD-9-CM	Diagnos	is	ICD-9-CM F	rocedu	ıre
AMI (inp	atient on	ly)		410.x1					
CABG (i	npatient (36.1, 30	• .	33510-3	33514, 33510	6-33519,	33521-3	33523, 33533-	33536	S2205-S2209
PCI	92980	, 92982, 92	2995	G0290		00.66, 3	36.06, 36.07		
Table IV	D-B: Coo	des to Ider	ntify IVD						
Descript	ion	ICD-9-(CM Diagno	osis					
IVD			-		9.2. 433	. 434. 44	0.1, 440.2, 44	0.4. 444	4. 445
	·	, ,	ntify Visit T	, ,	, , , , , , , , , , , , , , , , , , ,	, . 1	, · · · · · · · · · · · · · · · · · ·	,	,
			·	•					
Descript	ION	CPT	UB Rev	enue					
	9387, 99		7, 99401-9						345, 99347-99350, 051x, 0520-0523,

	0068 Ischemic vascular disease (IVD): Use of aspirin or another antithrombotic				
	Acute inpatient 99221-99223, 99231-99233, 99238, 99239, 99251-99255, 99261-99263, 99291 010x, 0110-0114, 0119, 0120-0124, 0129, 0130-0134, 0139, 0140-0144, 0149, 0150-0154, 0159, 016x, 020x-021x, 072x, 0987Medical record text Coronary artery disease				
	Stable angina				
	Lower extremity arterial disease/peripheral artery disease				
	Ischemia				
	Stroke				
	Artheroembolism				
	Renal artery atherosclerosis				
Exclusions	None				
Exclusion Details	None				
Risk Adjustment	No risk adjustment necessary				
Stratification	None				
Type Score	Rate/proportion better quality = higher score				
Algorithm	NA NA				

	0071 Acute myocardial infarction (AMI): Persistence of beta-blocker treatment after a heart attack				
Steward	National Committee for Quality Assurance, 1100 13th Street NW, Suite 1000, Washington, DC 20005				
Description	The percentage of patients age 18 years and older during the measurement year who were hospitalized and discharged alive July 1 of the year prior to the measurement year through June 30 of the measurement year with a diagnosis of acute myocardial infarction (AMI) and who received persistent beta-blocker treatment for six months after discharge.				
Туре	Process				
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Paper medical record/flow-sheet, Pharmacy data				
Level	Clinicians: Group, Clinicians: Individual, Health Plan				
Setting	All settings, Ambulatory Care: Clinic				
Numerator Statement	A 180-day course of treatment with beta-blockers post discharge.				
Numerator Details	Time Window: Six months after discharge from a hospital with AMI (with the discharge anywhere from July 1 of the year prior to the measurement year through June 30 of the measurement year).				
	Identify all patients in the denominator population whose dispensed days supply is ≥135 days in the 180 days following discharge. Persistence of treatment for this measure is defined as at least 75 percent of the days supply filled.				
	To determine continuity of treatment during the 180-day period, sum the number of allowed gap days to the number of treatment days for a maximum of 180 days (i.e., 135 treatment days + 45 gap days = 180 days); identify all prescriptions filled within 180 days of the Discharge Date.				
	To account for members who are on beta-blockers prior to admission, the organization should factor those prescriptions into adherence rates if the actual treatment days fall within the 180 days following discharge.				
	Table PBH-B Beta Blocker Medications: Noncardioselective beta-blockers (carteolol, carvedilol, labetalol, nadolol, penbutolol, pindolol, propranolol, timolol, sotalol), cardioselective beta-blockers (acebutolol, atenolol, betaxolol, bisoprolol, metoprolol, nebivolol), Antihypertensive combinations (atenolol-chlorthalidone, bendroflumethiazide-nadolol, bisoprolol-hydrochlorothiazide, hydrochlorothiazide-propranolol, hydrochlorothiazide-metoprolol, hydrochlorothiazide-timolol).				
Denominator Statement	Patients 18 years and older as of December 31 of the measurement year discharged alive from an acute inpatient setting with an AMI from July 1 of the year prior to the measurement year through June 30 of the measurement year.				

	0071 Acute myocardial infarction (AMI): Persistence of beta-blocker treatment after a heart attack				
Denominator Categories	Female; Male 18 years and older				
Denominator Details	Time Window: July 1 of the year prior to the measurement year through June 30 of the measurement year.				
	Patients 18 years and older as of December 31 of the measurement year discharged alive from an acute inpatient setting with an AMI from July 1 of the year prior to the measurement year through June 30 of the measurement year. If using health plan data, patient should have continuous medical and pharmacy benefit enrollment on the discharge date through 180 days after discharge, with no more than one gap in enrollment of up to 45 days within 180 days of the event. If the patient is a Medicaid beneficiary, the patient may not have more than 1 month gap in coverage and must be enrolled on the discharge date. If using non-health plan data, the patient must have a pharmacy claim or prescription written July 1 of the year prior to the measurement year through 180 days post-discharge to be included.				
	If a patient has more than one episode of AMI from July 1 of the year prior to the measurement year through June 30 of the measurement year, only the first discharge should be included.				
	Transfers to acute facilities: include hospitalizations in which the patient was transferred directly to another acute inpatient facility for any diagnosis. Count the discharge from the subsequent acute inpatient facility, not the inital discharge. The discharge date from the facility to which the patient was transferred must occur on or before June 30 of the measurement year.				
	Readmissions: If the patient was readmitted to an acute or nonacute care facility for any diagnosis, include the patient in the denominator and use the discharge date from the original hospitalization.				
	Description ICD-9-CM Diagnosis				
	AMI 410.x1				
Exclusions	Exclude patients who are identified as having a contraindication to beta-blocker therapy or previous adverse reaction to beta-blocker therapy. Also exclude from the denominator hospitalizations in which the patient was transferred directly to a nonacute care facility for any diagnosis.				
Exclusion Details	Exclude patients who are identified as having a contraindication to beta-blocker therapy or previous adverse reaction to beta-blocker therapy. Look as far back as possible in the patients' history through either administrative data or medical record review for evidence of contraindication or a previous adverse reaction to beta-blocker therapy.				
	Also exclude from the denominator hospitalizations in which the patient was transferred directly to a nonacute care facility for any diagnosis.				
	Table PBH-C: ICD-9 codes to identify exclusions: history of asthma: 493; hypotension: 458; heart block >1 degree: 426.0, 426.12, 426.13, 426.2-426.4, 426.51-426.54, 426.7; sinus bradycardia: 427.81; COPD: 491.2, 496, 506.4				

	0071 Acute myocardial infarction (AMI): Persistence of beta-blocker treatment after a heart attack
	Table PBH-D Medications to Identify Exclusions (hx of asthma): Bronchodilator combinations (budesonide-formoterol, fluticasone-salmeterol), inhaled corticosteroids (beclomethasone, budesonide, flunisolide, fluticasone, mometasone, triamcinolone, fluticasone CFC free).
Risk Adjustment	No risk adjustment necessary NA
Stratification	None
Type Score	Rate/proportion better quality = higher score
Algorithm	NA

	0073 IVD: Blood pressure management
Steward	National Committee for Quality Assurance, 1100 13th Street NW, Suite 1000, Washington, DC 20005
Description	The percentage of patients 18 years of age and older who were discharged alive with acute myocardial infarction (AMI), coronary artery bypass graft (CABG) or percutaneous coronary interventions (PCI) from January 1-November 1 of the year prior to the measurement year, or who had a diagnosis of ischemic vascular disease (IVD) during the measurement year and the year prior to the measurement year and who had BP reported as under control <140/90.
Туре	Outcome
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Paper medical record/flow-sheet
Level	Clinicians: Group, Clinicians: Individual
Setting	All settings, Ambulatory Care: Clinic
Numerator Statement	The numerator is the number of patients in the denominator whose most recent blood pressure is adequately controlled during the measurement year. For a patient's BP to be controlled, both the systolic and the diastolic BP must meet the desired threshold of <140/90 mm Hg.
Numerator Details	Time Window: 12 months
	The numerator is the number of patients in the denominator whose most recent blood pressure is adequately controlled during the measurement year. For a patient's BP to be controlled, both the systolic and the diastolic BP must meet the desired threshold of <140/90 mm Hg.
	Electronic Specification:
	If using electronic data to identify the most recent BP reading during the measurement year, calculate a numerator using the CPT Category II codes in Table IVD-G to determine compliance with the threshold. If CPT Category II codes are used to identify numerator compliance for this indicator, search for all codes in Table IVD-G and use the most recent code to evaluate whether the patient is numerator compliant. If a combination of data from internal electronic databases and CPT Category II codes is being used, search all sources and use the most recent result.
	If there are multiple BPs on the same date of service, use the lowest systolic and lowest diastolic BP on that date as the representative BP.
	The patient is noncompliant in the following circumstances.
	 The electronic result for the most recent BP test exceeds the desired threshold. The BP test result is missing. A BP test was not done during the measurement year.

	0073 IVD: Blood pressure managem	nent			
	Do not include readings that meet the	following criteria:			
	 Taken during an acute inpatient stay or an ED visit. Taken during an outpatient visit which was for the sole purpose of having a diagnostic test or surgical procedure performed. Taken the same day as major diagnostic or surgical procedure. Reported by or taken by the patient. Documentation of "VS within normal limits" or "vital signs normal". Medical Record Specification: 				
	To identify the representative blood pro	essure, follow these steps:			
	 Identify the most recent blood pressure reading noted during the measurement year. Do not include reading that meet the criteria as listed above under the electronic specification (i.e taken during an ED visit, etc.). Identify the lowest systolic and lowest diastolic reading from the most recent blood pressure notation in the medical record. If there are multiple readings for a single date, use the lowest systolic and the lowest diastoreading on that date as the representative blood pressure. The systolic and diastolic results do not need to from the same reading. Table IVD-G: Codes to Identify Systolic and Diastolic BP Levels 				
	Description	CPT Category II			
	Systolic pressure <140mm Hg	3076F			
	Systolic pressure ≥140 mm Hg	3077F			
	Diastolic pressure <80 mm Hg	3078F			
	Diastolic pressure 80-89 mm Hg	3079F			
	Diastolic pressure ≥ 90 mm Hg	3080F			
Denominator Statement	CABG or PCI on or between January	nber 31 of the measurement year who were discharged alive for AMI, 1 and November 1 of the year prior to the measurement year or who had a surement year and the year prior to the measurement year.			
Denominator Categories	Female; Male 18 years and older				
Denominator Details	Time Window: Between January 1st of measurement year.	of the year prior to the measurement year through December 31st of the			
	Patients 18 years or older as of Decen criteria:	nber 31 of the measurement year who met the following patient inclusion			
		from health plan data: Continuous medical benefit enrollment for the an one gap in continuous enrollment of up to 45 days during the			

0073 IVD: Blood pressure management

measurement year. To determine continuous enrollment for a Medicaid beneficiary for whom enrollment is verified monthly, there may not be more than a 1-month gap in coverage during each year of continuous enrollment. The patient must be enrolled as of December 31 of the measurement year.

• For calculating physician performance from non-health plan data. Any enrollment, claim or encounter transaction any time during the measurement year.

Event/ Diagnosis Event:

Discharged alive for AMI, CABG or PCI on or between January 1 and November 1 of the year prior to the measurement year. Use the codes listed in Table IVD-A to identify AMI, PCI and CABG. AMI and CABG cases should be from inpatient claims only. All cases of PCI should be included, regardless of setting (e.g., inpatient, outpatient, ED).

Diagnosis. Identify patients as having IVD who met at least one of the two criteria below, during both the measurement year and the year prior to the measurement year. Criteria need not be the same across both years.

• At least one outpatient visit (Table IVD-C) with an IVD diagnosis (Table IVD-B), or

• At least one acute inpatient visit (Table IVD-C) with an IVD diagnosis (Table IVD-B). Table IVD-A: Codes to Identify AMI,PCI, and CABG

Description CPT HCPCS ICD-9-CM Diagnosis ICD-9-CM Procedure

AMI (inpatient only) 410.x1

CABG (inpatient only) 33510-33514,

33516-33519,

33521-33523,

33533-33536 S2205-S2209 36.1, 36.2

PCI 92980, 92982, 92995 00.66, 36.06, 36.07

Table IVD-B: Codes to Identify IVD

Description ICD-9-CM Diagnosis

IVD 411, 413, 414.0, 414.2, 414.8, 414.9, 429.2, 433-434,

440.1, 440.2, 440.4, 444, 445

Table IVD-C: Codes to Identify Visit Type

Description CPT UB Revenue

Outpatient 99201-99205, 99211-99215, 99217-99220, 99241-99245, 99341-99345, 99347-99350, 99384-99387, 99394-99397, 99401-99404, 99411, 99412, 99420, 99429, 99455, 99456 051x, 0520-0523, 0526-0529, 057x-059x, 0982, 0983

Acute inpatient 99221-99223, 99231-99233, 99238, 99239, 99251-99255, 99261-99263, 99291

	0073 IVD: Blood pressure management
	010x, 0110-0114, 0119, 0120-0124, 0129, 0130-0134, 0139, 0140-0144, 0149, 0150-0154, 0159, 016x, 020x-021x, 072x, 0987
	Medical record data Documentation of IVD in the medical record includes:
	 IVD Ischemic heart disease Angina Coronary atherosclerosis Coronary artery occlusion Cardiovascular disease Occlusion or stenosis of precerebral arteries (including basilar, carotid and vertebral arteries) Atherosclerosis of renal artery Atherosclerosis of native arteries of the extremities Chronic total occlusion of artery of the extremities Arterial embolism and thrombosis Atheroembolism. Note: Use paper logs, patient registries or EMRs to identify the denominator, then use the medical record to confirm patient eligibility.
Exclusions	All patients with ESRD, who are pregnant or who had an admission to a non-acute inpatient setting during the measurement year.
Exclusion Details	 All patients with ESRD (Table CBP-C) on or prior to 12/31 of the measurement year. Documentation in the medical record must include a date noted indicating ESRD, dialysis or renal transplant meets the criterion for evidence of ESRD. All patients who are pregnant (Table CBP-C) during the measurement year. All patients who had an admission to a non-acute inpatient setting (Table FUH-B) any time during the measurement year. Table CBP-C Codes to Identify ESRD & Pregnancy Exclusions: Evidence of ESRD: CPT (36145, 36147, 36800, 36810, 36815, 36818, 36819, 36820, 36821, 36831-36833, 50300, 50320, , 50340, 50360, 50365, 50370, 50380, 90920, 90921, 90924, 90925, 90935, 90937, 90940, 90945, 90947, 90957-90962, 90965, 90966, 90969, 90970, 90989, 90993, 90997, 90999, 99512), HCPCS (G0257, G0308-G0319, G0322, G0323, G0326, G0327, G0392, G0393, S9339), ICD-9 diagnosis (585.5, 585.6, V42.0, V45.1, V56), ICD-9 Procedure (38.95, 39.27, 39.42, 39.43, 39.53, 39.93-39.95, 54.98, 55.6), UB Revenue (0367, 080x, 082x-085x, 088x), UB Type of Bill (72X), POS (65) Pregnancy: ICD-9 Diagnosis (630-679, V22, V23, V28) Table FUH-B to identify non-acute inpatient exclusions:
	Hospice: UB Rev (0115, 0125, 0135, 0145, 0155, 0650, 0656, 0658, 0659), UB Type Bill (81x, 82x), POS (34)
	SNF: UB Rev (019x), UB Type Bill (21x, 22x, 28x), POS (31, 32)
	Hospital Transitional Care: UB Type Bill (18x)
L	

	0073 IVD: Blood pressure management
	Rehabilitation: UB Rev (0118, 0128, 0138, 0148, 0158)
	Respite:UB Rev (0655)
	Intermediate Care Facility: POS (54)
	Residential Substance Abuse Treatment Facility: UB Rev (1002), POS (55)
	Psychiatric Residential Treatment Facility Center: HCPCS (T2048, H0017-19), UB Rev (1001), POS (56)
	Comprehensive Inpatient Rehabilitation Facility: POS (61)
Risk Adjustment	No risk adjustment necessary
Stratification	NA
Type Score	Rate/proportion better quality = higher score
Algorithm	NA

	0074 Chronic stable coronary artery disease: Lipid control
Steward	American Medical Association, 515 N. State St., Chicago, IL 60654
Description	Percentage of patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period who have a LDL-C result <100 mg/dL OR patients who have a LDL-C result ≥100 mg/dL and have a documented plan of care to achieve LDL-C <100mg/dL, including at a minimum the prescription of a statin.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Registry data URL www.pinnacleregistry.org Attachment PCPI_CAD-2_LipidControl NQF 0074.pdf
Level	Clinicians: Group, Clinicians: Individual
Setting	Ambulatory Care: Clinic, Ambulatory Care: Hospital Outpatient, Ambulatory Care: Office, Assisted Living, Group homes, Home, Nursing home (NH)/Skilled Nursing Facility (SNF)
Numerator Statement	Patients who have a LDL-C result <100 mg/dL OR Patients who have a LDL-C result ≥100 mg/dL and have a documented plan of care1 to achieve LDL-C <100 mg/dL, including at a minimum the prescription of a statin within a 12 month period. Definitions: *Documented plan of care may also include: documentation of discussion of lifestyle modifications (diet,
	exercise); scheduled re-assessment of LDL-C. *Prescribed may include prescription given to the patient for a statin at one or more visits in the measurement period OR patient already taking a statin as documented in current medication list. Numerator Instructions: The first numerator option can be reported for patients who have a documented LDL-C < 100 mg/dL at any time during the measurement period.
Numerator Details	Time Window:
	See attached for EHR Specifications. For Claims/Administrative: Report CPT II Code Patients who have LDL-C <100 mg/dL 3048F Most recent LDL-C <100 mg/dL OR

	0074 Chronic stable coronary artery disease: Lipid control
	Patients who have LDL-C =100 mg/dL and have a documented plan of care to achieve LDL-C <100 mg/dL,
	including prescription of lipid-lowering therapy
	3049F Most recent LDL-C 100-129 mg/dL OR
	3050F Most recent LDL-C greater than or equal to 130 mg/dL AND
	05XXF (code in development) Lipid lowering therapy plan of care documented AND
	• 4002F Statin therapy prescribed.
Denominator Statement	All patients aged 18 years and older with a diagnosis of coronary artery disease seen within a 12 month period.
Denominator Categories	Female; Male Aged 18 years and older
Denominator Details	Time Window: 12 consecutive months
	See attached for EHR Specifications.
	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, CPT).
Exclusions	Documentation of medical reason(s) for not prescribing a statin (e.g., allergy, intolerance to statin medication(s), other medical reasons).
	Documentation of patient reason(s) for not prescribing a statin (e.g., patient declined, other patient reasons).
	Documentation of system reason(s) for not prescribing a statin (e.g., financial reasons, other system reasons).
Exclusion	See attached for EHR Specifications.
Details	For Claims/Administrative:
	Documentation of medical reason(s) for not prescribing a statin (e.g., allergy, intolerance to statin medication(s), other medical reasons)
	 Append modifier to CPT II code 4XXXF-1P (in development). Documentation of patient reason(s) for not prescribing a statin (e.g., patient declined, other patient reasons)
	 Append modifier to CPT II code 4XXXF-2P (in development). Documentation of system reason(s) for not a statin (e.g., financial reasons, other system reasons)
	Append modifier to CPT II code 4XXXF-3P (in development).

	0074 Chronic stable coronary artery disease: Lipid control
Risk	No risk adjustment necessary
Adjustment	
Stratification	
Type Score	Rate/proportion better quality = higher score
Algorithm	See attached for calculation algorithm.

