## NATIONAL QUALITY FORUM

#### Measure Evaluation 4.1 December 2009

This form contains the measure information submitted by stewards. Blank fields indicate no information was provided. Attachments also may have been submitted and are provided to reviewers. The subcriteria and most of the footnotes from the <u>evaluation criteria</u> are provided in Word comments within the form and will appear if your cursor is over the highlighted area. Hyperlinks to the evaluation criteria and ratings are provided in each section.

**TAP/Workgroup** (if utilized): Complete all yellow highlighted areas of the form. Evaluate the extent to which each subcriterion is met. Based on your evaluation, summarize the strengths and weaknesses in each section.

Note: If there is no TAP or workgroup, the SC also evaluates the subcriteria (yellow highlighted areas).

**Steering Committee:** Complete all **pink** highlighted areas of the form. Review the workgroup/TAP assessment of the subcriteria, noting any areas of disagreement; then evaluate the extent to which each major criterion is met; and finally, indicate your recommendation for the endorsement. Provide the rationale for your ratings.

Evaluation ratings of the extent to which the criteria are met

C = Completely (unquestionably demonstrated to meet the criterion)

P = Partially (demonstrated to partially meet the criterion)

M = Minimally (addressed BUT demonstrated to only minimally meet the criterion)

N = Not at all (NOT addressed; OR incorrectly addressed; OR demonstrated to NOT meet the criterion)

NA = Not applicable (only an option for a few subcriteria as indicated)

(for NQF staff use) NQF Review #: 1541 NQF Project: Surgery Endorsement Maintenance 2010

### MEASURE DESCRIPTIVE INFORMATION

De.1 Measure Title: Blood Administration Documentation

**De.2 Brief description of measure:** Percentage of transfused units/doses (bags) of RBCs, plasma or platelets with documentation for all of the following:

1. Patient identification (ID) and transfusion order (Blood ID Number) confirmed prior to the initiation of blood 2. Date and time of transfusion

3. Blood pressure, pulse and temperature recorded pre, during and post transfusion

1.1-2 Type of Measure: Process

**De.3 If included in a composite or paired with another measure, please identify composite or paired measure** PBM-05 is a part of the Patient Blood Management (PBM) measure set: PBM-01 (Transfusion Consent), PBM-02 (RBC Transfusion Indication), PBM-03 (Plasma Transfusion Indication), PBM-04 (Platelet Transfusion Indication), PBM-06 (Preoperative Anemia Screening), PBM-07(Preoperative Blood Type Testing and Antibody Screening)

**De.4 National Priority Partners Priority Area:** Patient and family engagement, Care coordination, Safety **De.5 IOM Quality Domain:** Effectiveness, Patient-centered, Safety **De.6 Consumer Care Need:** Getting better, Living with illness

#### CONDITIONS FOR CONSIDERATION BY NQF

Four conditions must be met before proposed measures may be considered and evaluated for suitability as voluntary consensus standards:	NQF Staff
<ul> <li>A. The measure is in the public domain or an intellectual property (measure steward agreement) is signed. Public domain only applies to governmental organizations. All non-government organizations must sign a measure steward agreement even if measures are made publicly and freely available.</li> <li>A.1 Do you attest that the measure steward holds intellectual property rights to the measure and the right to use aspects of the measure owned by another entity (e.g., risk model, code set)? Yes A.2 Indicate if Proprietary Measure (as defined in measure steward agreement):</li