	0075 IVD: Complete lipid profile and Idl control <100
Steward	National Committee for Quality Assurance, 1100 13th Street NW, Suite 1000, Washington, DC 20005
Description	The percentage of patients 18 years of age and older who were discharged alive for acute myocardial infarction (AMI), coronary artery bypass graft (CABG) or percutaneous coronary interventions (PCI) from January 1-November 1 of the year prior to the measurement year, or who had a diagnosis of ischemic vascular disease (IVD) during the measurement year and the year prior to measurement year, who had each of the following during the measurement year. • Complete Lipid Profile
Туре	LDL-C control <100 mg/dL Outcome
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Lab data, Paper medical record/flow-sheet
Level	Clinicians: Group, Clinicians: Individual
Setting	All settings, Ambulatory Care: Clinic
Numerator	A complete lipid profile performed during the measurement year. A LDL-C control result of <100 mg/dL using the
Statement	most recent LDL-C screening test during the measurement year.
Numerator Details	Time Window: 12 months
	Electronic Specification:
	Complete Lipid Profile: A complete lipid profile performed during the measurement year (Table IVD-F) as identified by claim/encounter or electronic laboratory data.
	LDL-C Control: <100 mg/dL
	Use electronic laboratory data during the measurement year. Calculate a numerator by using the most recent LDL-C screening test. Use the CPT Category II codes in Table CMC-E to determine compliance. The patient is non compliant if: the electronic results for the most recent LDL-C test exceed the desired threshold, the electronic result for the most recent LDL-C test is missing or an LDL-C test was not done during the measurement year.
	Medical Record Specification:
	Complete Lipid Profile: A full lipid profile completed during the measurement year, with the date and result of each component of the profile documented. Identify the most recent visit of the doctor's office or clinic where a full lipid profile was documented and which occurred during the measurement year (but after the diagnosis of IVD was made). Each component of the lipid profile must be noted with the date of the test and results.
	LDL Control <100: The number of patients in the denominator whose LDL-C is adequately controlled during the measurement year. Use the most recent LDL-C level performed during the measurement year. At a minimum

	0075 IVD: Comple	ete lipid p	rofile and Idl control <100	
	documentation in t	he record	must include a note indicating the date when the test was performed and the result.	
	Table IVD-F: Codes to Identify a Complete Lipid Profile			
	Description	CPT	CPT Category II	
	Lipid panel	80061	3011F	
	OR			
	Description	CPT	LOINC	
	Total cholesterol	82465	2093-3, 14647-2	
	WITH			
	High density lipopr	otein (HD	L) 83701 2085-9, 14646-4, 18263-4	
	AND			
	Triglycerides	84478	2571-8, 12951-0, 14927-8, 47210-0	
	Table CMC-E: CP	Γ category	/ II codes to identify LDL-C levels	
	LDL-C <100: 3048	F		
	LDL-C 100-129: 30)49F		
	LDL-C ≥130: 3050	F		
Denominator Statement	AMI, CABG or PCI	on or bet	older as of December 31st of the measurement year who were discharged alive for ween January 1 and November 1 of the year prior to the measurement year or who g both the measurement year and the year prior to the measurement year.	
Denominator Categories	Female; Male 18 y	ears and	older	
Denominator Details	Time Window: Be measurement year		nuary 1 of the year prior to the measurement year and December 31st of the	
	Patients 18 years of criteria:	or older as	s of December 31 of the measurement year who met the following patient inclusion	
	measurement year measurement year	, with no i . To deter	mance generated from a health plan: Continuous medical benefit enrollment for the more than one gap in continuous enrollment of up to 45 days during the rmine continuous enrollment for a Medicaid beneficiary for whom enrollment is not be more than a 1-month gap in coverage during each year of continuous	

0075 IVD: Complete lipid profile and Idl control <100

enrollment. The patient must be enrolled as of December 31 of the measurement year.

For data on physician performance generated from non-health plan data: Any enrollment, claim or encounter transaction any time during the measurement year.

Event/ diagnosis: Event. Discharged alive for AMI, CABG or PCI on or between January 1 and November 1 of the year prior to the measurement year. Use the codes listed in Table IVD-A to identify AMI, PCI and CABG. AMI and CABG cases should be from inpatient claims only. All cases of PCI should be included, regardless of setting (e.g., inpatient, outpatient, ED).

Diagnosis. Identify patients as having IVD who met at least one of the two criteria below, during both the measurement year and the year prior to the measurement year. Criteria need not be the same across both years.

- At least one outpatient visit (Table IVD-C) with an IVD diagnosis (Table IVD-B), or
- At least one acute inpatient visit (Table IVD-C) with an IVD diagnosis (Table IVD-B).
 Medical record data
 Documentation of IVD in the medical record includes:
- IVD
- Ischemic heart disease
- Angina
- · Coronary atherosclerosis
- Coronary artery occlusion
- Cardiovascular disease
- Occlusion or stenosis of precerebral arteries (including basilar, carotid and vertebral arteries)
- Atherosclerosis of renal artery
- Atherosclerosis of native arteries of the extremities
- Chronic total occlusion of artery of the extremities
- Arterial embolism and thrombosis
- Atheroembolism.

Note: Use paper logs, patient registries or EMRs to identify the denominator, then use the medical record to confirm patient eligibility.

Exclusions None.

Table IVD-A: Codes to Identify AMI, PCI and CABG

Description CPT HCPCS ICD-9-CM Diagnosis ICD-9-CM Procedure

AMI (inpatient only) 410.x1

CABG (inpatient only) 33510-33514, 33516-33519, 33521-33523, 33533-33536 S2205-S2209

36.1, 36.2

PCI 92980, 92982, 92995 G0290 00.66, 36.06, 36.07

Table IVD-B: Codes to Identify IVD

Description ICD-9-CM Diagnosis

	0075 IVD: Complete lipid profile and Idl control <100
	IVD 411, 413, 414.0, 414.2, 414.8, 414.9, 429.2, 433, 434, 440.1, 440.2, 440.4, 444, 445
	Source: Table CMC-B in Cholesterol Management for Patients With Cardiovascular Conditions.
	Table IVD-C: Codes to Identify Visit Type
	Description CPT UB Revenue
	Outpatient 99201-99205, 99211-99215, 99217-99220, 99241-99245, 99341-99345, 99347-99350, 99384-99387, 99394-99397, 99401-99404, 99411, 99412, 99420, 99429, 99455, 99456 051x, 0520-0523, 0526-0529, 057x-059x, 0982, 0983
	Acute inpatient 99221-99223, 99231-99233, 99238, 99239, 99251-99255, 99261-99263, 99291 010x, 0110-0114, 0119, 0120-0124, 0129, 0130-0134, 0139, 0140-0144, 0149, 0150-0154, 0159, 016x, 020x-021x, 072x, 0987
Exclusions	None
Exclusion Details	None
Risk	No risk adjustment necessary
Adjustment	NA
Stratification	NA
Type Score	Rate/proportion better quality = higher score
Algorithm	NA NA

	0076 Optimal vascular care		
Steward	MN Community Measurement, 3433 Broadway Street NE, Suite 455, Minneapolis MN 55413		
Description	Percentage of adult patients ages 18 to 75 who have ischemic vascular disease with optimally managed modifiable risk factors (LDL, blood pressure, tobacco-free status, daily aspirin use).		
Туре	Outcome		
Data Source	Electronic Clinical Data : Electronic Health Record, Electronic Clinical Data : Registry, Paper Records		
	URL www.mncm.org/site/?p=resources URL www.mncm.org/site/?p=resources		
Level	Clinician: Group/Practice Clinic site location		
Setting	Ambulatory Care: Clinic/Urgent Care, Ambulatory Care: Clinician Office		
Numerator Statement	Patients ages 18 to 75 with ischemic vascular disease (IVD) who meet all of the following targets from the most recent visit during the measurement period: LDL less than 100, Blood Pressure less than 140/90, Tobacco-Free Status, Daily Aspirin Use (unless contraindicated).		
	Please note: On 7/27/2010, the blood pressure component of this measure was changed for patients with a comorbidity of diabetes (target less than 140/90). MNCM's technical advisory group recommended this changed based on ACCORD results, ICSI's most recent guideline changes (July 2010), and the national meaningful use measures for diabetes blood pressure control. A target of less than 140/90 allows for individualization of patient goals.		
	On March 9, 2011, the measurement and reporting committee reviewed recent ICSI guideline changes for blood pressure targets for stable coronary artery disease and hypertension and additionally considered the request of the NQF cardiovascular committee and decided to change the blood pressure target to < 140/90 for all IVD patients.		
	Values are collected as the most recent during the measurement period (January 1 through December 31), with the exception of the LDL value which is collected over a 15 month time span to allow a greater window of time for patients that may not complete a cholesterol test within the 12 month time frame, but do complete a cholesterol test within 15 months (October 1 of the previous year through December 31 of the measurement year).		
Numerator Details	Time Window:		
	Numerator for the LDL Component:		
	LDL Date [Date (mm/dd/yyyy)] AND		
	LDL Value [Numeric]		
	Numerator calculation: numerator compliant is LDL during the last 15 months AND LDL value is less than 100.		
	I		

0076 Optimal vascular care
Enter the date of the most recent LDL test prior to and including 12/31/YYYY (measurement period).
Enter the value of the most recent LDL test prior to and including 12/31/ YYYY (measurement period).
Numerator for the Blood Pressure Component:
Blood Pressure Date [Date (mm/dd/yyyy)] AND
BP Systolic [Numeric] AND
BP Diastolic [Numeric]
Numerator calculation: numerator compliant is BP during the measurement period AND the following targets: Systolic <140 AND Diastolic <90.
Enter the date of the most recent Blood Pressure (BP) test prior to and including 12/31/YYYY (measurement period).
Numerator for the Tobacco Component:
Tobacco Status Documentation Date [Date (mm/dd/yyyy)] AND
Tobacco Status [Numeric]
1 = Tobacco Free (patient does not use tobacco) 2 = No Documentation 3 = Current Tobacco User
Numerator calculation: Numerator compliant is Value 1 = Tobacco Free AND valid date
Enter the most recent date (prior to and including 12/31/YYYY (measurement period) that the patient's tobacco status was documented.
Numerator for the Aspirin Component:
Aspirin Use or Documented Contraindication for the use of aspirin.
Aspirin (ASA) Date [Date (mm/dd/yyyy)]
Enter the most recent date of documented ASA or anti-platelet prior to and including 12/31/YYYY (measurement period).
FYI: any documented date in the measurement period of ASA or an anti-platelet is acceptable; the date does not need to be the most recent.
The following are accepted ASA or anti-platelet medications:
 Aspirin (ASA) Plavix (clopidogrel) Ticlid (ticlopidine) Pravigard (aspirin/pravastatin) Aggrenox (aspirin/dypyridamole) Low dose enteric-coated 81 mg ASA (Ecotrin or Bayer)

	0076 Optimal vascular care
	OR
	Aspirin (ASA) Contraindication Date [Date (mm/dd/yyyy)].
	If patient has a documented contraindication to ASA, enter the date of the contraindication. Any valid contraindication date will be given credit. Auditor must be able to validate this date.
	Accepted contraindications:
	 Anticoagulant use, Lovenox (Enoxaparin) or Coumadin (Warfarin) Any history of gastrointestinal (GI)* or intracranial bleed (ICB)
	• Allergy to ASA. *Gastroesophogeal reflux disease (GERD) is not automatically considered a contraindication but may be included if specifically documented as a contraindication by the physician.
	The following may be exclusions if specifically documented by the physician:
	 Use of non-steroidal anti-inflammatory agents Documented risk for drug interaction Uncontrolled hypertension defined as >180 systolic, >110 diastolic
Denominator	 Other provider documented reason for not being on ASA therapy. Patients ages 18 to 75 with ischemic vascular disease who have at least two visits for this condition over the last
Statement	two years (established patient) with at least one visit in the last 12 months.
Denominator Categories	Female; Male Ages 18 to 75 during the measurement period
Denominator Details	Time Window:
	Birth date [Date (mm/dd/yyyy)]
	Ischemic vascular disease ICD-9 codes:
	410-410.92 Acute Myocardial Infarction (AMI)
	411-411.89 Post Myocardial Infarction Syndrome
	412 Old AMI
	413-413.9 Angina Pectoris
	414.0-414.07 Coronary Arthrosclerosis
	414.2 Chronic Total Occlusion of Coronary Artery
	414.8 Other Chronic Ischemic Heart Disease (IHD)

	0076 Optimal vascular care
	414.3 Atherosclerosis due to lipid rich plaque
	414.9 Chronic IHD
	429.2 Cardiovascular (CV) disease, unspecified
	433-433.91 Occlusion and stenosis of pre-cerebral arteries
	434-434.91 Occlusion of cerebral arteries
	440.1 Atherosclerosis of renal artery
	440.2-440.29 Atherosclerosis of native arteries of the extremities, unspecified
	440.4 Chronic Total Occlusion of Artery of the Extremities
	444-444.9 Arterial embolism and thrombosis
	445-445.8 Atheroembolism.
Exclusions	Valid exclusions include patients who only had one coded visit to the clinic during the last two years, patients who had died during the measurement period, patients who were in hospice during the measurement period, patients who were permanent nursing home residents during the measurement period, or patients who were coded with IVD in error.
Exclusion	Patient was a permanent nursing home resident home during the measurement period.
Details	Patient was in hospice at any time during the measurement period.
	Patient died prior to the end of the measurement period.
	Documentation that diagnosis was coded in error.
Risk	Case-mix adjustment
Adjustment	Attachment MNCM Case Mix Risk Adjustment June 2010-634242034150216836.docx
Stratification	
Type Score	Weighted score/composite/scale better quality = higher score
Algorithm	

	0079 Heart failure: Left ventricular ejection fraction assessment (outpatient setting)
Steward	American Medical Association, 515 N State St., Chicago, IL 60654
Description	Percentage of patients aged 18 years and older with a diagnosis of heart failure for whom the quantitative or qualitative results of a recent or prior (any time in the past) LVEF assessment is documented within a 12 month period.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Paper medical record/flow-sheet, Registry data
	URL www.pinnacleregistry.org Attachment NQF 0079_PCPI_HF-1_LVEF Assessment.pdf
Level	Clinicians: Group, Clinicians: Individual
Setting	Ambulatory Care: Clinic, Ambulatory Care: Hospital Outpatient, Ambulatory Care: Office, Assisted Living, Group homes, Home, Nursing home (NH)/Skilled Nursing Facility (SNF)
Numerator	Patients for whom the quantitative or qualitative results of a recent or prior (any time in the past) LVEF
Statement	assessment is documented* within a 12 month period.
	*Documentation must include documentation in a progress note of the results of an LVEF assessment, regardless of when the evaluation of ejection fraction was performed.
	Qualitative results correspond to numeric equivalents as follows:
	Hyperdynamic: corresponds to LVEF greater than 70%
	Normal: corresponds to LVEF 50% to 70% (midpoint 60%)
	Mild dysfunction: corresponds to LVEF 40% to 49% (midpoint 45%)
	Moderate dysfunction: corresponds to LVEF 30% to 39% (midpoint 35%)
	Severe dysfunction: corresponds to LVEF less than 30%.
Numerator Details	Time Window: Once during the measurement period.
	See attached for EHR Specifications.
	For Claims/Administrative: Report CPT Category II Code 3021F- Left ventricular ejection fraction (LVEF) < 40% or documentation of moderately or severely depressed left ventricular systolic dysfunction
	OR
	CPT Category II Code 3022F- Left ventricular ejection fraction (LVEF) ≥ 40% or documentation as normal

	0079 Heart failure: Left ventricular ejection fraction assessment (outpatient setting)
	function or mildly depressed left ventricular systolic function.
Denominator Statement	All patients aged 18 years and older with a diagnosis of heart failure.
Denominator Categories	Female; Male 18 years of age and older
Denominator Details	Time Window: 12 consecutive months
	See attached for EHR Specifications.
	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, SNOMED, CPT).
Exclusions	None
Exclusion Details	
Risk Adjustment	No risk adjustment necessary
Stratification	
Type Score	Rate/proportion better quality = higher score
Algorithm	See attached for calculation algorithm.

	0081 Heart failure: Angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB)
	therapy for left ventricular systolic dysfunction
Steward	American Medical Association, 515 N State St., Chicago, IL 60654
Description	Percentage of patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40% who were prescribed ACE inhibitor or ARB therapy either within a 12 month period when seen in the outpatient setting or at hospital discharge.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Paper medical record/flow-sheet, Registry data
	URL www.pinnacleregistry.org Attachment NQF 0081_PCPI_HF-7_ACE ARB for LVSD.pdf
Level	Clinicians: Group, Clinicians: Individual
Setting	Ambulatory Care: Clinic, Ambulatory Care: Hospital Outpatient, Ambulatory Care: Office, Assisted Living, Group homes, Home, Hospital, Nursing home (NH) /Skilled Nursing Facility (SNF)
Numerator	Patients who were prescribed* ACE inhibitor or ARB therapy either within a 12 month period when seen in the
Statement	outpatient setting or at hospital discharge.
	*Prescribed may include prescription given to the patient for ACE inhibitor or ARB therapy at one or more visits in the measurement period OR patient already taking ACE inhibitor or ARB therapy as documented in current medication list.
Numerator Details	Time Window: Once during the measurement period (outpatient/nursing home) OR at each hospital discharge.
	See attached for EHR Specifications.
	For Claims/Administrative: Report CPT Category II Code 4009F- Angiotensin converting enzyme (ACE) inhibitor or Angiotensin Receptor Blocker (ARB) therapy prescribed.
	All patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40%.
Statement	LVEF < 40% corresponds to qualitative documentation of moderate dysfunction or severe dysfunction.
Denominator Categories	Female; Male 18 years of age and older
Denominator Details	Time Window: 12 consecutive months
	Note: For the inpatient setting (CPT 99239, 99239), the diagnosis refers to the principal discharge diagnosis. The

	0081 Heart failure: Angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) therapy for left ventricular systolic dysfunction
	principal diagnosis is typically the first listed on the inpatient claim form with secondary or attributed diagnoses to follow in descending order of importance.
	ICD-9-CM Diagnosis Code:
	Note: Although this measure is limited to patients with left ventricular systolic dysfunction, diastolic ICD-9-CM codes are included to provide invariability in coding among measures.
	See attached for EHR Specifications.
	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, CPT)
	AND
	Report CPT Category II Code (in development)
	3021F- Left ventricular ejection fraction (LVEF) < 40% or qualitative documentation of moderate dysfunction or severe dysfunction.
Exclusions	Documentation of medical reason(s) for not prescribing ACE inhibitor or ARB therapy; Append modifier to CPT II code 4009F-1P.
	Documentation of patient reason(s) for not prescribing ACE inhibitor or ARB; Append modifier to CPT II code 4009F-2P.
	Documentation of system reason(s) for not prescribing ACE inhibitor or ARB; Append modifier to CPT II code 4009F-3P.
Exclusion	See attached for EHR specifications.
Details	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, SNOMED, CPT).
Risk Adjustment	No risk adjustment necessary
Stratification	
Type Score	Rate/proportion better quality = higher score
Algorithm	See attached for calculation algorithm.

	0083 Heart failure: Beta-blocker therapy for left ventricular systolic dysfunction
Steward	American Medical Association, 515 N State St., Chicago, IL 60654
Description	Percentage of patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40% who were prescribed beta-blocker therapy either within a 12 month period when seen in the outpatient setting or at hospital discharge.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Clinical Data, Electronic Health/Medical Record, Paper medical record/flow-sheet, Registry data
	URL www.pinnacleregistry.org Attachment NQF 0083_PCPI_HF-6_Beta Blocker for LVSD.pdf
Level	Clinicians: Group, Clinicians: Individual
Setting	Ambulatory Care: Clinic, Ambulatory Care: Hospital Outpatient, Ambulatory Care: Office, Assisted Living, Group homes, Home, Hospital, Nursing home (NH)/Skilled Nursing Facility (SNF)
Numerator Statement	Patients who were prescribed* beta-blocker therapy** either within a 12 month period when seen in the outpatient setting or at hospital discharge.
	*Prescribed may include prescription given to the patient for beta-blocker therapy at one or more visits in the measurement period OR patient already taking beta-blocker therapy as documented in current medication list.
	**Beta-blocker therapy should include bisoprolol, carvedilol, or sustained release metoprolol succinate.
Numerator Details	Time Window: Once during the measurement period.
	See attached for EHR Specifications.
	For Claims/Administrative: Report CPT Category II Code: 4006F- Beta-blocker therapy prescribed.
Denominator Statement	All patients aged 18 years and older with a diagnosis of heart failure with a current or prior LVEF < 40%.
Statement	LVEF < 40% corresponds to qualitative documentation of moderate dysfunction or severe dysfunction.
Denominator Categories	Female; Male 18 years and older
Denominator Details	Time Window: 12 consecutive months
	See attached for EHR Specifications.

	0083 Heart failure: Beta-blocker therapy for left ventricular systolic dysfunction
	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, SNOMED, CPT) AND
	Report CPT Category II Code (in development)3021F- Left ventricular ejection fraction (LVEF) < 40% or documentation of moderately or severely depressed left ventricular systolic function.
Exclusions	Documentation of medical reason(s) for not prescribing beta-blocker therapy.
	Documentation of patient reason(s) for not prescribing beta-blocker therapy.
	Documentation of system reason(s) for not prescribing beta-blocker therapy.
Exclusion Details	See attached for EHR Specifications.
Details	For Claims/Administrative: See coding tables attached for coding (ICD-9-CM, ICD-10-CM, SNOMED, CPT)
	Append modifier to CPT II code 4006F-1P
	 Append modifier to CPT II code 4006F-2P Append modifier to CPT II code 4006F-3P.
Risk Adjustment	No risk adjustment necessary
Stratification	
Type Score	Rate/proportion better quality = higher score
Algorithm	See attached for calculation algorithm

	0132 Aspirin at arrival for acute myocardial infarction (AMI)
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of acute myocardial infarction (AMI) patients who received aspirin within 24 hours before or after hospital arrival.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet
	URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1135 267770141 URL Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228
	760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator	AMI patients who received aspirin within 24 hours before or after hospital arrival.
Statement	
Numerator Details	Time Window: 24 hours before hospital arrival through 24 hours after hospital arrival.
	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-77 through 1-78. Appendices Appendix C - Medication Tables – pages Appendix C-3 through Appendix C-6. Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-1-1 through AMI-1-5.
Denominator Statement	AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91).
Denominator Categories	Female; Male Greater than or equal to 18 years old
Denominator Details	Time Window: From hospital arrival to time of hospital discharge.
	ICD-9-CM Principal Diagnosis codes:

	0132 Aspirin at arrival for acute myocardial infarction (AMI)
	410.00: Anterolateral wall, acute myocardial infarction-episode of care unspecified
	410.01: Anterolateral wall, acute myocardial infarction-initial episode
	410.10: Other anterior wall, acute myocardial infarction-episode of care unspecified
	410.11: Other anterior wall, acute myocardial infarction-initial episode
	410.20: Inferolateral wall, acute myocardial infarction-episode of care unspecified
	410.21: Inferolateral wall, acute myocardial infarction-initial episode
	410.30: Inferoposterior wall, acute myocardial infarction-episode of care unspecified
	410.31: Inferoposterior wall, acute myocardial infarction-initial episode
	410.40: Other inferior wall, acute myocardial infarction-episode of care unspecified
	410.41: Other inferior wall, acute myocardial infarction-initial episode
	410.50: Other lateral wall, acute myocardial infarction-episode of care unspecified
	410.51: Other lateral wall, acute myocardial infarction-initial episode
	410.60: True posterior wall, acute myocardial infarction-episode of care unspecified
	410.61: True posterior wall, acute myocardial infarction-initial episode
	410.70: Subendocardial, acute myocardial infarction-episode of care unspecified
	410.71: Subendocardial, acute myocardial infarction-initial episode
	410.80: Other specified sites, acute myocardial infarction-episode of care unspecified
	410.81: Other specified sites, acute myocardial infarction-initial episode
	410.90: Unspecified site, acute myocardial infarction-episode of care unspecified
	410.91: Unspecified site, acute myocardial infarction-initial episode
Exclusions	Exclusions:
	 <18 years of age Patients who have a length of stay greater than 120 days Patients enrolled in clinical trials Discharged to another hospital on day of or day after arrival Discharged on day of arrival Expired on day of or day after arrival Left against medical advice on day of or day after arrival Patients with comfort measures only documented on day of or day after arrival

	0132 Aspirin at arrival for acute myocardial infarction (AMI)
	Patients with a documented reason for no aspirin on arrival.
Exclusion	Refer to
Details	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	• Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-69 through 1-71, 1-90, 1-98 through 1-104, 1-117, 1-118 through 1-120, 1-204, and 1-324 through 1-326.
	• Appendices Appendix C - Medication Tables PDF – pages Appendix C-3 through Appendix C-6 plus Appendix C-9, and Appendix H - Miscellaneous Tables – pages Appendix H-5.
	• Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-1-1 through AMI-1-5.
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	Refer to
3	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228
	760129036: Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) - pages AMI-
	5 plus AMI-1-1 through AMI-1-5.

	0133 PCI mortality (risk-adjusted)©
Steward	American College of Cardiology, 2400 N Street, NW, Washington, DC 20037
Description	Risk adjusted PCI mortality rate.
Туре	Outcome
Data Source	Registry data
	URL http://www.ncdr.com/WebNCDR/ELEMENTS.ASPX URL http://www.ncdr.com/WebNCDR/ELEMENTS.ASPX
Level	Facility/Agency
Setting	Hospital
Numerator Statement	Patients 18 years of age and older with a PCI procedure performed during admission who expired.
Numerator Details	Time Window: One year
	PCI=yes
	Coding instructions: indicate if the patient had a percutaneous coronary intervention (PCI)
	Selections: yes/no
	Supporting definitions: PCI: A percutaneous coronary intervention (PCI) is the placement of an angioplasty guide wire, balloon, or other device (e.g., stent, atherectomy, brachytherapy, or thrombectomy catheter) into a native coronary artery or coronary bypass graft for the purpose of mechanical coronary revascularization. Source: NCDR
	Discharge status = deceased
	Selections: Alive/deceased
	Coding instructions: Indicate whether the patient was alive or deceased at discharge.
Denominator Statement	Patients 18 years of age and older with a PCI procedure performed during admission.
Denominator Categories	Female; Male > 18 years of age
Denominator	Time Window: One year (quarterly to include previous four quarters of data)

	0133 PCI mortality (risk-adjusted)©
Details	
	PCI=yes
	Coding instructions: indicate if the patient had a percutaneous coronary intervention (PCI)
	Selections: yes/no
	Supporting definitions: PCI: A percutaneous coronary intervention (PCI) is the placement of an angioplasty guide wire, balloon, or other device (e.g., stent, atherectomy, brachytherapy, or thrombectomy catheter) into a native coronary artery or coronary bypass graft for the purpose of mechanical coronary revascularization. Source: NCDR
	Age: patients must be 18 years of age to be included in the registry.
Exclusions	NCDR Registry patients who did not have a PCI (Patient admissions with a diagnostic cath only during that admission);
	2. Data submissions that do not pass the data quality and completeness reports;
	3. Procedure variables for subsequent PCIs during the same admission (if the patient had more than one PCI procedure during that admission);
	Patient admissions with PCI who transferred to another facility on discharge;
	5. Patient admissions with PCI who have more than two variables in the risk model that are missing.
Exclusion Details	1. PCI = yes
Details	2. All data submissions must pass the data quality and completeness reports to be included. Note: If one or two variables are missing, the value is imputed for certain characteristics (see appendix 2 of the NCDR CathPCI Registry PCI Risk Adjusted Morality Model 2008 for more information). If the value is missing for more than two variables, the patient record is excluded. However, in our data quality program, all variables in the risk model have a high "inclusion" criteria. This means that, when a hospital submits data to us, they need to have a high level of completeness (around 99%) for those variables. If they are not able to meet the criteria in our data quality program, they do not receive risk adjusted mortality for the records they submitted for that quarter.
	3. PCI = yes for more than one procedure during the same admission.
	Discharge location = transferred to another facility
Risk	Risk-adjustment devised specifically for this measure/condition
Adjustment	Attachment Contemporary Mortality Risk Prediction for PCI (2).pdf
Stratification	N/A

	0133 PCI mortality (risk-adjusted)©
Type Score	Weighted score/composite/scale better quality = lower score
Algorithm	

	0135 Evaluation of left ventricular systolic function (LVS)
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of heart failure patients with documentation in the hospital record that left ventricular systolic (LVS) function was evaluated before arrival, during hospitalization, or is planned for after discharge.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet
	URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1135 267770141 URL Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228
	760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator	HF patients with documentation in the hospital record that LVS function was evaluated before arrival, during
Statement	hospitalization, or is planned for after discharge.
Numerator Details	Time Window: From hospital arrival to time of hospital discharge.
	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-254 through 1-256. Section 2 - Measurement Information Section 2.2 – Heart Failure (HF) – pages HF-2-1 through HF-2-5.
Denominator Statement	HF patients (ICD-9-CM principal diagnosis of HF: 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, 428.0, 428.1, 428.20, 428.21, 428.22, 428.23, 428.30, 428.31, 428.32, 428.33, 428.40, 428.41, 428.42, 428.43, 428.9).
Denominator Categories	Female; Male Greater than or equal to 18 years old
Denominator Details	Time Window: From hospital arrival to time of hospital discharge.
	ICD-9-CM Principal Diagnosis codes:
	402.01: Hypertensive heart disease, malignant, with heart failure
<u> </u>	I

0135 Evaluation of left ventricular systolic function (L\	/S)
402.11: Hypertensive heart disease, benign, with heart fail	ure
402.91: Hypertensive heart disease, unspecified, with hear	rt failure
404.01: Hypertensive heart and chronic kidney disease, m disease stage I through stage IV, or unspecified	alignant, with heart failure and with chronic kidney
404.03: Hypertensive heart and chronic kidney disease, m disease stage V or end stage renal disease	alignant, with heart failure and with chronic kidney
404.11: Hypertensive heart and chronic kidney disease, be stage I through stage IV, or unspecified	enign, with heart failure and with chronic kidney disease
404.13: Hypertensive heart and chronic kidney disease, be stage V or end stage renal disease	enign, with heart failure and chronic kidney disease
404.91: Hypertensive heart and chronic kidney disease, ur disease stage I through stage IV, or unspecified	nspecified, with heart failure and with chronic kidney
404.93: Hypertensive heart and chronic kidney disease, ur stage V or end stage renal disease	nspecified, with heart failure and chronic kidney disease
428.0: Congestive heart failure, unspecified	
428.1: Left heart failure	
428.20: Unspecified systolic heart failure	
428.21: Acute systolic heart failure	
428.22: Chronic systolic heart failure	
428.23: Acute on chronic systolic heart failure	
428.30: Unspecified diastolic heart failure	
428.31: Acute diastolic heart failure	
428.32: Chronic diastolic heart failure	
428.33: Acute on chronic diastolic heart failure	
428.40: Unspecified combined systolic and diastolic heart	failure
428.41: Acute combined systolic and diastolic heart failure	
428.42: hronic combined systolic and diastolic heart failure	e
428.43: Acute on chronic combined systolic and diastolic h	neart failure

	0135 Evaluation of left ventricular systolic function (LVS)
	428.9: Heart failure, unspecified.
Exclusions	Exclusions:
	 <18 years of age Patients who have a length of stay greater than 120 days Discharged to another hospital Expired Left against medical advice Discharged to home for hospice care Discharged to a health care facility for hospice care Patients enrolled in clinical trials Patients with comfort measures only documented Reasons for no LVS function evaluation documented by a physician, advanced practice nurse, or physician assistant
	• Patients who had a left ventricular assistive device (LVAD) or heart transplant procedure during hospital stay (ICD-9-CM procedure code of LVAD or Heart Transplant: 33.6, 37.51, 37.52, 37.53, 37.54, 37.60, 37.62, 37.63, 37.65, 37.66, 37.68).
Exclusion	Refer to
Details	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: • Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-90, 1-98 through 1-117 through 1-120, 1-201, 1-204 through 1-205, and 1-254 through 1-256. • Section 2 - Measurement Information Section 2.2 – Heart Failure (HF) – pages HF-5 plus HF-2-1 through HF-
	2-5
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 2 - Measurement Information Section 2.2 – Heart Failure (HF) – pages HF-5 plus HF-2-4 through HF-2-5.

	0137 ACEI or ARB for left ventricular systolic dysfunction—acute myocardial infarction (AMI) patients
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of acute myocardial infarction (AMI) patients with left ventricular systolic dysfunction (LVSD) who are prescribed an ACEI or ARB at hospital discharge. For purposes of this measure, LVSD is defined as chart documentation of a left ventricular ejection fraction (LVEF) less than 40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet URL
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=113 5267770141 URL Refer to
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=122 8760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator Statement	AMI patients who are prescribed an ACEI or ARB at hospital discharge.
Numerator Details	Time Window: From hospital arrival to time of hospital discharge.
	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=122 8760129036:
	• Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-18 through 1-19 plus pages 1-67 through 1-68.
	 Appendices Appendix C - Medication Tables – pages Appendix C-6 through Appendix C-7 plus pages Appendix C-11 through Appendix C-12. Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-3-1 through AMI-3-6.
Denominator Statement	AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41,
	410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.90, 410.91); with chart documentation of a left ventricular ejection fraction (LVEF) < 40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction.
Denominator	Female; Male Greater than or equal to 18 years old

	0137 ACEI or ARB for left ventricular systolic dysfunction—acute myocardial infarction (AMI) patients
Categories	
Denominator Details	Time Window: From hospital arrival to time of hospital discharge
	ICD-9-CM Principal Diagnosis codes:
	410.00: Anterolateral wall, acute myocardial infarction-episode of care unspecified
	410.01: Anterolateral wall, acute myocardial infarction-initial episode
	410.10: Other anterior wall, acute myocardial infarction-episode of care unspecified
	410.11: Other anterior wall, acute myocardial infarction-initial episode
	410.20: Inferolateral wall, acute myocardial infarction-episode of care unspecified
	410.21: Inferolateral wall, acute myocardial infarction-initial episode
	410.30: Inferoposterior wall, acute myocardial infarction-episode of care unspecified
	410.31: Inferoposterior wall, acute myocardial infarction-initial episode
	410.40: Other inferior wall, acute myocardial infarction-episode of care unspecified
	410.41: Other inferior wall, acute myocardial infarction-initial episode
	410.50: Other lateral wall, acute myocardial infarction-episode of care unspecified
	410.51: Other lateral wall, acute myocardial infarction-initial episode
	410.60: True posterior wall, acute myocardial infarction-episode of care unspecified
	410.61: True posterior wall, acute myocardial infarction-initial episode
	410.70: Subendocardial, acute myocardial infarction-episode of care unspecified
	410.71: Subendocardial, acute myocardial infarction-initial episode
	410.80: Other specified sites, acute myocardial infarction-episode of care unspecified
	410.81: Other specified sites, acute myocardial infarction-initial episode
	410.90: Unspecified site, acute myocardial infarction-episode of care unspecified
	410.91: Unspecified site, acute myocardial infarction-initial episode.
	LVSD - Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=122

	0137 ACEI or ARB for left ventricular systolic dysfunction—acute myocardial infarction (AMI) patients
	8760129036:
	Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-257 through 1-260.
Exclusions	Exclusions:
	• <18 years of age
	Patients who have a length of stay greater than 120 days
	Discharged to another hospital
	• Expired
	Left against medical advice
	Discharged to home for hospice care Discharged to a hospital care facility for hospical care.
	 Discharged to a health care facility for hospice care Patients with comfort measures only documented
	Patients with comfort measures only documented Patients enrolled in clinical trials
	Patients with a documented reason for no ACEI and no ARB at discharge.
Exclusion	Refer to
Details	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=122
Details	8760129036:
	• Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-90, 1-98 through 1-104 1 117 through 1 120 1 204 1 257 through 1 260 and 1 215 through 1 220
	104, 1-117 through 1-120, 1-204, 1-257 through 1-260, and 1-315 through 1-320. • Appendices Appendix C - Medication Tables PDF – pages Appendix C-6 through Appendix C-7 plus pages
	Appendix C-11 through Appendix C-12, and Appendix H - Miscellaneous Tables – page Appendix H-5.
	• Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus
	AMI-3-1 through AMI-3-6.
Risk	No risk adjustment necessary
Adjustment	
	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	Refer to
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=122
	8760129036: Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages
	AMI-5 plus AMI-3-1 through AMI-3-6.

	0142 Aspirin prescribed at discharge for AMI
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of acute myocardial infarction (AMI) patients who are prescribed aspirin at hospital discharge.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet
	URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1135 267770141 URL Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator Statement	AMI patients who are prescribed aspirin at hospital discharge
Numerator Details	Time Window: From hospital arrival to time of hospital discharge.
	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-75 through 1-76. Appendices Appendix C - Medication Tables – pages Appendix C-3 through Appendix C-6. Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-2-1 through AMI-2-5.
Denominator Statement	AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91).
Denominator Categories	Female; Male Greater than or equal to 18 years old
Denominator Details	Time Window: From hospital arrival to time of hospital discharge.
	ICD-9-CM Principal Diagnosis codes:

	0142 Aspirin prescribed at discharge for AMI
	410.00: Anterolateral wall, acute myocardial infarction-episode of care unspecified
	410.01: Anterolateral wall, acute myocardial infarction-initial episode
	410.10: Other anterior wall, acute myocardial infarction-episode of care unspecified
	410.11: Other anterior wall, acute myocardial infarction-initial episode
	410.20: Inferolateral wall, acute myocardial infarction-episode of care unspecified
	410.21: Inferolateral wall, acute myocardial infarction-initial episode
	410.30: Inferoposterior wall, acute myocardial infarction-episode of care unspecified
	410.31: Inferoposterior wall, acute myocardial infarction-initial episode
	410.40: Other inferior wall, acute myocardial infarction-episode of care unspecified
	410.41: Other inferior wall, acute myocardial infarction-initial episode
	410.50: Other lateral wall, acute myocardial infarction-episode of care unspecified
	410.51: Other lateral wall, acute myocardial infarction-initial episode
	410.60: True posterior wall, acute myocardial infarction-episode of care unspecified
	410.61: True posterior wall, acute myocardial infarction-initial episode
	410.70: Subendocardial, acute myocardial infarction-episode of care unspecified
	410.71: Subendocardial, acute myocardial infarction-initial episode
	410.80: Other specified sites, acute myocardial infarction-episode of care unspecified
	410.81: Other specified sites, acute myocardial infarction-initial episode
	410.90: Unspecified site, acute myocardial infarction-episode of care unspecified
	410.91: Unspecified site, acute myocardial infarction-initial episode.
Exclusions	Exclusions:
	 <18 years of age Patients who have a length of stay greater than 120 days Patients enrolled in clinical trials Discharged to another hospital Expired Left against medical advice Discharged to home for hospice care Discharged to a health care facility for hospice care

	0142 Aspirin prescribed at discharge for AMI
	 Patients with comfort measures only documented Patients with a documented reason for no aspirin at discharge.
Exclusion	Refer to
Details	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	• Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-69 through 1-71, 1-90, 1-98 through 1-104, 1-117, 1-118 through 1-120, 1-204, and 1-321 through 1-323.
	• Appendices Appendix C - Medication Tables PDF – pages Appendix C-3 through Appendix C-6 plus Appendix C-9, and Appendix H - Miscellaneous Tables – page Appendix H-5.
	• Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-2-1 through AMI-2-5.
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-2-1 through AMI-2-5.

	0160 Beta-blocker prescribed at discharge for AMI
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of acute myocardial infarction (AMI) patients who are prescribed a beta-blocker at hospital discharge.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet
	URL
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1135 267770141 URL Refer to
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator	AMI patients who are prescribed a beta-blocker at hospital discharge.
Statement	
Numerator	Time Window: From hospital arrival to time of hospital discharge.
Details	
	 Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-88 through 1-89. Appendices Appendix C - Medication Tables – pages Appendix C-7 through Appendix C-9.
	 Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5-1 through AMI-5-5.
Denominator	AMI patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal
Statement	diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91)
	Female; Male Greater than or equal to 18 years old
Categories	
Denominator Details	Time Window: From hospital arrival to time of hospital discharge.
	ICD-9-CM Principal Diagnosis codes:
	410.00: Anterolateral wall, acute myocardial infarction-episode of care unspecified

	0160 Beta-blocker prescribed at discharge for AMI
	410.01: Anterolateral wall, acute myocardial infarction-initial episode
	410.10: Other anterior wall, acute myocardial infarction-episode of care unspecified
	410.11: Other anterior wall, acute myocardial infarction-initial episode
	410.20: Inferolateral wall, acute myocardial infarction-episode of care unspecified
	410.21: Inferolateral wall, acute myocardial infarction-initial episode
	410.30: Inferoposterior wall, acute myocardial infarction-episode of care unspecified
	410.31: Inferoposterior wall, acute myocardial infarction-initial episode
	410.40: Other inferior wall, acute myocardial infarction-episode of care unspecified
	410.41: Other inferior wall, acute myocardial infarction-initial episode
	410.50: Other lateral wall, acute myocardial infarction-episode of care unspecified
	410.51: Other lateral wall, acute myocardial infarction-initial episode
	410.60: True posterior wall, acute myocardial infarction-episode of care unspecified
	410.61: True posterior wall, acute myocardial infarction-initial episode
	410.70: Subendocardial, acute myocardial infarction-episode of care unspecified
	410.71: Subendocardial, acute myocardial infarction-initial episode
	410.80: Other specified sites, acute myocardial infarction-episode of care unspecified
	410.81: Other specified sites, acute myocardial infarction-initial episode
	410.90: Unspecified site, acute myocardial infarction-episode of care unspecified
	410.91: Unspecified site, acute myocardial infarction-initial episode
Exclusions	Exclusions:
	 <18 years of age Patients who have a length of stay greater than 120 days Patients enrolled in clinical trials Discharged to another hospital Expired Left against medical advice Discharged to home for hospice care Discharged to a health care facility for hospice care Patients with comfort measures only documented Patients with a documented reason for no beta-blocker at discharge.

	0160 Beta-blocker prescribed at discharge for AMI
Exclusion Details	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-90, 1-98 through 1-104, 1-117, 1-118 through 1-120, 1-204, and 1-327 through 1-330. Appendices Appendix C - Medication Tables PDF – pages Appendix C-7 through Appendix C-9, and Appendix H - Miscellaneous Tables – page Appendix H-5. Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-5-1 through AMI-5-5.
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-5-1 through AMI-5-5.

	0162 ACEI or ARB for left ventricular systolic dysfunction—heart failure (HF) patients
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of heart failure (HF) patients with left ventricular systolic dysfunction (LVSD) who are prescribed an ACEI or ARB at hospital discharge. For purposes of this measure, LVSD is defined as chart documentation of a left ventricular ejection fraction (LVEF) less than 40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet URL
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1135 267770141 URL Refer to
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator Statement	HF patients who are prescribed an ACEI or ARB at hospital discharge.
Numerator Details	Time Window: From hospital arrival to time of hospital discharge
	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-18 through 1-19 plus pages 1-67 through 1-68. Appendices Appendix C - Medication Tables – pages Appendix C-6 through Appendix C-7 plus pages
	Appendix C-11 through Appendix C-12. Section 2 - Measurement Information Section 2.2 – Heart Failure (HF) – pages HF-3-1 through HF-3-5.
Denominator Statement	HF patients (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of HF: 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, 428.0, 428.1, 428.20, 428.21, 428.22, 428.23, 428.30, 428.31, 428.32, 428.33, 428.40, 428.41, 428.42, 428.43, 428.9); with chart documentation of a left ventricular ejection fraction (LVEF) < 40% or a narrative description of left ventricular systolic (LVS) function consistent with moderate or severe systolic dysfunction.
Denominator Categories	Female; Male Greater than or equal to 18 years old

	0162 ACEI or ARB for left ventricular systolic dysfunction—heart failure (HF) patients
Denominator Details	Time Window: From hospital arrival to time of hospital discharge.
	ICD-9-CM Principal Diagnosis codes:
	402.01: Hypertensive heart disease, malignant, with heart failure
	402.11: Hypertensive heart disease, benign, with heart failure
	402.91: Hypertensive heart disease, unspecified, with heart failure
	404.01: Hypertensive heart and chronic kidney disease, malignant, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.03: Hypertensive heart and chronic kidney disease, malignant, with heart failure and with chronic kidney disease stage V or end stage renal disease
	404.11: Hypertensive heart and chronic kidney disease, benign, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.13: Hypertensive heart and chronic kidney disease, benign, with heart failure and chronic kidney disease stage V or end stage renal disease
	404.91: Hypertensive heart and chronic kidney disease, unspecified, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.93: Hypertensive heart and chronic kidney disease, unspecified, with heart failure and chronic kidney disease stage V or end stage renal disease
	428.0: Congestive heart failure, unspecified
	428.1: Left heart failure
	428.20: Unspecified systolic heart failure
	428.21: Acute systolic heart failure
	428.22: Chronic systolic heart failure
	428.23: Acute on chronic systolic heart failure
	428.30: Unspecified diastolic heart failure
	428.31: Acute diastolic heart failure
	428.32: Chronic diastolic heart failure
	428.33: Acute on chronic diastolic heart failure

	0162 ACEI or ARB for left ventricular systolic dysfunction—heart failure (HF) patients
	428.40: Unspecified combined systolic and diastolic heart failure
	428.41: Acute combined systolic and diastolic heart failure
	428.42: Chronic combined systolic and diastolic heart failure
	428.43: Acute on chronic combined systolic and diastolic heart failure
	428.9: Heart failure, unspecified
	LVSD - Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-257 through 1-260.
Exclusions	Exclusions:
	 Patients who had a left ventricular assistive device (LVAD) or heart transplant procedure during hospital stay (ICD-9-CM procedure code of LVAD or Heart Transplant: 33.6, 37.51, 37.52, 37.53, 37.54, 37.60, 37.62, 37.63, 37.65, 37.66, 37.68) <18 years of age Patients who have a length of stay greater than 120 days
	Discharged to another hospitalExpired
	Left against medical advice
	 Discharged to home for hospice care Discharged to a health care facility for hospice care
	Patients enrolled in clinical trials
	Patients with comfort measures only documented
Exclusion	Patients with a documented reason for no ACEI and no ARB at discharge. Refer to
Details	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-90, 1-98 through 1-104, 1-117 through 1-120, 1-201, 1-204 through 1-205, 1-257 through 1-260, and 1-315 through 1-320. Appendices Appendix C - Medication Tables PDF – pages Appendix C-6 through Appendix C-7 plus pages Appendix C-11 through Appendix C-12, and Appendix H - Miscellaneous Tables – page Appendix H-5. Section 2 - Measurement Information Section 2.2 – Heart Failure (HF) – pages HF-5 plus HF-3-1 through HF-3-5
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score

	0162 ACEI or ARB for left ventricular systolic dysfunction—heart failure (HF) patients
Algorithm	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 2 - Measurement Information Section 2.2 – Heart Failure (HF) – pages HF-5 plus HF-3-4 through HF-3-5.

	0163 Primary PCI received within 90 minutes of hospital arrival
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of acute myocardial infarction (AMI) patients with ST-segment elevation or LBBB on the ECG closest to arrival time receiving primary percutaneous coronary intervention (PCI) during the hospital stay with a time from hospital arrival to PCI of 90 minutes or less.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet
	URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1135 267770141 URL Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Lavel	
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator Statement	AMI patients whose time from hospital arrival to primary Percutaneous Coronary Intervention (PCI) is 90 minutes or less.
Numerator Details	Time Window: From hospital arrival through 90 minutes after hospital arrival.
	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-69 through 1-74 and 1-172 through 1-176. Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-8a-1 through AMI-8a-7.
Denominator Statement	Principal diagnosis of AMI (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91); and PCI procedure (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal or other procedure code for PCI: 00.66); and ST-segment elevation or LBBB on the ECG performed closest to hospital arrival; and PCI performed within 24 hours after hospital arrival.
Denominator Categories	Female; Male Greater than or equal to 18 years old
Denominator	Time Window: From hospital arrival through 24 hours after hospital arrival.

	0163 Primary PCI received within 90 minutes of hospital arrival
Details	
	ICD-9-CM Principal Diagnosis codes:
	410.00: Anterolateral wall, acute myocardial infarction-episode of care unspecified
	410.01: Anterolateral wall, acute myocardial infarction-initial episode
	410.10: Other anterior wall, acute myocardial infarction-episode of care unspecified
	410.11: Other anterior wall, acute myocardial infarction-initial episode
	410.20: Inferolateral wall, acute myocardial infarction-episode of care unspecified
	410.21: Inferolateral wall, acute myocardial infarction-initial episode
	410.30: Inferoposterior wall, acute myocardial infarction-episode of care unspecified
	410.31: Inferoposterior wall, acute myocardial infarction-initial episode
	410.40: Other inferior wall, acute myocardial infarction-episode of care unspecified
	410.41: Other inferior wall, acute myocardial infarction-initial episode
	410.50: Other lateral wall, acute myocardial infarction-episode of care unspecified
	410.51: Other lateral wall, acute myocardial infarction-initial episode
	410.60: True posterior wall, acute myocardial infarction-episode of care unspecified
	410.61: True posterior wall, acute myocardial infarction-initial episode
	410.70: Subendocardial, acute myocardial infarction-episode of care unspecified
	410.71: Subendocardial, acute myocardial infarction-initial episode
	410.80: Other specified sites, acute myocardial infarction-episode of care unspecified
	410.81: Other specified sites, acute myocardial infarction-initial episode
	410.90: Unspecified site, acute myocardial infarction-episode of care unspecified
	410.91: Unspecified site, acute myocardial infarction-initial episode
	ICD-9-CM Principal or Other Procedure code: 00.66: Percutaneous transluminal coronary angioplasty [PTCA] or coronary atherectomy
	First PCI Date, First PCI Time, and Initial ECG Interpretation—Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228

	0163 Primary PCI received within 90 minutes of hospital arrival
	760129036:
	• Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-172 through 1-176 and 1-228 through 1-231.
Exclusions	Exclusions:
	<18 years of agePatients who have a length of stay greater than 120 days
	Patients enrolled in clinical trials
	 Patients received as a transfer from an inpatient or outpatient department of another hospital Patients received as a transfer from the emergency/observation department of another hospital Patients received as a transfer from an ambulatory surgery center
	 Patient administered fibrinolytic agent prior to PCI PCI described as non-primary by physician, advanced practice nurse, or physician assistant Patients who did not receive PCI within 90 minutes and had a reason for delay documented by a physician, advanced practice nurse, or physician assistant (e.g., social, religious, initial concern or refusal, cardiopulmonary arrest, balloon pump insertion, respiratory failure requiring intubation).
Exclusion	Refer to
Details	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-69 through 1-74, 1-90, 1-98 through 1-100, 1-117, 1-166, 1-172 through 1-176, 1-201, 1-204 through 1-205, 1-228 through 1-231, 1-266 through 1-267, 1-310 through 1-312, and 1-392 through 1-393. Appendices Appendix C - Medication Tables PDF – page Appendix C-9. Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-8a-1 through AMI-8a-7.
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	Refer to
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-8a-1 through AMI-8a-7.

	0164 Fibrinolytic therapy received within 30 minutes of hospital arrival
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850
Description	Percentage of acute myocardial infarction (AMI) patients with ST-segment elevation or LBBB on the ECG closest to arrival time receiving fibrinolytic therapy during the hospital stay and having a time from hospital arrival to fibrinolysis of 30 minutes or less.
Туре	Process
Data Source	Electronic Health/Medical Record, Paper medical record/flow-sheet
	URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1135 267770141 URL Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 1 - Data Dictionary Alphabetical Data Dictionary.
Level	Facility/Agency, Population: National, Program: QIO
Setting	Hospital
Numerator Statement	AMI patients whose time from hospital arrival to fibrinolysis is 30 minutes or less.
Numerator Details	Time Window: From hospital arrival through 30 minutes after hospital arrival.
	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: • Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-69 through 1-74 and 1-167 through 1-170. • Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-7a-1
Denominator Statement	through AMI-7a-6. Principal diagnosis of AMI (International Classification of Diseases, 9th revision, Clinical Modification [ICD-9-CM] principal diagnosis code of AMI: 410.00, 410.01, 410.10, 410.11, 410.20, 410.21, 410.30, 410.31, 410.40, 410.41, 410.50, 410.51, 410.60, 410.61, 410.70, 410.71, 410.80, 410.81, 410.90, 410.91); and ST-segment elevation or LBBB on the ECG performed closest to hospital arrival; and fibrinolytic therapy within 6 hours after hospital arrival; and fibrinolytic therapy is primary reperfusion therapy.
Denominator Categories	Female; Male Greater than or equal to 18 years old
Denominator Details	Time Window: From hospital arrival through 6 hours after hospital arrival.

0164 Fibrinolytic therapy received within 30 minutes of hospital arrival
ICD-9-CM Principal Diagnosis codes:
410.00: Anterolateral wall, acute myocardial infarction-episode of care unspecified
410.01: Anterolateral wall, acute myocardial infarction-initial episode
410.10: Other anterior wall, acute myocardial infarction-episode of care unspecified
410.11: Other anterior wall, acute myocardial infarction-initial episode
410.20: Inferolateral wall, acute myocardial infarction-episode of care unspecified
410.21: Inferolateral wall, acute myocardial infarction-initial episode
410.30: Inferoposterior wall, acute myocardial infarction-episode of care unspecified
410.31: Inferoposterior wall, acute myocardial infarction-initial episode
410.40: Other inferior wall, acute myocardial infarction-episode of care unspecified
410.41: Other inferior wall, acute myocardial infarction-initial episode
410.50: Other lateral wall, acute myocardial infarction-episode of care unspecified
410.51: Other lateral wall, acute myocardial infarction-initial episode
410.60: True posterior wall, acute myocardial infarction-episode of care unspecified
410.61: True posterior wall, acute myocardial infarction-initial episode
410.70: Subendocardial, acute myocardial infarction-episode of care unspecified
410.71: Subendocardial, acute myocardial infarction-initial episode
410.80: Other specified sites, acute myocardial infarction-episode of care unspecified
410.81: Other specified sites, acute myocardial infarction-initial episode
410.90: Unspecified site, acute myocardial infarction-episode of care unspecified
410.91: Unspecified site, acute myocardial infarction-initial episode
Fibrinolytic Administration, Fibrinolytic Administration Date, Fibrinolytic Administration Time, and Initial ECG Interpretation - Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
• Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-166 through 1-170 and 1-228 through 1-

	0164 Fibrinolytic therapy received within 30 minutes of hospital arrival
	231.
Exclusions	Exclusions:
	 <18 years of age Patients who have a length of stay greater than 120 days Patients enrolled in clinical trials Patients received as a transfer from an inpatient or outpatient department of another hospital Patients received as a transfer from the emergency/observation department of another hospital Patients received as a transfer from an ambulatory surgery center Patients who did not receive fibrinolytic therapy within 30 minutes and had a reason for delay documented by a physician, advanced practice nurse, or physician assistant (e.g., social, religious, initial concern or refusal, cardiopulmonary arrest, balloon pump insertion, respiratory failure requiring intubation).
Exclusion	Refer to
Details	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036:
	 Section 1 - Data Dictionary Alphabetical Data Dictionary – pages 1-20 through 1-21, 1-69 through 1-74, 1-90, 1-98 through 1-100, 1-117, 1-166 through 1-170, 1-204, 1-228 through 1-231, 1-307 through 1-309, and 1-392 through 1-393. Appendices Appendix C - Medication Tables PDF – page Appendix C-9. Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-7a-1 through AMI-7a-6.
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	Refer to http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier4&cid=1228 760129036: Section 2 - Measurement Information Section 2.1 - Acute Myocardial Infarction (AMI) – pages AMI-5 plus AMI-
	7a-1 through AMI-7a-6.

	0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-02-01, Baltimore, MD 21244-9045
Description	The measure estimates a hospital-level risk-standardized mortality rate (RSMR), defined as death from any cause within 30 days after the index admission date, for patients discharged from the hospital with a principal diagnosis of HF.
Туре	Outcome
Data Source	Electronic administrative data/claims
	URL N/A www.qualitynet.org N/A URL Condition Category/ICD-9 Code Map available at: http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1182 785083979
Level	Facility/Agency
Setting	Hospital
Numerator Statement	This outcome measure does not have a traditional numerator and denominator like a core process measure (e.g., percentage of adult patients with diabetes aged 18-75 years receiving one or more hemoglobin A1c tests per year); thus, we are using this field to define the outcome.
	The outcome for this measure is 30-day all-cause mortality. We define mortality as death from any cause within 30 days of the index admission date for patients discharged from the hospital with a principal diagnosis of HF.
Numerator Details	Time Window: Patients who die within 30 days of the index admission date.
	Measure includes deaths from any cause within 30 days from admission date of index hospitalization.
Denominator Statement	Note: This outcome measure does not have a traditional numerator and denominator like a core process measure; thus, we are using this field to define the patient cohort and to define exclusions to the patient cohort.
	The cohort includes admissions for Medicare FFS beneficiaries age 65 years or older discharged from the hospital with a principal diagnosis of HF (ICD-9-CM codes 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, and 428.xx) and with a complete claims history for the 12 months prior to admission. Patients who are transferred from one acute care facility to another must have a principal discharge diagnosis of HF at both hospitals. The initial hospital for a transferred patient is designated as the responsible institution for the episode.
	If a patient has more than one HF admission in a year, one hospitalization is randomly selected for inclusion in the measure.

	0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization
Denominator Categories	Female; Male The target population is age 65 years or older
Denominator Details	Time Window: This measure was developed with 12 months of data. Currently the measure is publicly reported with three years of index hospitalizations.
	The denominator includes patients aged 65 and older admitted to non-federal acute care hospitals for an HF defined by a principal discharge diagnosis of (ICD-9-CM codes 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, and 428.xx) and with a complete claims history for the 12 months prior to admission.
	ICD-9-CM codes that define the patient cohort:
	402.01 Hypertensive heart disease, malignant, with heart failure
	402.11 Hypertensive heart disease, benign, with heart failure
	402.91 Hypertensive heart disease, unspecified, with heart failure
	404.01 Hypertensive heart and chronic kidney disease, malignant, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.03 Hypertensive heart and chronic kidney disease, malignant, with heart failure and with chronic kidney disease stage V or end stage renal disease
	404.11 Hypertensive heart and chronic kidney disease, benign, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.13 Hypertensive heart and chronic kidney disease, benign, with heart failure and chronic kidney disease stage V or end stage renal disease
	404.91 Hypertensive heart and chronic kidney disease, unspecified, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.93 Hypertensive heart and chronic kidney disease, unspecified, with heart failure and chronic kidney disease stage V or end stage renal disease
	428.0 Congestive heart failure, unspecified
	428.1 Left heart failure
	428.20 Unspecified systolic heart failure
	428.21 Acute systolic heart failure
	428.22 Chronic systolic heart failure

	0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization
	428.23 Acute on chronic systolic heart failure
	428.30 Unspecified diastolic heart failure
	428.31 Acute diastolic heart failure
	428.32 Chronic diastolic heart failure
	428.33 Acute on chronic diastolic heart failure
	428.40 Unspecified combined systolic and diastolic heart failure
	428.41 Acute combined systolic and diastolic heart failure
	428.42 Chronic combined systolic and diastolic heart failure
	428.43 Acute on chronic combined systolic and diastolic heart failure
	428.9 Heart Failure, unspecified.
Exclusions	The measures exclude admissions for patients:
Exclusion Details	 who were discharged on the day of admission or the following day and did not die or get transferred (because it is less likely they had a significant HF diagnosis); who were transferred from another acute care hospital (because the death is attributed to the hospital where the patient was initially admitted); with inconsistent or unknown mortality status or other unreliable data (e.g. date of death precedes admission date); enrolled in the Medicare Hospice program any time in the 12 months prior to the index hospitalization including the first day of the index admission (since it is likely these patients are continuing to seek comfort measures only); who were discharged alive and against medical advice (AMA) (because providers did not have the opportunity to deliver full care and prepare the patient for discharge); that were not the first hospitalization in the 30 days prior to a patient's death. We use this criteria to prevent attribution of a death to two admissions. See "Denominator Exclusions" section.
Risk Adjustment	Risk-adjustment devised specifically for this measure/condition URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1163 010421830
Stratification	Results of this measure will not be stratified.
Type Score	Rate/proportion better quality = lower score

	0229 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following heart failure (HF) hospitalization
Algorithm	

	0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization
Steward	Centers for Medicare & Medicaid Services (CMS), 7500 Security Boulevard, Mail Stop S3-02-01, Baltimore, MD 21244-9045
Description	The measure estimates a hospital-level risk-standardized mortality rate (RSMR), defined as death from any cause within 30 days after the index admission date, for patients discharged from the hospital with a principal diagnosis of AMI.
Туре	Outcome
Data Source	Electronic administrative data/claims URL Condition Category/ICD-9 Code Map available at: http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1182 785083979
Level	Facility/Agency
Setting	Hospital
Numerator Statement	This outcome measure does not have a traditional numerator and denominator like a core process measure (e.g., percentage of adult patients with diabetes aged 18-75 years receiving one or more hemoglobin A1c tests per year); thus, we are using this field to define the outcome.
	The outcome for this measure is 30-day all-cause mortality. We define mortality as death from any cause within 30 days of the index admission date for patients discharged from the hospital with a principal diagnosis of AMI.
Numerator Details	Time Window: Patients who die within 30 days of the index admission date.
	Measure includes deaths from any cause within 30 days from admission date of index hospitalization.
Denominator Statement	Note: This outcome measure does not have a traditional numerator and denominator like a core process measure; thus, we are using this field to define the patient cohort.
	The cohort includes admissions for Medicare FFS beneficiaries age 65 years or older discharged from the hospital with a principal diagnosis of AMI (ICD-9-CM codes 410.xx except for 410.x2) and with a complete claims history for the 12 months prior to admission. Patients who are transferred from one acute care facility to another must have a principal discharge diagnosis of AMI at both hospitals. The initial hospital for a transferred patient is designated as the responsible institution for the episode.
	If a patient has more than one AMI admission in a year, one hospitalization is randomly selected for inclusion in the measure.
Denominator	Female; Male The target population is age 65 years or older

	0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization
Categories	
Denominator Details	Time Window: This measure was developed with 12 months of data. Currently the measure is publicly reported with three years of index hospitalizations.
	The denominator includes patients aged 65 and older admitted to non-federal acute care hospitals for an AMI defined by a principal discharge diagnosis of ICD-9-CM code 410.xx, excluding those with 410.x2 (AMI, subsequent episode of care) and with a complete claims history for the 12 months prior to admission.
	ICD-9-CM codes that define the patient cohort:
	410.00 AMI (anterolateral wall) – episode of care unspecified
	410.01 AMI (anterolateral wall) – initial episode of care
	410.10 AMI (other anterior wall) – episode of care unspecified
	410.11 AMI (other anterior wall) – initial episode of care
	410.20 AMI (inferolateral wall) – episode of care unspecified
	410.21 AMI (inferolateral wall) – initial episode of care
	410.30 AMI (inferoposterior wall) – episode of care unspecified
	410.31 AMI (inferoposterior wall) – initial episode of care
	410.40 AMI (other inferior wall) – episode of care unspecified
	410.41 AMI (other inferior wall) – initial episode of care
	410.50 AMI (other lateral wall) – episode of care unspecified
	410.51 AMI (other lateral wall) – initial episode of care
	410.60 AMI (true posterior wall) – episode of care unspecified
	410.61 AMI (true posterior wall) – initial episode of care
	410.70 AMI (subendocardial) – episode of care unspecified
	410.71 AMI (subendocardial) – initial episode of care
	410.80 AMI (other specified site) – episode of care unspecified
	410.81 AMI (other specified site) – initial episode of care

	0230 Hospital 30-day, all-cause, risk-standardized mortality rate (RSMR) following acute myocardial infarction (AMI) hospitalization
	410.90 AMI (unspecified site) – episode of care unspecified
	410.91 AMI (unspecified site) – initial episode of care
	Note: We do not include 410.x2 (AMI, subsequent episode of care).
Exclusions	The measures exclude admissions for patients:
	• who were discharged on the day of admission or the following day and did not die or get transferred (because it is less likely they had a significant AMI);
	• who were transferred from another acute care hospital (because the death is attributed to the hospital where the patient was initially admitted);
	• with inconsistent or unknown mortality status or other unreliable data (e.g. date of death precedes admission date); enrolled in the Medicare Hospice program any time in the 12 months prior to the index hospitalization including the first day of the index admission (since it is likely these patients are continuing to seek comfort measures only);
	 who were discharged alive and against medical advice (AMA) (because providers did not have the opportunity to deliver full care and prepare the patient for discharge); that were not the first hospitalization in the 30 days prior to a patient's death. We use this criteria to prevent
	attribution of a death to two admissions.
Exclusion Details	See "Denominator Exclusions" section.
Risk	Risk-adjustment devised specifically for this measure/condition.
Adjustment	URL
	http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1163 010421830
Stratification	Results of this measure will not be stratified.
Type Score	Rate/proportion better quality = lower score
Algorithm	

	0277 Congestive heart failure admission rate (PQI 8)
Steward	Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, MD 20850
Description	Percent of county population with an admissions for CHF.
Туре	Outcome
Data Source	Electronic administrative data/claims
	URL http://www.qualityindicators.ahrq.gov/software.htm http://www.qualityindicators.ahrq.gov/downloads/winqi/AHRQ_QI_Windows_Software_Documentation_V41a.pdf
Level	Population: Counties or cities, Population: states
Setting	Ambulatory Care: Office
Numerator Statement	All discharges of age 18 years and older with ICD-9-CM principal diagnosis code for CHF.
Numerator Details	Time Window: Time period is user defined. Users of the measure typically use a 12 month time period.
	All discharges of age 18 years and older with ICD-9-CM principal diagnosis code for CHF.
	Include ICD-9-CM diagnosis codes:
	39891
	RHEUMATIC HEART FAILURE
	4280
	CONGESTIVE HEART FAILURE
	4281
	LEFT HEART FAILURE
	42820
	SYSTOLIC HRT FAILURE NOS OCT02-
	42821
	AC SYSTOLIC HRT FAILURE OCT02-
	42822

0277 Congestive heart failure admission rate (PQI 8)
CHR SYSTOLIC HRT FAILURE OCT02-
42823
AC ON CHR SYST HRT FAIL OCT02-
42830
DIASTOLC HRT FAILURE NOS OCT02-
42831
AC DIASTOLIC HRT FAILURE OCT02-
42832
CHR DIASTOLIC HRT FAIL OCT02-
42833
AC ON CHR DIAST HRT FAIL OCT02-
42840
SYST/DIAST HRT FAIL NOS OCT02-
42841
AC SYST/DIASTOL HRT FAIL OCT02-
42842
CHR SYST/DIASTL HRT FAIL OCT02-
42843
AC/CHR SYST/DIA HRT FAIL OCT02-
4289
HEART FAILURE NOS
Include ICD-9-CM diagnosis codes ONLY for discharges before 2002Q3 (ending September 30, 2002):
40201
MAL HYPERT HRT DIS W CHF
40211

0277 Congestive heart failure admission rate (PQI 8)
BENIGN HYP HRT DIS W CHF
40291
HYPERTEN HEART DIS W CHF
40401
MAL HYPER HRT/REN W CHF
40403
MAL HYP HRT/REN W CHF/RF
40411
BEN HYPER HRT/REN W CHF
40413
BEN HYP HRT/REN W CHF/RF
40491
HYPER HRT/REN NOS W CHF
40493
HYP HT/REN NOS W CHF/RF
Exclude cases:
 transfer from a hospital (different facility) transfer from a skilled Nursing Facility (SNF) or Intermediate Care Facility (ICF) transfer from another health care facility MDC 14 (pregnancy, childbirth, and puerperium) with a cardiac procedure code. ICD-9-CM Cardiac procedure codes:
0050
IMPL CRT PACEMAKER SYS OCT02-
0051
IMPL CRT DEFIBRILLAT OCT02-
0052

0277 Congestive heart failure admission rate (PQI 8)
IMP/REP LEAD LF VEN SYS OCT02-
0053
IMP/REP CRT PACEMKR GEN OCT02-
0054
IMP/REP CRT DEFIB GENAT OCT02-
0056
INS/REP IMPL SENSOR LEAD OCT06-
0057
IMP/REP SUBCUE CARD DEV OCT06-
0066
PTCA OCT06-
1751
IMPLANTATION OF RECHARGEABLE CARDIAC CONTRACTILITY MODULATION [CCM], TOTAL SYSTEM OCT09-
1752
IMPLANTATION OR REPLACEMENT OF CARDIAC CONTRACTILITY MODULATION [CCM] RECHARGEABLE PULSE GENERATOR ONLY OCT09-
3500
CLOSED VALVOTOMY NOS
3501
CLOSED AORTIC VALVOTOMY
3502
CLOSED MITRAL VALVOTOMY
3503
CLOSED PULMON VALVOTOMY
3504

0277 Congestive heart failure admission rate (PQI 8)
CLOSED TRICUSP VALVOTOMY
3510
OPEN VALVULOPLASTY NOS
3511
OPN AORTIC VALVULOPLASTY
3512
OPN MITRAL VALVULOPLASTY
3513
OPN PULMON VALVULOPLASTY
3514
OPN TRICUS VALVULOPLASTY
3520
REPLACE HEART VALVE NOS
3521
REPLACE AORT VALV-TISSUE
3522
REPLACE AORTIC VALVE NEC
3523
REPLACE MITR VALV-TISSUE
3524
REPLACE MITRAL VALVE NEC
3525
REPLACE PULM VALV-TISSUE
3526
REPLACE PULMON VALVE NEC

0277 Congestive heart failure admission rate (PQI 8)
PROS REPAIR ATRIA DEF-CL
3553
PROST REPAIR VENTRIC DEF
3554
PROS REP ENDOCAR CUSHION
3555
PROS REP VENTRC DEF-CLOS OCT06-
3560
GRFT REPAIR HRT SEPT NOS
3561
GRAFT REPAIR ATRIAL DEF
3562
GRAFT REPAIR VENTRIC DEF
3563
GRFT REP ENDOCAR CUSHION
3570
HEART SEPTA REPAIR NOS
3571
ATRIA SEPTA DEF REP NEC
3572
VENTR SEPTA DEF REP NEC
3573
ENDOCAR CUSHION REP NEC
3581
TOT REPAIR TETRAL FALLOT

0277 Congestive heart failure admission rate (PQI 8)
3582
TOTAL REPAIR OF TAPVC
3583
TOT REP TRUNCUS ARTERIOS
3584
TOT COR TRANSPOS GRT VES
3591
INTERAT VEN RETRN TRANSP
3592
CONDUIT RT VENT-PUL ART
3593
CONDUIT LEFT VENTR-AORTA
3594
CONDUIT ARTIUM-PULM ART
3595
HEART REPAIR REVISION
3596
PERC HEART VALVULOPLASTY
3598
OTHER HEART SEPTA OPS
3599
OTHER HEART VALVE OPS
3601
PTCA-1 VESSEL W/O AGENT
3602

0277 Congestive heart failure admission rate (PQI 8)
PTCA-1 VESSEL WITH AGNT
3603
OPEN CORONRY ANGIOPLASTY
3604
INTRCORONRY THROMB INFUS
3605
PTCA-MULTIPLE VESSEL
3606
INSERT OF COR ART STENT OCT95-
3607
INS DRUG-ELUT CORONRY ST OCT02-
3609
REM OF COR ART OBSTR NEC
3610
AORTOCORONARY BYPASS NOS
3611
AORTOCOR BYPAS-1 COR ART
3612
AORTOCOR BYPAS-2 COR ART
3613
AORTOCOR BYPAS-3 COR ART
3614
AORTCOR BYPAS-4+ COR ART
3615
1 INT MAM-COR ART BYPASS

0277 Congestive heart failure admission rate (PQI 8)
3616
2 INT MAM-COR ART BYPASS
3617
ABD-CORON ART BYPASS OCT96-
3619
HRT REVAS BYPS ANAS NEC
362
ARTERIAL IMPLANT REVASC
363
OTH HEART REVASCULAR
3631
OPEN CHEST TRANS REVASC
3632
OTH TRANSMYO REVASCULAR
3633
ENDO TRANSMYO REVASCULAR OCT06-
3634
PERC TRANSMYO REVASCULAR OCT06-
3639
OTH HEART REVASULAR
3691
CORON VESS ANEURYSM REP
3699
HEART VESSLE OP NEC
3731

0277 Congestive heart failure admission rate (PQI 8)
PERICARDIECTOMY
3732
HEART ANEURYSM EXCISION
3733
EXC/DEST HRT LESION OPEN
3734
EXC/DEST HRT LES OTHER
3735
PARTIAL VENTRICULECTOMY
3736
EXCISION OR DESTRUCTION OF LEFT ATRIAL APPENDAGE (LAA) OCT08-
3741
IMPLANT PROSTH CARD SUPPORT DEV OCT06
375
HEART TRANSPLANTATION (NOT VALID AFTER OCT 03)
3751
HEART TRANPLANTATION OCT03-
3752
IMPLANT TOT REP HRT SYS OCT03-
3753
REPL/REP THORAC UNIT HRT OCT03-
3754
REPL/REP OTH TOT HRT SYS OCT03-
3755
REMOVAL OF INTERNAL BIVENTRICULAR HEART REPLACEMENT SYSTEM OCT08-
3751 HEART TRANPLANTATION OCT03- 3752 IMPLANT TOT REP HRT SYS OCT03- 3753 REPL/REP THORAC UNIT HRT OCT03- 3754 REPL/REP OTH TOT HRT SYS OCT03- 3755

0277 Congestive heart failure admission rate (PQI 8)
3760
IMPLANTATION OR INSERTION OF BIVENTRICULAR EXTERNAL HEART ASSIST SYSTEM OCT08-
3761
IMPLANT OF PULSATION BALLOON
3762
INSERTION OF NON-IMPLANTABLE HEART ASSIST SYSTEM
3763
REPAIR OF HEART ASSIST SYSTEM
3764
REMOVAL OF HEART ASSIST SYSTEM
3765
IMPLANT OF EXTERNAL HEART ASSIST SYSTEM
3766
INSERTION OF IMPLANTABLE HEART ASSIST SYSTEM
3770
INT INSERT PACEMAK LEAD
3771
INT INSERT LEAD IN VENT
3772
INT INSERT LEAD ATRI-VENT
3773
INT INSER LEAD IN ATRIUM
3774
INT OR REPL LEAD EPICAR
AHRQ Quality Indicators Web Site: http://www.qualityindicators.ahrq.gov

0277 Congestive heart failure admission rate (PQI 8)
Prevention Quality Indicators Technical Specifications Version 4.2– 2010
PQI #8 Congestive Heart Failure (CHF) Admission Rate Page 3
3775
REVISION OF LEAD
3776
REPL TV ATRI-VENT LEAD
3777
REMOVAL OF LEAD W/O REPL
3778
INSER TEAM PACEMAKER SYS
3779
REVIS OR RELOCATE POCKET
3780
INT OR REPL PERM PACEMKR
3781
INT INSERT 1-CHAM, NON
3782
INT INSERT 1-CHAM, RATE
3783
INT INSERT DUAL-CHAM DEV
3785
REPL PACEM W 1-CHAM, NON
3786
REPL PACEM 1-CHAM, RATE
3787

	0277 Congestive heart failure admission rate (PQI 8)
	REPL PACEM W DUAL-CHAM
	3789
	REVISE OR REMOVE PACEMAK
	3794
	IMPLT/REPL CARDDEFIB TOT
	3795
	IMPLT CARDIODEFIB LEADS
	3796
	IMPLT CARDIODEFIB GENATR
	3797
	REPL CARDIODEFIB LEADS
	3798
	REPL CARDIODEFIB GENRATR
Denominator Statement	Population in Metro Area or county, age 18 years and older.
Denominator Categories	Female; Male 18 and older
Denominator Details	Time Window: Time period is user defined. Users of the measure typically use a 12 month time period.
	Population in Metro Area or county, age 18 years and older.
Exclusions	None
Exclusion Details	Not applicable
Risk	Risk adjustment method widely or commercially available.
Adjustment	URL http://qualityindicators.ahrq.gov/downloads/pqi/PQI_Risk_Adjustment_Tables_(Version_4_2).pdf
Stratification	Observed rates may be stratified by gender, age (5-year age groups), race / ethnicity

	0277 Congestive heart failure admission rate (PQI 8)
Type Score	Rate/proportion better quality = lower score
Algorithm	

	0286 Aspirin at arrival
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-01-02, Baltimore, MD 21244-1850
Description	Percentage of emergency department acute myocardial infarction (AMI) patients or chest pain patients (with Probable Cardiac Chest Pain) without aspirin contraindications who received aspirin within 24 hours before ED arrival or prior to transfer.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Health/Medical Record, Paper medical record/flow-sheet URL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899
Lovel	81244
Level	Facility/Agency, Population: National
Setting	Ambulatory Care: Emergency Dept, Ambulatory Care: Hospital Outpatient, Hospital
Numerator Statement	Emergency Department AMI or Chest Pain patients (with Probable Cardiac Chest Pain) who received aspirin within 24 hours before ED arrival or prior to transfer.
Numerator Details	Time Window: During the measurement period.
	Patients with:
	 An E/M Code for emergency department encounter as defined in Appendix A, Table 1.0, and Patients discharged/transferred to a short term general hospital for inpatient care, or to a Federal healthcare facility, and
	 An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1 or an ICD-9-CM Principal or Other Diagnosis Codes for Angina, Acute Coronary Syndrome, or Chest Pain as defined in Appendix A, OP Table 1.1a with Probable Cardiac Chest Pain and Patients with Aspirin Received.
Denominator Statement	Emergency Department AMI or Chest Pain patients (with Probable Cardiac Chest Pain) without aspirin contraindications.
Denominator Categories	Female; Male 18 years of age and older
Denominator Details	Time Window: During the measurement period.
	Patients with:

	0286 Aspirin at arrival
	 An E/M Code for emergency department encounter as defined in Appendix A, Table 1.0, and Patients discharged/transferred to a short term general hospital for inpatient care, or to a Federal healthcare facility, and An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1 or an ICD-9-CM Principal or Other Diagnosis Codes for Angina, Acute Coronary Syndrome, or Chest Pain as defined in Appendix A, OP Table 1.1a with Probable Cardiac Chest Pain
Exclusions	 Excluded Populations: Patients less than 18 years of age Patients with a documented reason for no aspirin on arrival.
Exclusion Details	Specifications available at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244
Risk Adjustment	No risk adjustment necessary N/A
Stratification	Specifications available at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244
Type Score	Rate/proportion better quality = higher score
Algorithm	Specifications available at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244

	0288 Fibrinolytic therapy received within 30 minutes of ed arrival
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-01-02, Baltimore, MD 21244-1850
Description	Emergency Department acute myocardial infarction (AMI) patients receiving fibrinolytic therapy during the ED stay and having a time from ED arrival to fibrinolysis of 30 minutes or less.
Туре	Process
Data Source	Electronic administrative data/claims, Electronic Health/Medical Record, Paper medical record/flow-sheet
	URL
	http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244 URL
	http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244
Level	Facility/Agency, Population: National
Setting	Ambulatory Care: Emergency Dept, Ambulatory Care: Hospital Outpatient, Hospital
Numerator Statement	Emergency Department AMI patients whose time from ED arrival to fibrinolysis is 30 minutes or less.
Numerator Details	Time Window: During the measurement period.
	Patients with:
	 An E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0, and Patients discharged/transferred to a short-term general hospital for inpatient care, or to a Federal healthcare facility, and
	 An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1, and ST-segment elevation or LBBB on the ECG performed closest to ED arrival, and Fibrinolytic Administration as defined in the Data Dictionary.
Denominator Statement	Emergency Department AMI patients with ST-segment elevation or LBBB on ECG who received fibrinolytic therapy.
Denominator Categories	Female; Male 18 years of age and older
Denominator Details	Time Window: During the measurement period.
	Patients with:

	0288 Fibrinolytic therapy received within 30 minutes of ed arrival
	 An E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0, and Patients discharged/transferred to a short-term general hospital for inpatient care, or to a Federal healthcare facility, and An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1, and ST-segment elevation or LBBB on the ECG performed closest to ED arrival, and Fibrinolytic Administration as defined in the Data Dictionary.
Exclusions	Excluded Populations:
	 Patients less than 18 years of age Patients who did not receive Fibrinolytic Administration within 30 minutes AND had a Reason for Delay in Fibrinolytic Therapy as defined in the Data Dictionary.
Exclusion	See specifications at
Details	http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244
Risk	No risk adjustment necessary
Adjustment	N/A
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	See specifications at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244

	September for Madicage 9 Medicaid Comices, 7500 County Parlament Mail Clare CO 04 00 Pattern vs MD 04044			
18	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-01-02, Baltimore, MD 212 1850			
· ·	Median time from emergency department arrival to ECG (performed in the ED prior to transfer) for acute myocardial infarction (AMI) or Chest Pain patients (with probable cardiac chest pain).			
Туре	Process			
Data Source E	Electronic administrative data/claims, Electronic Health/Medical Record, Paper medical record/flow-sheet			
ht	JRL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899			
Level Fa	Facility/Agency, Population: National			
Setting A	Ambulatory Care: Emergency Dept, Ambulatory Care: Hospital Outpatient, Hospital			
Statement T	Continuous Variable Statement: Time (in minutes) from emergency department arrival to ECG (performed in the ED prior to transfer) for acute myocardial infarction (AMI) or Chest Pain patients (with Probable Cardiac Chest Pain).			
•	ncluded Populations: PICD-9-CM Principal or Other Diagnosis Code for AMI as defined in Appendix A1, OP Table 6.1 or an ICD-9-CM Principal or Other Diagnosis Code for Angina, Acute Coronary Syndrome, or Chest Pain as defined in Appendix A1, OP Table 6.1a, and E/M Code for emergency department encounter as defined in Appendix A1, OP Table 1.0a, and Patients receiving an ECG as defined in the Appendix A1, and Patients discharged/transferred to a short term general hospital for inpatient care, to a Federal healthcare facility, or to a Critical Access Hospital.			
Numerator Topotails	ime Window: During the measurement period.			
P	Patients with:			
•	An E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0, and Patients discharged/transferred to a short term general hospital for inpatient care, or to a Federal healthcare facility, and An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1 or an ICD-9-CM Principal or Other Diagnosis Codes for Angina, Acute Coronary Syndrome, or Chest Pain as defined in Appendix A, OP Table 1.1a, and Patients receiving an ECG as defined in the Data Dictionary.			
Denominator C Statement	Continuous Variable Statement:			

	0289 Median time to ECG				
	myocardial infarction (AMI) or Chest Pain patients (with Probable Cardiac Chest Pain).				
Denominator Categories	Female; Male 18 years of age and older				
Denominator Details	Time Window: During the measurement period.				
	Patients with:				
	 An E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0, and Patients discharged/transferred to a short term general hospital for inpatient care, or to a Federal healthcare facility, and An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1 or an ICD-9-CM Principal or Other Diagnosis Codes for Angina, Acute Coronary Syndrome, or Chest Pain as defined in Appendix A, OP Table 1.1a, and 				
	Patients receiving an ECG as defined in the Data Dictionary				
Exclusions	Patients less than 18 years of age				
Exclusion	Specifications available at				
Details	http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244				
Risk	No risk adjustment necessary				
Adjustment	N/A				
Stratification	N/A				
Type Score	Continuous variable better quality = lower score				
Algorithm	Specifications available at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244				

	0290 Median time to transfer to another facility for acute coronary intervention			
Steward	Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-01-02, Baltimore, MD 21244-1850			
Description	Median time from emergency department arrival to time of transfer to another facility for acute coronary intervention.			
Туре	Process			
Data Source	Electronic administrative data/claims, Electronic Health/Medical Record, Paper medical record/flow-sheet			
	URL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244 URL			
	http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244			
Level	Can be measured at all levels, Facility/Agency, Population: National			
Setting	Ambulatory Care: Emergency Dept, Ambulatory Care: Hospital Outpatient, Hospital			
Numerator Statement	Continuous Variable Statement: Time (in minutes) from emergency department arrival to transfer to another facility for acute coronary intervention Included Populations:			
	 ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 6.1, and E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0a, and Patients discharged/transferred to a short-term general hospital for inpatient care, to a Federal healthcare facility, or to a Critical Access Hospital, and Patients not receiving Fibrinolytic Administration as defined in the Data Dictionary, and Patients with Transfer for Acute Coronary Intervention as defined in the Data Dictionary. 			
Numerator Details	Time Window: During the measurement period.			
	Patients with:			
	 An E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0, and Patients discharged/transferred to a short-term general hospital for inpatient care, or to a Federal healthcare facility, and An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1, and ST-segment elevation or LBBB on the ECG performed closest to ED arrival, and Patients with Transfer for Acute Coronary Intervention as defined in the Data Dictionary. 			
Denominator Statement	·			

	0290 Median time to transfer to another facility for acute coronary intervention				
Denominator Categories	Female; Male 18 years of age and older				
Denominator Details	Time Window: During the measurement period.				
	Patients with:				
	 An E/M Code for emergency department encounter as defined in Appendix A, OP Table 1.0, and Patients discharged/transferred to a short-term general hospital for inpatient care, or to a Federal healthcare facility, and An ICD-9-CM Principal Diagnosis Code for AMI as defined in Appendix A, OP Table 1.1, and ST-segment elevation or LBBB on the ECG performed closest to ED arrival, and Patients with Transfer for Acute Coronary Intervention as defined in the Data Dictionary. 				
Exclusions	 Patients less than 18 years of age Patients receiving Fibrinolytic Administration as defined in the Data Dictionary. 				
Exclusion	Specifications available at				
Details	http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=119628981244				
Risk	No risk adjustment necessary				
Adjustment	N/A				
Stratification	Specifications available at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244				
Type Score	Continuous variable better quality = lower score				
Algorithm	Specifications available at http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier2&cid=11962899 81244				

Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Mail Stop S3-02-01, Baltimore, MD 21244-9045 The measure estimates a hospital 30-day risk-standardized readmission rate (RSRR), defined as readmission for any cause within 30 days after the date of discharge of the index admission for patients discharged from the hospital with a principal diagnosis of heart failure (HF). Outcome Electronic administrative data/claims URL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=12190698 55841 N/A URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1182 785083979 Facility/Agency			
any cause within 30 days after the date of discharge of the index admission for patients discharged from the hospital with a principal diagnosis of heart failure (HF). Outcome Electronic administrative data/claims URL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=12190698 55841 N/A URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1182 785083979			
Electronic administrative data/claims URL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=12190698 55841 N/A URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1182 785083979			
URL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=12190698 55841 N/A URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1182 785083979			
55841 N/A URL http://www.qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1182 785083979			
Facility/Agency			
1			
Hospital			
This outcome measure does not have a traditional numerator and denominator like a core process measure (e.g., percentage of adult patients with diabetes aged 18-75 years receiving one or more hemoglobin A1c tests per year); thus, we are using this field to define the outcome.			
The outcome for this measure is 30 day all-cause readmission. We define this as readmission for any cause within 30 days from the date of discharge of the index HF admission.			
In addition, if a patient has one or more admissions within 30 days of discharge from the index admission, only one was counted as a readmission.			
Time Window: Defined as readmission for any cause within 30 days from the date of discharge of the index admission.			
Measure includes readmissions to any acute care hospital for any cause within 30 days of the index HF admission discharge date.			
Note: This outcome measure does not have a traditional numerator and denominator like a core process measure; thus, we are using this field to define the patient cohort and to define exclusions to the patient cohort.			
The cohort includes admissions for Medicare fee-for service (FFS) beneficiaries age 65 years or older discharged from the hospital with a principal diagnosis of HF (ICD-9-CM codes 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, and 428.xx) and with a complete claims history for the 12 months prior to admission.			

	0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure hospitalization
Denominator Categories	Female; Male The target population is age 65 years or older
Denominator Details	Time Window: This measure was developed with 12 months of data. Currently the measure is publicly reported with three years of index hospitalizations.
	The denominator includes patients aged 65 and older admitted to non-federal acute care hospitals for HF defined by a principal discharge diagnosis of the following (ICD-9-CM codes 402.01, 402.11, 402.91, 404.01, 404.03, 404.11, 404.13, 404.91, 404.93, and 428.xx) and with a complete claims history for the 12 months prior to admission.
	ICD-9-CM codes that define the patient cohort:
	402.01 Hypertensive heart disease, malignant, with heart failure
	402.11 Hypertensive heart disease, benign, with heart failure
	402.91 Hypertensive heart disease, unspecified, with heart failure
	404.01 Hypertensive heart and chronic kidney disease, malignant, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.03 Hypertensive heart and chronic kidney disease, malignant, with heart failure and with chronic kidney disease stage V or end stage renal disease
	404.11 Hypertensive heart and chronic kidney disease, benign, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.13 Hypertensive heart and chronic kidney disease, benign, with heart failure and chronic kidney disease stage V or end stage renal disease
	404.91 Hypertensive heart and chronic kidney disease, unspecified, with heart failure and with chronic kidney disease stage I through stage IV, or unspecified
	404.93 Hypertensive heart and chronic kidney disease, unspecified, with heart failure and chronic kidney disease stage V or end stage renal disease
	428.0 Congestive heart failure, unspecified
	428.1 Left heart failure
	428.20 Unspecified systolic heart failure
	428.21 Acute systolic heart failure
	428.22 Chronic systolic heart failure

	0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure hospitalization	
	428.23 Acute on chronic systolic heart failure	
	428.30 Unspecified diastolic heart failure	
	428.31 Acute diastolic heart failure	
	428.32 Chronic diastolic heart failure	
	428.33 Acute on chronic diastolic heart failure	
	428.40 Unspecified combined systolic and diastolic heart failure	
	428.41 Acute combined systolic and diastolic heart failure	
	428.42 Chronic combined systolic and diastolic heart failure	
	428.43 Acute on chronic combined systolic and diastolic heart failure	
	428.9 Heart Failure, unspecified.	
Exclusions	We excluded admissions for patients:	
Exclusion	 with an in-hospital death (because they are not eligible for readmission); without at least 30 days post-discharge enrollment in Medicare FFS (because the 30-day readmission outcome cannot be assessed in this group); transferred to another acute care facility (When a patient is transferred from one acute care hospital to another, these multiple contiguous hospitalizations are considered one episode of care. Readmissions for transferred patients are attributed to the hospital that ultimately discharges the patient to a non-acute care setting.); discharged against medical advice (AMA) (because providers did not have the opportunity to deliver full care and prepare the patient for discharge); admitted with HF within 30 days of discharge from an index admission. (Admissions within 30 days of discharge of an index admission will be considered readmissions. No admission is counted as a readmission and an index admission. The next eligible admission after the 30-day time period following an index admission will be considered another index admission.) See "Denominator Exclusions" section. 	
Details		
Risk Adjustment	Risk-adjustment devised specifically for this measure/condition. URL http://qualitynet.org/dcs/ContentServer?c=Page&pagename=QnetPublic%2FPage%2FQnetTier3&cid=1219069 55841	
Stratification	Results of this measure will not be stratified.	
Type Score	Rate/proportion better quality = lower score	

	0330 Hospital 30-day, all-cause, risk-standardized readmission rate following heart failure hospitalization
Algorithm	

	0355 Bilateral cardiac catheterization rate (IQI 25)		
Steward	Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, MD 20850		
Description	Percent of discharges with heart catheterizations in any procedure field with simultaneous right and left heart (bilateral) heart catheterizations.		
Туре	Outcome		
Data Source	Electronic administrative data/claims		
	URL http://www.qualityindicators.ahrq.gov/software.htm URL http://www.qualityindicators.ahrq.gov/downloads/winqi/AHRQ_QI_Windows_Software_Documentation_V41a.pdf		
Level	Facility/Agency		
Setting	Hospital		
Numerator Statement	Discharges with ICD-9-CM procedure code for right and left heart catheterization in any procedure code field.		
Numerator Details	Time Window: Inpatient hospitalization		
	ICD-9-CM right and left heart catheterization procedure code:		
	3723 RT/LEFT HEART CARD CATH		
	Exclude cases:		
	 with valid indications for right-sided catheterization ICD-9-CM Indications for Right Heart Catheterization diagnosis codes: 		
	3910 ACUTE RHEUMATIC PERICARD		
	3911 ACUTE RHEUMATIC ENDOCARD		
	3912 AC RHEUMATIC MYOCARDITIS		
	3918 AC RHEUMAT HRT DIS NEC		
	3919 AC RHEUMAT HRT DIS NOS		
	3920 RHEUM CHOREA W HRT INVOL		
	3929 RHEUMATIC CHOREA NOS		
	393 CHR RHEUMATIC PERICARD		

	ateral cardiac catheterization rate (IQI 25)
3940	MITRAL STENOSIS
3941	RHEUMATIC MITRAL INSUFF
3942	MITRAL STENOSIS W INSUFF
3949	MITRAL VALVE DIS NEC/NOS
3960	MITRAL/AORTIC STENOSIS
3961	MITRAL STENOS/AORT INSUF
3962	MITRAL INSUF/AORT STENOS
3963	MITRAL/AORTIC VAL INSUFF
3968	MITR/AORTIC MULT INVOLV
3969	MITRAL/AORTIC V DIS NOS
3970	TRICUSPID VALVE DISEASE
3971	RHEUM PULMON VALVE DIS
3979	RHEUM ENDOCARDITIS NOS
3980	RHEUMATIC MYOCARDITIS
39890	RHEUMATIC HEART DIS NOS
39891	RHEUMATIC HEART FAILURE
39899	RHEUMATIC HEART DIS NEC
40200	MAL HYPERTEN HRT DIS NOS
40201	MAL HYPERT HRT DIS W CHF
40210	BEN HYPERTEN HRT DIS NOS
40211	BENIGN HYP HRT DIS W CHF
40290	HYPERTENSIVE HRT DIS NOS
40291	HYPERTEN HEART DIS W CHF
40400	MAL HY HT/REN W/O HF/RF
40401	MAL HYPER HRT/REN W HF
	3941 3942 3949 3960 3961 3962 3963 3968 3969 3970 3971 3979 3980 39890 39890 40200 40201 40210 40211 40290 40291 40400

0355 Bi	lateral cardiac catheterization rate (IQI 25)
40402	MAL HY HT/REN W REN FAIL
40403	MAL HYP HRT/REN W HF/RF
40410	BEN HY HT/REN W/O HF/RF
40411	BEN HYPER HRT/REN W HF
40412	BEN HY HT/REN W REN FAIL
40413	BEN HYP HRT/REN W HF/RF
40490	HY HT/REN NOS W/O HF/RF
40491	HYPER HRT/REN NOS W HF
40492	HY HT/REN NOS W REN FAIL
74684	OBSTRUCT HEART ANOM NEC
74685	CORONARY ARTERY ANOMALY
74686	CONGENITAL HEART BLOCK
74687	MALPOSITION OF HEART
74689	CONG HEART ANOMALY NEC
7469	CONG HEART ANOMALY NOS
7470	PATENT DUCTUS ARTERIOSUS
74710	COARCTATION OF AORTA
74711	INTERRUPT OF AORTIC ARCH
74720	CONG ANOM OF AORTA NOS
74721	ANOMALIES OF AORTIC ARCH
74722	AORTIC ATRESIA/STENOSIS
74729	CONG ANOM OF AORTA NEC
7473	PULMONARY ARTERY ANOM
74740	GREAT VEIN ANOMALY NOS
40493	HYP HRT/REN NOS W HF/RF

0355 Bi	lateral cardiac catheterization rate (IQI 25)
4150	ACUTE COR PULMONALE
4151	PULM EMBOLISM/INFARCT-
41511	IATROGENIC PULMON. EMBOLISM
41512	SEPTIC PULMONARY EMBOLSM
41519	OTHER PULMON EMBOLISM
4160	PRIM PULM HYPERTENSION
4161	KYPHOSCOLIOTIC HEART DIS
4168	CHR PULMON HEART DIS NEC
4169	CHR PULMON HEART DIS NOS
4170	ARTERIOVEN FISTU PUL VES
4171	PULMON ARTERY ANEURYSM
4178	PULMON CIRCULAT DIS NEC
4179	PULMON CIRCULAT DIS NOS
4200	AC PERICARDIT IN OTH DIS
42090	ACUTE PERICARDITIS NOS
42091	AC IDIOPATH PERICARDITIS
42099	ACUTE PERICARDITIS NEC
4210	AC/SUBAC BACT ENDOCARD
4211	AC ENDOCARDIT IN OTH DIS
4219	AC/SUBAC ENDOCARDIT NOS
4220	AC MYOCARDIT IN OTH DIS
42290	ACUTE MYOCARDITIS NOS
42291	IDIOPATHIC MYOCARDITIS
42292	SEPTIC MYOCARDITIS
42293	TOXIC MYOCARDITIS

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0355 Bi	lateral cardiac catheterization rate (IQI 25)
42820	SYSTOLIC HRT FAILURE NOS
42821	AC SYSTOLIC HRT FAILURE
42822	CHR SYSTOLIC HRT FAILURE
42823	AC ON CHR SYST HRT FAIL
42830	DIASTOLC HRT FAILURE NOS
42831	AC DIASTOLIC HRT FAILURE
42832	CHR DIASTOLIC HRT FAIL
42833	AC ON CHR DIAST HRT FAIL
42840	SYST/DIAST HRT FAIL NOS
42841	AC SYST/DIASTOL HRT FAIL
42842	CHR SYST/DIASTL HRT FAIL
42843	AC/CHR SYST/DIA HRT FAIL
4289	HEART FAILURE NOS
7450	COMMON TRUNCUS
74510	COMPL TRANSPOS GREAT VES
74511	DOUBLE OUTLET RT VENTRIC
74512	CORRECT TRANSPOS GRT VES
74519	TRANSPOS GREAT VESS NEC
7452	TETRALOGY OF FALLOT
7453	COMMON VENTRICLE
7454	VENTRICULAR SEPT DEFECT
7455	SECUNDUM ATRIAL SEPT DEF
74560	ENDOCARD CUSHION DEF NOS
74561	OSTIUM PRIMUM DEFECT
74569	ENDOCARD CUSHION DEF NEC
	42820 42821 42822 42823 42830 42831 42832 42833 42840 42841 42842 42843 4289 7450 74510 74511 74512 74519 7452 7453 7454 7455 74560 74561

0355 Bil	ateral cardiac catheterization rate (IQI 25)
7457	COR BILOCULARE
7458	SEPTAL CLOSURE ANOM NEC
7459	SEPTAL CLOSURE ANOM NOS
74600	PULMONARY VALVE ANOM NOS
74601	CONG PULMON VALV ATRESIA
74602	CONG PULMON VALVE STENOS
74609	PULMONARY VALVE ANOM NEC
7461	CONG TRICUSP ATRES/STEN
7462	EBSTEIN'S ANOMALY
7463	CONG AORTA VALV STENOSIS
7464	CONG AORTA VALV INSUFFIC
7465	CONGEN MITRAL STENOSIS
7466	CONG MITRAL INSUFFICIENC
7467	HYPOPLAS LEFT HEART SYND
74681	CONG SUBAORTIC STENOSIS
74682	COR TRIATRIATUM
74683	INFUNDIB PULMON STENOSIS
74741	TOT ANOM PULM VEN CONNEC
74742	PART ANOM PULM VEN CONN
74749	GREAT VEIN ANOMALY NEC
7475	UMBILICAL ARTERY ABSENCE
74760	UNSP PRPHERL VASC ANOMAL
74761	GSTRONTEST VESL ANOMALY
74762	RENAL VESSEL ANOMALY
74763	UPR LIMB VESSEL ANOMALY

0355 Bilateral cardiac catheterization rate (IQI 25)
74764 LWR LIMB VESSEL ANOMALY
74769 OTH SPCF PRPH VSCL ANOML
74781 CEREBROVASCULAR ANOMALY
74782 SPINAL VESSEL ANOMALY
74783 PERSISTENT FETAL CIRC OCT02-
74789 CIRCULATORY ANOMALY NEC
7479 CIRCULATORY ANOMALY NOS
Discharges with ICD-9-CM procedure code for heart catheterizations in any procedure code field.
Female; Male 18 and older
Time Window: User defined; Most users use one calendar year
All discharges, age 18 years and older, with heart catheterization in any procedure field.
ICD-9-CM heart catheterization procedure codes:
3722 LEFT HEART CARDIAC CATH
3723RT/LEFT HEART CARD CATH
Include only cases with any diagnosis of coronary artery disease. ICD-9-CM coronary artery disease diagnosis codes:
41000 AMI ANTEROLATERAL, UNSPEC
41001 AMI ANTEROLATERAL, INIT
41002 AMI ANTEROLATERAL, SUBSEQ
41010 AMI ANTERIOR WALL, UNSPEC
41011 AMI ANTERIOR WALL, INIT
41012 AMI ANTERIOR WALL, SUBSEQ
41020 AMI INFEROLATERAL, UNSPEC

	0355 Bilateral cardiac catheterization rate (IQI 25)
	41021 AMI INFEROLATERAL, INIT
	41022 AMI INFEROLATERAL, SUBSEQ
	41030 AMI INFEROPOST, UNSPEC
	41031 AMI INFEROPOST, INITIAL
	41032 AMI INFEROPOST, SUBSEQ
	41040 AMI INFERIOR WALL, UNSPEC
	41041 AMI INFERIOR WALL, INIT
	41042 AMI INFERIOR WALL, SUBSEQ
	41050 AMI LATERAL NEC, UNSPEC
	41051 AMI LATERAL NEC, INITIAL
	41052 AMI LATERAL NEC, SUBSEQ
	41060 TRUE POST INFARCT, UNSPEC
	41061 TRUE POST INFARCT, INIT
	41062 TRUE POST INFARCT, SUBSEQ
	41070 SUBENDO INFARCT, UNSPEC
	41071 SUBENDO INFARCT, INITIAL
	41072 SUBENDO INFARCT, SUBSEQ
	41080 AMI NEC, UNSPECIFIED
	41081 AMI NEC, INITIAL
	41082 AMI NEC, SUBSEQUENT
	41090 AMI NOS, UNSPECIFIED
	41091 AMI NOS, INITIAL
	41092 AMI NOS, SUBSEQUENT
	4110 POST MI SYNDROME
	4111 INTERMED CORONARY SYND
<u> </u>	

	0355 Bilateral cardiac catheterization rate (IQI 25)
	41181 CORONARY OCCLSN W/O MI
	41189 AC ISCHEMIC HRT DIS NEC
	412 OLD MYOCARDIAL INFARCT
	4130 ANGINA DECUBITUS
	4131 PRINZMETAL ANGINA
	4139 ANGINA PECTORIS NEC/NOS
	4140 COR ATHEROSCLEROSIS OCT94-
	41400 COR ATH UNSP VSL NTV/GFT OCT94-
	41401 CRNRY ATHRSCL NATVE VSSL OCT94-
	41402 CRN ATH ATLG VN BPS GRFT OCT94-
	41403 CRN ATH NONATLG BLG GRFT OCT94-
	41404 COR ATH ARTRY BYPAS GRFT OCT96-
	41405 COR ATH BYPASS GRAFT NOS OCT96-
	41406 COR ATH NATV ART TP HRT OCT02-
	41407 COR ATH BPS GRAFT TP HRT OCT03-
	41410 ANEURYSM, HEART (WALL)
	41411 CORONARY VESSEL ANEURYSM
	41412 DISSECTION COR ARTERY OCT02-
	41419 ANEURYSM OF HEART NEC
	4143 CORONARY ATHEROSCLEROSIS DUE TO LIPID RICH PLAQUE OCT08-
	4148 CHR ISCHEMIC HRT DIS NEC
	4149 CHR ISCHEMIC HRT DIS NOS
Exclusions	None
Exclusion Details	Not applicable

	0355 Bilateral cardiac catheterization rate (IQI 25)
Risk	No risk adjustment necessary
Adjustment	None
Stratification	Observed (raw) rates may be stratified by gender, age groups, race/ethnicity categories and payer categories.
	Risk adjustment of the data is recommended using age and sex. Reliability adjustment is also recommended.
Type Score	Rate/proportion better quality = lower score
Algorithm	

	0358 Congestive heart failure (CHF) mortality rate (IQI 16)
Steward	Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, MD 20850
Description	Percent of discharges with principal diagnosis code of CHF with in-hospital mortality.
Туре	Outcome
Data Source	Electronic administrative data/claims
	URL None http://www.qualityindicators.ahrq.gov/software.htm URL http://www.qualityindicators.ahrq.gov/downloads/winqi/AHRQ_QI_Windows_Software_Documentation_V41a.pdf
Level	Facility/Agency
Setting	Hospital
Numerator Statement	Number of deaths (DISP = 20) among cases meeting the inclusion and exclusion rules for the denominator.
Numerator Details	Time Window: Time window can be determined by user, but is generally a calendar year.
	Number of deaths (DISP = 20) among cases meeting the inclusion and exclusion rules for the denominator.
Denominator Statement	All discharges, age 18 years and older, with a principal diagnosis code of CHF.
Denominator Categories	Female; Male 18 and older
Denominator Details	Time Window: Time window can be determined by user, but is generally a calendar year.
	All discharges, age 18 years and older, with a principal diagnosis code of CHF.
	ICD-9-CM CHF diagnosis codes:
	39891
	RHEUMATIC HEART FAILURE
	40201
	MAL HYPERT HRT DIS W CHF
	40211

ENIGN HYP HRT DIS W CHF 0291 YPERTEN HEART DIS W CHF 0401
YPERTEN HEART DIS W CHF 0401
0401
ALLIVOED LIDT/DEN W. CLIE
AL HYPER HRT/REN W CHF
0403
AL HYP HRT/REN W CHF&RF
0411
EN HYPER HRT/REN W CHF
0413
EN HYP HRT/REN W CHF&RF
0491
YPER HRT/REN NOS W CHF
0493
YP HT/REN NOS W CHF&RF
280
ONGESTIVE HEART FAILURE
281
EFT HEART FAILURE
2820
YSTOLIC HEART FAILURE NOS OCT02-
2821
C SYSTOLIC HRT FAILURE OCT02-
2822
HR SYSTOLIC HRT FAILURE OCT02-

	0358 Congestive heart failure (CHF) mortality rate (IQI 16)
	42823
	AC ON CHR SYST HRT FAIL OCT02-
	4289
	HEART FAILURE NOS
	42830
	DIASTOLIC HRT FAILURE NOS OCT02-
	42831
	AC DIASTOLIC HRT FAILURE OCT02-
	42832
	CHR DIASTOLIC HRT FAIL OCT02-
	42833
	AC ON CHR DIAST HRT FAIL OCT02-
	42840
	SYST/DIAST HRT FAIL NOS OCT02-
	42841
	AC SYST/DIASTOL HRT FAIL OCT02-
	42842
	CHR SYST/DIASTL HRT FAIL OCT02-
	42843
	AC/CHR SYST/DIA HRT FAIL OCT02-
	Exclude cases:
	 missing discharge disposition (DISP = missing), gender (SEX = missing), age (AGE = missing), quarter (DQTR = missing), year (YEAR = missing) or principal diagnosis (DX1 = missing) transferring to another short-term hospital (DISP = 2) MDC 14 (pregnancy, childbirth, and puerperium).
Exclusions	missing discharge disposition (DISP = missing)
	transferring to another short-term hospital (DISP = 2)

	0358 Congestive heart failure (CHF) mortality rate (IQI 16)
	MDC 14 (pregnancy, childbirth, and puerperium).
Exclusion Details	 Exclude cases: missing discharge disposition (DISP = missing), gender (SEX = missing), age (AGE = missing), quarter (DQTR = missing), year (YEAR = missing) or principal diagnosis (DX1 = missing) transferring to another short-term hospital (DISP = 2) MDC 14 (pregnancy, childbirth, and puerperium).
Risk Adjustment	Risk adjustment method widely or commercially available. URL http://qualityindicators.ahrq.gov/downloads/iqi/IQI_Risk_Adjustment_Tables_(Version_4_2).pdf
Stratification	Gender, age (5-year age groups), race / ethnicity, primary payer, custom
Type Score	Rate/proportion better quality = lower score
Algorithm	

	1522 ACE/ARB therapy at discharge for ICD implant patients with LVSD
Steward	American College of Cardiology Foundation (ACCF), 2400 N Street NW, Washington, DC 20037
Description	Proportion of ICD implant patients with a diagnosis of LVSD who are prescribed ACE-I or ARB therapy at discharge.
Туре	Process
Data Source	Electronic Clinical Data: Registry
	URL http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX URL http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX
Level	Facility
Setting	Ambulatory Care: Clinic/Urgent Care, Hospital/Acute Care Facility
Numerator Statement	Count of patients with ACE-I or ARB therapy prescribed at discharge.
Numerator Details	Time Window: 1 year
	Discharge medications = ACE inhibitor (any) = yes or ARB (any)= yes
Denominator Statement	Count of patients with an ICD implant with moderate or severe LVSD (LVEF<40%) without contraindication to ACE inhibitors and ARBs.
Denominator Categories	Female; Male All patients
Denominator Details	Time Window: 1 year
	Procedure type = initial generator implant = yes or generator change = yes
	Generator type includes single chamber, dual chamber, and biventricular (CRT-D) ICD
	Most recent LVEF<40%
Exclusions	 Patients who expired prior to discharge. Patients with ACE-I and ARB therapy contraindicated or blinded.
Exclusion Details	Discharge status = deceased ACE inhibitor (any) = contraindicated or blinded **AND** ARB (any) = contraindicated or blinded.

	1522 ACE/ARB therapy at discharge for ICD implant patients with LVSD
	Contraindicated supporting definition:
	Medication was not prescribed because of a contraindication.
	Contraindications must be documented explicitly by the physician, or clearly evidenced within the medical record
	Blinded supporting definition:
	Patient was in research study or clinical trial and administration of this specific medication is unknown
Risk	N/A
Adjustment	
Stratification	N/A
Type Score	Rate/proportion better quality = higher score
Algorithm	

	1524 Assessment of thromboembolic risk factors (CHADS2)
Steward	American College of Cardiology Foundation/ American Heart Association/American Medical Association's Physician Consortium for Performance Improvement, 2400 N. Street NW, Washington, DC 20037
Description	Patients with nonvalvular atrial fibrillation or atrial flutter in whom assessment of thromboembolic risk factors using the CHADS2 risk criteria has been documented.
Туре	Process
Data Source	Electronic Clinical Data, Electronic Clinical Data: Electronic Health Record, Electronic Clinical Data: Registry, Paper Records
	URL Journal—see Appendix E http://content.onlinejacc.org/cgi/content/full/51/8/865 https://www.pinnacleregistry.org/Documents/PINNACLE_DataCollectionForm_1.2.pdf Journal—see Appendix E URL https://www.pinnacleregistry.org/Documents/PINNACLE_DataCollectionForm_1.2.pdf
Level	Clinician: Group/Practice, Clinician: Individual
Setting	Ambulatory Care: Clinician Office
Numerator Statement	Patients with nonvalvular atrial fibrillation or atrial flutter in whom assessment of all of the specified thromeboembolic risk factors is documented.
	For patients with nonvalvular atrial fibrillation or atrial flutter, assessment of thromboembolic risk should include the following factors:
	Electronic Specifications:
	Risk factors:
	Prior stroke or transient ischemic attack> High risk
	Age = 75 years> Moderate risk
	Hypertension> Moderate risk
	Diabetes mellitus> Moderate risk
	Heart failure or impaired LV systolic function> Moderate risk
Numerator Details	Time Window: Reporting year
Denominator Statement	All patients 18 years of age or older with nonvalvular atrial fibrillation or atrial flutter other than those specifically excluded
Denominator	Female; Male 18 years or older

	1524 Assessment of thromboembolic risk factors (CHADS2)
Categories	
Denominator Details	Time Window: Reporting year
	For Claims/Administrative: Denominator (Eligible Population): All patients aged 18 years and older with a diagnosis of nonvalvular AF or
	atrial flutter
	ICD-9 diagnosis codes: 427.31, 427.32
	AND
	Not ICD-9 diagnosis codes: 394.0, 394.2 (mitral stenosis); 996.02, 996.71, V42.2, V43.3 (prosthetic heart valve)
	AND
	CPT E/M Service Code: 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, 99241, 99242,
	99243, 99245
	Numerator: Patients with an assessment of all of the specified thromboembolic risk factors documented during
	the 12 month reporting period
	CPT Category II code: 1180F- All specified thromboembolic risk factors assessed
	Denominator Exclusion: Documentation of medical reason(s) for not having an assessment of all of the specified
	thromboembolic risk factors documented during the 12 month reporting period
	Append modifier to CPT Category II code: 1180F-1P
Exclusions	 Patients with mitral stenosis or prosthetic heart valves. Patients with transient or reversible causes of atrial fibrillation (e.g., pneumonia or hyperthyroidism). Postoperative patients. Patients who are pregnant. Medical reason(s) documented by a physician, nurse practitioner, or physician assistant for not assessing risk
	factors. Examples of medical reasons for not assessing risk factors include but are not limited to the following: o allergy to warfarin and other anticoagulant drugs that are FDA approved for the prevention of thromboembolism o risk of bleeding
Exclusion Details	None
Risk Adjustment	No risk adjustment necessary

	1524 Assessment of thromboembolic risk factors (CHADS2)
	None
Stratification	None
Type Score	Rate/proportion better quality = higher score
Algorithm	

1525 Chronic anticoagulation therapy
American College of Cardiology Foundation/American Heart Association/American Medical Association's Physician Consortium for Performance Improvement, 2400 N. Street NW, Washington, DC 20037
Prescription of warfarin or another anticoagulant drug that is FDA approved for the prevention of thromboembolism for all patients with nonvalvular atrial fibrillation or atrial flutter at high risk for thromboembolism, according to CHADS2 risk stratification.
Process
Electronic Clinical Data, Electronic Clinical Data: Electronic Health Record, Electronic Clinical Data: Registry, Paper Records
URL Journal- see Appendix E http://content.onlinejacc.org/cgi/content/full/51/8/865 https://www.pinnacleregistry.org/Documents/PINNACLE_DataCollectionForm_1.2.pdf Journal- see Appendix E URL https://www.pinnacleregistry.org/Documents/PINNACLE_DataCollectionForm_1.2.pdf
Clinician: Individual
Ambulatory Care: Clinician Office
All patients with nonvalvular atrial fibrillation or atrial flutter at high risk of thromboembolism (i.e., those with any high-risk factor or more than 1 moderate-risk factor) who are prescribed warfarin OR another anticoagulant drug that is FDA approved for the prevention of thromboembolism.
Time Window: Reporting year
Patients with nonvalvular AF or atrial flutter for whom assessment of the specified thromboembolic risk factors documented one or more high-risk factor or more than one moderate-risk factor.
Female; Male 18 years or older
Time Window: Reporting year
Claims/Administrative: Denominator (Eligible Population): All patients aged 18 years and older with a diagnosis of nonvalvular AF or
atrial flutter at high risk for thromboembolism
ICD-9 diagnosis codes: 427.31, 427.32
AND
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	1525 Chronic anticoagulation therapy
	Not ICD-9 diagnosis codes: 394.0, 394.2 (mitral stenosis); 996.02, 996.71, V42.2, V43.3 (prosthetic heart valve)
	AND
	CPT E/M Service Code: 99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, 99241, 99242,
	99243, 99245
	AND (Report a CPT Category II code for risk of thromboembolism)
	 CPT Category II code: 3552F- High risk for thromboembolism CPT Category II code: 3551F- Intermediate risk for thromboembolism CPT Category II code: 3550F- Low risk for thromboembolism NOTE: ONLY PATIENTS AT HIGH RISK FOR THROMBOEMBOLISM ARE INCLUDED IN THE MEASURE'S
	DENOMINATOR WHEN CALCULATING PERFORMANCE
	Numerator: Patients who were prescribed warfarin during the 12 month reporting period
	CPT Category II code: 4012F-Warfarin therapy prescribed Denominator Exclusion: Documentation of medical reason(s) for not prescribing warfarin during the 12 month
	reporting period
	Append modifier to CPT Category II code: 4012F-1P Documentation of patient reason(s) for not prescribing warfarin during the 12 month reporting period
	Append modifier to CPT Category II code: 4012F-2P Electronic Specifications:
	The assessment of patients with nonvalvular AF for thromboembolic risk factors should include the following criteria:
	Risk factors:
	 Prior stroke or transient ischemic attack> High risk Age = 75 years> Moderate risk Hypertension> Moderate risk Diabetes mellitus> Moderate risk Heart failure or impaired LV systolic function> Moderate risk
Exclusions	 Patients with mitral stenosis or prosthetic heart valves. Patients at low risk for thromboembolism (i.e., those with none of the risk factors listed above). Patients with only one moderate risk factor. Postoperative patients. Patients with transient or reversible causes of AF (e.g., pneumonia or hyperthyroidism). Patients who are pregnant. Medical reason(s) documented by a physician, nurse practitioner, or physician assistant for not prescribing warfarin or another anticoagulant drug that is FDA approved for the prevention of thromboembolism. Examples of medical reasons include, but are not limited to:

	1525 Chronic anticoagulation therapy
	 Allergy Risk of bleeding. Documentation of patient reason(s) for not prescribing warfarin or another anticoagulant drug that is FDA approved for the prevention of thromboembolism (e.g., economic, social, and/or religious impediments, noncompliance or patient refusal).
Exclusion	None
Details	
Risk Adjustment	No risk adjustment necessary N/A
Stratification	None
Type Score	Rate/proportion better quality = higher score
Algorithm	

	1528 Beta blocker at discharge for ICD implant patients with a previous MI
Steward	American College of Cardiology Foundation (ACCF), 2400 N Street NW, Washington, DC 20037
Description	Proportion of ICD implant patients with a diagnosis of previous MI who are prescribed a Beta Blocker at discharge.
Туре	Process
Data Source	National Cardiovascular Data Registry (NCDR)® ICD RegistryTM http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX URL http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX URL http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX
Level	Affects large numbers; Frequently performed procedure; Leading cause of morbidity/mortality; High resource use; Severity of illness.
Setting	Facility
Numerator Statement	Count of patients discharged on beta-blocker therapy.
Numerator Details	Time Window: 1 year
	discharge medication of beta blocker (any)= yes
Denominator Statement	Count of patients with an ICD implant without contraindication to beta-blockers.
Denominator Categories	Female; Male All Patients
Denominator Details	Time Window:
	1 year
Exclusions	Procedure type = initial generator implant = yes or generator change = yes
	Generator type includes single chamber, dual chamber, and biventricular (CRT-D) ICD
	Previous MI = yes
Exclusion Details	Patients who expired. Beta-blocker therapy contraindicated or blinded.
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	1528 Beta blocker at discharge for ICD implant patients with a previous MI
	Contraindicated supporting definition:
	Medication was not prescribed because of a contraindication.
	Contraindications must be documented explicitly by the physician, or clearly evidenced within the medical record.
	Blinded supporting definition:
	Patient was in research study or clinical trial and administration of this specific medication is unknown.
Risk Adjustment	N/A
Stratification	Discharge status = deceased
	Beta blocker (any) = contraindicated or blinded
Type Score	Rate/proportion
Algorithm	better quality = higher score
1	

1529 Beta blocker at discharge for ICD implant patients with LVSD
American College of Cardiology Foundation, 2400 N Street NW, Washington, DC 20037
Proportion of ICD implant patients with a diagnosis of LVSD who are prescribed beta-blocker therapy on discharge.
Process
National Cardiovascular Data Registry (NCDR)® ICD RegistryTM http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX URL http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX URL http://www.ncdr.com/WebNCDR/ICD/ELEMENTS.ASPX
Affects large numbers; Frequently performed procedure; Leading cause of morbidity/mortality; High resource use; Severity of illness.
Facility/Agency
Count of patients with beta blocker therapy prescribed on discharge.
Time Window: 1 year
discharge medication of beta blocker (any) = yes
Count of patients with an ICD implant with LVSD without contraindication to beta blockers
Female; Male All Patients
Time Window:
1 year
Procedure type = initial generator implant = yes or generator change = yes
Most recent LVEF<40%
Patients who expired. Beta blocker therapy contraindicated or blinded.

	1529 Beta blocker at discharge for ICD implant patients with LVSD
Risk	N/A
Adjustment	
Stratification	
Type Score	Rate/proportion
Algorithm	better quality = higher score

637

	0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge
Steward	American College of Cardiology Foundation, 2400 N. Street NW, Washington, DC 20037
Description	Patients undergoing PCI who receive prescriptions for all medications (aspirin, P2Y12 and satins) for which they are eligible for at discharge.
Туре	Composite with component measures combined at patient level.
Data Source	Registry Data http://www.ncdr.com/WebNCDR/ELEMENTS.ASPX
Level	Facility
Setting	Hospital
Numerator Statement	Patients who receive all medications for which they are eligible. 1. Aspirin prescribed at discharge (if eligible for aspirin as described in denominator) AND 2. P2Y12 agent (clopidogrel, prasurgel, or ticlopidine) prescribed at discharge (if eligible for P2Y12 as
	described in denominator) AND 3. Statin prescribed at discharge (if eligible for statin as described in denominator).
Numerator Details	Time Window: 1 year
	Count of patients with PCI procedures with
	[((ASA=yes) AND (ASA not contraindicated or blinded) AND
	((p2Y12=yes) AND (p2Y12 not contraindicated or blinded) AND
	(patient with PCI procedure with stents implanted)) AND
	((statin=yes) and (statin not contraindicated or blinded))]
	AND
	[Discharge status=alive) AND
	(Discharge Location=home, extended care facility, nursing home, other)]

	0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge
Denominator	All patients surviving hospitalization who are eligible to receive any one of the three medication classes:
Statement	3. Eligible for aspirin (ASA): Patients undergoing PCI who do not have contraindication to aspirin documented OR
	 Eligibility for P2Y12 agent (clopidogrel, prasurgel, or ticlopidine): Patients undergoing PCI with stenting who do not have a contraindication to P2Y12 agent documented OR
	5. Eligibility for statin therapy: Patients undergoing PCI who do not have a contraindication to stain therapy.
Denominator Categories	Female and Male 18 years of age and older
Denominator Details	Time Window: 1 year
	Count of patients with PCI procedures with
	[(ASA not contraindicated or blinded) OR
	[((p2Y12 not contraindicated or blinded) AND (patient with PCI procedure with stents implanted)) OR
	(statin not contraindicated or blinded))]]
	AND
	[Discharge status=alive) AND
	(Discharge Location=home, extended care facility, nursing home, other)]
Exclusions	Discharge statue of expired; not eligible for aspirin, P2Y12, or statin (contraindicated or blinded to all 3 medications).
Exclusion Details	N/A
Risk Adjustment	
Stratification	N/A
Type Score	Non-weighted score/composite/scale better quality = Higher score

0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge
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Denominator: Count of patients with PCI procedures with
[(ASA not contraindicated or blinded) OR
[((p2Y12 not contraindicated or blinded) AND (patient with PCI procedure with stents implanted)) OR
(statin not contraindicated or blinded))]]
AND
[Discharge status=alive) AND
(Discharge Location=home, extended care facility, nursing home, other)]
Numerator: Count of patients with PCI procedures with
[((ASA=yes) AND (ASA not contraindicated or blinded) AND
((p2Y12=yes) AND (p2Y12 not contraindicated or blinded) AND
(patient with PCI procedure with stents implanted)) AND
((statin=yes) and (statin not contraindicated or blinded))]
AND
[Discharge status=alive) AND
(Discharge Location=home, extended care facility, nursing home, other)]

639

0965 Patients with an ICD Implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge
American College of Cardiology Foundation, 2400 N. Street NW, Washington, DC 20037
Proportion of patients with an ICD implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge (all-or-none composite measures of two medications classes).
Composite with component measures combined at patient-level.
Registry Data
http://www.ncdr.com/WebNCDR/ELEMENTS.ASPX
Facility
Hospital
Patients who receive all medications for which they are eligible.
ACE/ARB prescribed at discharge (if eligible for ACE/ARB as described in denominator) AND
2. Beta blockers prescribed at discharge (if eligible for beta blockers as described in denominator)
Time Window: 1 year
Count of ICD implants patients with
[(ACE/ARB=yes) AND [(EF<40) AND (ACE/ARB not contraindicated or blinded)]] AND
[[Beta blocker=yes) AND [(EF<40) AND/OR (previous MI)]] AND (beta blockers not contraindicated or blinded)]
AND
[Discharge status=alive) AND (Discharge Against Medical Advice=No)]
All patients with an ICD implant surviving hospitalization who are eligible to receive any of the two medication classes:
Eligible for ACE/ARB: Patients who have a ejection fraction (EF) of 40% AND do not have a documented contraindication to ACE/ARB documented OR

	0965 Patients with an ICD Implant who receive prescriptions for all medications (ACE/ARB and beta
	blockers) for which they are eligible for at discharge
	2. Eligibility for beta blockers: Patients who do not have documented contraindication to beta blocker therapy
	and have either:
	a. EF of 40% OR b. A previous myocardial infarction 9MI)
Donominator	, , , , , , , , , , , , , , , , , , , ,
Categories	Female and Male 18 years of age and older
Denominator Details	Time Window: 1 year
	Count of ICD implants patients with
	[(EF<40) AND (ACE/ARB not contraindicated or blinded)]] OR
	[(EF<40) AND/OR (previous MI)]] AND (beta blockers not contraindicated or blinded)]
	AND
	[Discharge status=alive) AND (Discharge Against Medical Advice=No)]
Exclusions	Discharge status of expired; not eligible for either ACE/ARB or beta blockers.
Exclusion	Medication prescribed at discharge coded as 'contraindicated" or 'blinded" for beta blocker or ACE/ARB.
Details	Discharge status = deceased.
Risk	N/A
Adjustment	
Stratification	N/A
Type Score	Non-weighted score/composite/scale better quality = Higher score
Algorithm	Denominator: Count of ICD implants patients with
	[(EF<40) AND (ACE/ARB not contraindicated or blinded)]] OR
	[(EF<40) AND/OR (previous MI)]] AND (beta blockers not contraindicated or blinded)]
	AND

0965 Patients with an ICD Implant who receive prescriptions for all medications (ACE/ARB and beta blockers) for which they are eligible for at discharge
[Discharge status=alive) AND (Discharge Against Medical Advice=No)]
Numerator: Count of ICD implants patients with
[(ACE/ARB=yes) AND [(EF<40) AND (ACE/ARB not contraindicated or blinded)]] AND
[[Beta blocker=yes) AND [(EF<40) AND/OR (previous MI)]] AND (beta blockers not contraindicated or blinded)]
AND
[Discharge status=alive) AND (Discharge Against Medical Advice=No)]

641

APPENDIX B – NATIONAL VOLUNTARY CONSENSUS STANDARDS: CARDIOVASCULAR ENDORSEMENT MAINTENANCE 2010 STEERING COMMITTEE AND NQF STAFF

Raymond Gibbons, MD (Chair)

Mayo Clinic, Rochester, MN

Mary George, MD, MSPH (Vice Chair)

Centers for Disease Control and Prevention, Atlanta, GA

Carol Allred, RN

National Coalition for Women with Heart Disease, Harker Heights, TX

Rochelle Ayala, MD, FACP

Memorial Healthcare System, Hollywood, FL

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APPENDIX C—ENDORSED CARDIOVASCULAR CONSENSUS STANDARDS (AFTER 2008)

Coronary Artery Disease (CAD)—Secondary Prevention

Measure Number	Title	Description	Measure Steward
0543	Coronary artery disease and medication possession ratio for statin therapy	Medication Possession Ratio (MPR) for statin therapy for individuals over 18 years of age with coronary artery disease.	Centers for Medicare & Medicaid Services
0551	ACE inhibitor/angiotensin receptor blocker use and persistence among members with coronary artery disease at high risk for coronary events	To assess the use of and persistence to ACE inhibitors or angiotensin receptor blockers (ARBs) among members with CAD or other atherosclerotic vascular disease (i.e., peripheral arterial disease, atherosclerotic aortic disease and carotid artery disease) who are at high risk for coronary events during a one-year period. High-risk comorbidities are defined as heart failure, hypertension, diabetes, or chronic kidney disease (excluding stage V and patients on dialysis).	Health Benchmarks, Inc, IMS Health
0569	Adherence to lipid lowering medication	To ensure that members who are taking medications to treat hyperlipidemia filled an adequate supply of medications over a predefined time period.	Health Benchmarks, Inc, IMS Health
0583	Dyslipidemia new med 12-week lipid test	This measure identifies patients age 18 or older who started lipid-lowering medication during the measurement year and had a lipid panel checked within 3 months after starting drug therapy.	Resolution Health, Inc.
0594	Post MI: ACE inhibitor or ARB therapy	This measure identifies patients with ST elevation MI (STEMI), or non-ST elevation MI (NSTEMI) plus a history of hypertension, heart failure, and/or diabetes prior to the measurement year who are taking an ACEI or an ARB during the measurement year.	Resolution Health, Inc.
0611	Hyperlipidemia (primary prevention)—lifestyle changes and/or lipid lowering therapy	Percentage of patients with coronary artery disease risk factors who have an elevated LDL and who have initiated therapeutic lifestyle changes or are taking a lipid-lowering agent	ActiveHealth Management
0613	MI—use of beta blocker therapy	Percentage of patients who had a myocardial infarction (MI) and are taking a beta blocker.	ActiveHealth Management
0616	Atherosclerotic disease—lipid panel monitoring	Percentage of patients with coronary artery, cerebrovascular, or peripheral vascular disease that have been screened for	ActiveHealth Management

		dyslipidemia with a lipid profile.	
0631	Secondary prevention of cardiovascular events—use of aspirin or antiplatelet therapy	Percentage of patients with ischemic vascular disease (IVD) that are taking aspirin or an antiplatelet agent.	ActiveHealth Management
0636	Atherosclerotic disease and LDL greater than 100—use of lipid lowering agent	Percentage of adult patients with atherosclerotic disease and an LDL greater than 100 that are taking a lipid lowering agent.	ActiveHealth Management
Acute My	yocardial Infarction (AMI)—Emerg		
660	Troponin results for emergency department acute myocardial infarction (AMI) patients or chest pain patients (with probable cardiac chest pain) received within 60 minutes of arrival	Emergency Department acute myocardial infarction (AMI) patients or chest pain patients (with Probable Cardiac Chest Pain) with an order for Troponin during the stay and having a time from ED arrival to completion of Troponin results within 60 minutes of arrival.	Centers for Medicare & Medicaid Services
Acute My	yocardial Infarction (AMI)—Hospit	al	
0639	Statin prescribed at discharge	Percent of acute myocardial infarction (AMI) patients 18 years of age or older who are prescribed a statin medication at hospital discharge.	Centers for Medicare & Medicaid Services
704	Proportion of AMI patients that have a potentially avoidable complication (during the index stay or in the 30-day post-discharge period)	Percent of adult population aged 18-65 years who were admitted to a hospital with acute myocardial infarction (AMI), were followed for one month after discharge, and had one or more potentially avoidable complications (PACs).	Bridges to Excellence
730	Acute myocardial infarction (AMI) mortality rate	Number of deaths per 100 discharges with a principal diagnosis code of acute myocardial infarction.	Agency for Healthcare Research and Quality
0505	Thirty-day all-cause risk standardized readmission rate following acute myocardial infarction (AMI) hospitalization.	Hospital-specific 30-day all-cause risk standardized readmission rate following hospitalization for AMI among Medicare beneficiaries aged 65 years or older at the time of index hospitalization.	Centers for Medicare & Medicaid Services
698	30-day post-hospital AMI discharge care transition composite measure	This measure scores a hospital on the incidence among its patients during the month following discharge from an inpatient stay having a primary diagnosis of heart failure for three types of events: readmissions, ED visits, and evaluation and	Centers for Medicare & Medicaid Services

		management (E&M) services.	
Percuta	neous Coronary Interventions (PC	U , ,	<u>. </u>
0588	Stent drug-eluting clopidogrel	This measure identifies patients undergoing percutaneous coronary intervention (PCI) with placement of a drug-eluting intracoronary stent during the first 9 months of the measurement year, who filled a prescription for clopidogrel in the 3 months following stent placement.	Resolution Health, Inc.
695	Hospital 30-day risk- standardized readmission rates following percutaneous coronary intervention (PCI)	This measure estimates hospital risk-standardized 30-day readmission rates following PCI in patients at least 65 years of age. As PCI patients may be readmitted electively for staged revascularization procedures, we will exclude such elective readmissions from the measure. The measure uses clinical data available in the National Cardiovascular Disease Registry (NCDR) CathPCI Registry for risk adjustment that has been linked with the administrative claims data used to identify readmissions.	Centers for Medicare & Medicaid Services
0536	30-day all-cause risk- standardized mortality rate following percutaneous coronary intervention (PCI) for patients with ST segment elevation myocardial infarction (STEMI) or cardiogenic shock	Hospital-specific 30-day all-cause risk-standardized mortality rate following Percutaneous Coronary Intervention (PCI) among patients aged 18 years or older with ST segment elevation myocardial infarction (STEMI) or cardiogenic shock at the time of procedure.	Centers for Medicare & Medicaid Services
0535	30-day all-cause risk- standardized mortality rate following percutaneous coronary intervention (PCI) for patients without ST segment elevation myocardial infarction (STEMI) and without cardiogenic shock	Hospital-specific 30-day all-cause risk-standardized mortality rate following Percutaneous Coronary Intervention (PCI) among patients aged 18 years or older without ST segment elevation myocardial infarction (STEMI) and without cardiogenic shock at the time of procedure.	Centers for Medicare & Medicaid Services
0355	Bilateral cardiac catheterization rate (IQI 25)	Percent of discharges with heart catheterizations in any procedure field with simultaneous right and left heart (bilateral) heart catheterizations.	Agency for Healthcare Research and Quality

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Cardiac	Imaaina
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Cardiac	iiiayiiiy		
669	Cardiac imaging for preoperative risk assessment for non-cardiac low-risk surgery	This measure calculates the percentage of low-risk, non-cardiac surgeries performed at a hospital outpatient facility with a Stress Echocardiography, SPECT MPI or Stress MRI study performed in the 30 days prior to the surgery at a hospital outpatient facility (e.g., endoscopic, superficial, cataract surgery, and breast biopsy procedures). Results are to be segmented and reported by hospital outpatient facility where the imaging procedure was performed.	Centers for Medicare & Medicaid Services
670	Cardiac stress imaging not meeting appropriate use criteria: Preoperative evaluation in low risk surgery patients	Percentage of stress SPECT MPI, stress echo, CCTA, or CMR performed in low risk surgery patients for preoperative evaluation.	American College of Cardiology Foundation
671	Cardiac stress imaging not meeting appropriate use criteria: routine testing after percutaneous coronary interventions (PCI)	Percentage of all stress SPECT MPI, stress echo, CCTA and CMR performed routinely after PCI, with reference to timing of test after PCI and symptom status.	American College of Cardiology Foundation
672	Cardiac stress imaging not meeting appropriate use criteria: testing in asymptomatic, low risk patients	Percentage of all stress SPECT MPI, stress echo, CCTA, and CMR performed in asymptomatic, low CHD risk patients for initial detection and risk assessment	American College of Cardiology Foundation
Cardiac	Rehabilitation		
0642	Cardiac rehabilitation patient referral from an inpatient setting	Percentage of patients admitted to a hospital with a primary diagnosis of an acute myocardial infarction or chronic stable angina or who during hospitalization have undergone coronary artery bypass (CABG) surgery, a percutaneous coronary intervention (PCI), cardiac valve surgery (CVS), or cardiac transplantation who are referred to an early outpatient cardiac rehabilitation/secondary prevention program.	ACCF/AHA Task Force on Performance Measures
0643	Cardiac rehabilitation patient referral from an outpatient setting	Percentage of patients evaluated in an outpatient setting who in the previous 12 months have experienced an acute myocardial infarction or chronic stable angina or who have undergone coronary artery bypass (CABG) surgery, a percutaneous coronary intervention (PCI), cardiac valve surgery (CVS), or	ACCF/AHA Task Force on Performance Measures

		cardiac transplantation, who have not already participated in an early outpatient cardiac rehabilitation/secondary prevention program for the qualifying event, and who are referred to an	
		outpatient cardiac rehabilitation/secondary prevention program.	
Atrial Fil	brillation		
0600	New atrial fibrillation: Thyroid function test	This measure identifies patients with new-onset atrial fibrillation during the measurement year who have had a thyroid function test 6 weeks before or after the diagnosis of atrial fibrillation.	Resolution Health, Inc.
0624	Atrial fibrillation—warfarin therapy	Percentage of adult patients with atrial fibrillation and major stroke risk factors on warfarin.	ActiveHealth Management
0578	Ambulatory initiated amiodarone therapy: TSH test	This measure identifies the percentage of patients who had a TSH baseline measurement at the start of amiodarone therapy.	Resolution Health, Inc.
ICD Imp			
694	Hospital risk-standardized complication rate following implantation of implantable cardioverter-defibrillator (ICD)	This measure provides hospital specific risk-standardized rates of procedural complications following the implantation of an ICD in patients at least 65 years of age. The measure uses clinical data available in the National Cardiovascular Data Registry (NCDR) ICD Registry for risk adjustment that has been linked with administrative claims data used to identify procedural complications.	Centers for Medicare & Medicaid Services
Hyperte	nsion		
0605	Patient(s) that had a serum creatinine in last 12 reported months	This measure identifies patients with hypertension (HTN) that had a serum creatinine in last 12 reported months.	Ingenix
Heart Fa	ilure—Hospital		ļ.
699	30-day post-hospital HF discharge care transition composite measure	This measure scores a hospital on the incidence among its patients during the month following discharge from an inpatient stay having a primary diagnosis of heart failure for three types of events: readmissions, ED visits, and evaluation and management (E&M) services.	Centers for Medicare & Medicaid Services
0358	Congestive heart failure mortality (IQI 16) (risk adjusted)	Percent of in-hospital death for discharges, 18 years and older, with ICD-9-CM principle diagnosis code of CHF.	Agency for Healthcare Research and Quality
0330	30-day all-cause risk	Hospital-specific, risk-standardized, 30-day all-cause	Centers for

	standardized readmission rate following heart failure hospitalization (risk adjusted)	readmission rates for Medicare fee-for-service patients discharged from the hospital with a principal diagnosis of heart failure (HF).	Medicare & Medicaid Services
Heart Fa	ailure—Outpatient		<u>J</u>
0521	Heart failure symptoms addressed	Percent of patients exhibiting symptoms of heart failure for whom appropriate actions were taken.	Centers for Medicare & Medicaid Services
0610	Heart failure—use of ACE inhibitor (ACEI) or angiotensin receptor blocker (ARB) therapy	Percentage of patients with heart failure that are on an ACEI or ARB.	ActiveHealth Management
0615	Heart failure—use of beta blocker therapy	Percentage of adult patients with heart failure that are on a beta blocker.	ActiveHealth Management

APPENDIX D—GAPS IN THE CARDIOVASCULAR PORTFOLIO

The measures in the cardiovascular portfolio have been assigned to appropriate domains reflecting the priorities and goals of NQF, the National Priorities Partnership, and the National Quality Strategy. Large gaps in the areas of patient and family engagement and patient-reported outcomes persist. Additional measures are needed to address access and affordability.

					Domains			
	Cross- Cutting	Care Coordination & Management	Population Health	Safety	Affordability	Access	Patient & Family Engagement	Patient- Reported Outcomes
Cardiovascular TOPIC AREA	including across settings, disciplines, populations, & combinations Composites Emphasis on longitudinal measures over specified timeframe (e.g., 30 day, etc.)	including communication, pre-, intra-, post-op care; secondary prevention; intermediate outcomes, recovery & rehabilitation, end-of-life care	including prevention, healthfullifestyle behaviors, populations at risk & disparities (age, race/ ethnicity, gender, geographic & socio-economic)	including morbidity & mortality, complications of care, prevention of adverse events including HAI, medication adherence/use	including cost/efficiency direct/indirect cost, overuse, underuse, appropriateness	including access, timeliness	including Patient/ Family Engagement (perspective, knowledge, understanding, motivation, risk behavior), Satisfaction, Self- management, Shared Decision making	including Health Status (quality of life, functional status, productivity, burden on patient & family)
Coronary Artery Disease (CAD)— Secondary Prevention								
0073 IVD: Blood pressure management		Х	Х	Х				
0068 IVD: Use of aspirin or antithrombotics		Χ	_	Χ				
0067 CAD: Antiplatelet therapy		Χ		Х				
0631 Secondary prevention of cardiovascular		Χ		Χ				

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Cardiovascular TOPIC AREA	including across settings, disciplines, populations, & combinations Composites Emphasis on longitudinal measures over specified timeframe (e.g., 30 day, etc.)	including communication, pre-, intra-, post-op care; secondary prevention; intermediate outcomes, recovery & rehabilitation, end-of-life care	including prevention, healthfullifestyle behaviors, populations at risk & disparities (age, race/ ethnicity, gender, geographic & socio-economic)	including morbidity & mortality, complications of care, prevention of adverse events including HAI, medication adherence/use	including cost/efficiency direct/indirect cost, overuse, underuse, appropriateness	including access, timeliness	including Patient/ Family Engagement (perspective, knowledge, understanding, motivation, risk behavior), Satisfaction, Self- management, Shared Decision making	including Health Status (quality of life, functional status, productivity, burden on patient & family)
Events—Use of Aspirin or Antiplatelet Therapy	,							
0611 Hyperlipidemia (primary prevention) — lifestyle changes and/or lipid lowering Therapy		Х		Х				
0583 Dyslipidemia new med 12-week lipid test		Х		Х				
0569 Adherence to lipid-lowering medication		Χ		Χ				
0543 Coronary artery disease and medication possession ratio for statin therapy		Х		Х				
0075 IVD—Complete lipid profile and LDL control <100		Х		Х				
0074 Chronic stable coronary artery disease: lipid control		Х	X	Х				
0616 Atherosclerotic disease—lipid panel monitoring		Х						
0636 Atherosclerotic disease and LDL greater than 100—use of lipid lowering agent		Х	Х	Х				
0066 CAD: ACEI/ARB therapy		Х		Х				
0551 ACE inhibitor/angiotensin receptor Blocker Use and Persistence Among Members with Coronary Artery Disease at		Х		Х				
High Risk for Coronary Events								

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Cardiovascular TOPIC AREA	including across settings, disciplines, populations, & combinations Composites Emphasis on longitudinal measures over specified timeframe (e.g., 30 day, etc.)	including communication, pre-, intra-, post-op care; secondary prevention; intermediate outcomes, recovery & rehabilitation, end-of-life care	including prevention, healthfullifestyle behaviors, populations at risk & disparities (age, race/ ethnicity, gender, geographic & socio-economic)	including morbidity & mortality, complications of care, prevention of adverse events including HAI, medication adherence/use	including cost/efficiency direct/indirect cost, overuse, underuse, appropriateness	including access, timeliness	including Patient/ Family Engagement (perspective, knowledge, understanding, motivation, risk behavior), Satisfaction, Self- management, Shared Decision making	including Health Status (quality of life, functional status, productivity, burden on patient & family)
0594 Post MI: ACE inhibitor or ARB therapy	,	Х		Χ				
0071 AMI—Persistence of beta blocker		Х		Χ				
therapy								
0613 MI—Use of beta blocker therapy		Х		Χ				
0076 - Optimal vascular care		Х	Х	Х				
Acute Myocardial Infarction (AMI) —								
Emergency Department								
0289 Median to ECG				Х		X		
0660 Troponin results for emergency		Х		Х				
department acute myocardial infarction (AMI)								
patients or chest pain patients (with probable								
cardiac chest pain) received within 60 minutes								
of arrival		V		V				
0132 Aspirin at arrival for AMI		X		X				
0286 Aspirin at arrival						V		
0163 Primary PCI within 90 minutes of arrival		X		X		X		
0164 Fibrinolytic therapy received within 30 minutes		, x		X		X		
0288 Fibrinolytic therapy received within 30		Х		Х		Х		
minutes of ED arrival		٨		Χ		Χ		
0287 Median time to fibrinolysis		Х		Х		Χ		

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0290 Median time to transfer to another facility		Х				Х		
AMI —Hospital								
0160 Beta blocker prescribed at discharge		Х		Х				
0142 Aspirin prescribed at discharge for AMI		Х		Х				
0137 ACEI/ARB at discharge for AMI		Х		Χ				
0639 Statin prescribed at discharge		Х		Х				
0704 Proportion of AMI patients that have a potentially avoidable complication (during the index stay or in the 30-day post-discharge period)		Х		Х				
0730 Acute myocardial infarction (AMI) mortality rate [in patient]		Х		Х				
0230 AMI 30-day mortality		Х		Х				
0505 Thirty-day all-cause risk standardized readmission rate following acute myocardial infarction (AMI) hospitalization.		Х						
0698 30-day post-hospital AMI discharge care		Х		Х				

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transition composite measure	Cic.j							
Percutaneous Coronary Interventions (PCI)								
0588 Stent drug-eluting clopidogrel		Х						
0964 Therapy with aspirin, P2Y12 inhibitor and statin at discharge		Х		Х				
0695 Hospital 30-day risk-standardized readmission rates following percutaneous coronary intervention (PCI)		Х						
0535 30-day all-cause risk-standardized mortality rate following percutaneous coronary intervention (PCI) for patients without ST segment elevation myocardial infarction (STEMI) and without cardiogenic shock		X		X				
0536 30-day all-cause risk-standardized mortality rate following Percutaneous Coronary Intervention (PCI) for patients with ST segment elevation myocardial infarction		Х		Х				

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(STEMI) or cardiogenic shock								
0734 30-day all cause risk-standardized percutaneous coronary intervention (PCI) mortality rate for patients with ST segment elevation myocardial infarction (STEMI) or cardiogenic shock		Х		Х				
0355 Bilateral cardiac catheterization rate (IQI 25)					Х			
Cardiac Imaging								
0669 Cardiac imaging for preoperative risk assessment for non-cardiac low-risk surgery		Х						
0670 Cardiac stress imaging not meeting appropriate use criteria: preoperative evaluation in low risk surgery patients					Х			
0671 Cardiac stress imaging not meeting appropriate use criteria: routine testing after					Х			

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percutaneous coronary interventions (PCI)								
0672 Cardiac stress imaging not meeting appropriate use criteria: testing in asymptomatic, low risk patients					Х			
Cardiac Rehabilitation								
0642 Cardiac rehabilitation patient referral from an inpatient setting		Х						
0643 Cardiac rehabilitation patient referral from an outpatient setting		Х						
Atrial Fibrillation								
0600 New atrial fibrillation: Thyroid function test		X						
1524 Assessment of thromboembolic risk factors (CHADS 2)		X		Х				
1525 Chronic anticoagulation therapy		Х		Х				
0624 Atrial fibrillation—warfarin therapy		Х		Х				
0578 Ambulatory initiated amiodarone		Х		Χ				

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Cardiovascular TOPIC AREA	including across settings, disciplines, populations, & combinations Composites Emphasis on longitudinal measures over specified timeframe (e.g., 30 day, etc.)	including communication, pre-, intra-, post-op care; secondary prevention; intermediate outcomes, recovery & rehabilitation, end-of-life care	including prevention, healthfullifestyle behaviors, populations at risk & disparities (age, race/ ethnicity, gender, geographic & socio-economic)	including morbidity & mortality, complications of care, prevention of adverse events including HAI, medication adherence/use	including cost/efficiency direct/indirect cost, overuse, underuse, appropriateness	including access, timeliness	including Patient/ Family Engagement (perspective, knowledge, understanding, motivation, risk behavior), Satisfaction, Self- management, Shared Decision making	including Health Status (quality of life, functional status, productivity, burden on patient & family)
therapy: TSH test								
ICD Implants								
1522 ACE/ARB therapy at discharge for ICD implant patients with LVSD		Х		Х				
1528 Beta blocker at discharge for ICD implant patients with a previous MI		Х		Χ				
1529 Beta blocker at discharge for ICD implant patients with LVSD		Х		Х				
0965 Therapy with ACE/ARB and beta blocker at discharge following ICD implantation		Х		Х				
0694 Hospital risk-standardized complication rate following implantation of implantable cardioverter-defibrillator (ICD)	Х	Х		Х				
Hypertension								
0605 Patient(s) that had a serum creatinine in last 12 reported months.		Х						
0018 Controlling high blood pressure		Х	Х	Х				
Heart Failure—Hospital								
0135 Evaluation of left ventricular systolic dysfunction		Х						
0162 ACEI or ARB for left ventricular		Χ		Χ				

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0358 Congestive heart failure mortality (IQI 16) (risk adjusted)	,	Х		Х				
0229 Hospital 30-day, all-cause, risk standardized mortality rate (RSMR) following heart failure hospitalization		Х		Х				
0330 30-day, all-cause risk standardized readmission rate following heart failure hospitalization (risk adjusted)		Х	Х	Х				
0330 30-day all-cause risk standardized readmission rate following heart failure hospitalization (risk adjusted)		X		Х				
0699 30-day post-hospital HF discharge care transition composite measure		Х		Х				
Heart Failure—Outpatient								
0521 Heart failure symptoms addressed		Х						
0079 Heart failure: left ventricular ejection fraction assessment (outpatient setting)		Х						
0081 Heart failure: ACEI or ARB therapy for		Х		Х				

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left ventricular systolic dysfunction								
0610 Heart failure—use of ACE inhibitor (ACEI) or angiotensin receptor blocker (ARB) therapy		Х		Х				
0083 Heart failure: Beta-blocker therapy for left ventricular systolic dysfunction		X		Х				
0615 Heart failure—use of beta blocker Therapy		X		X				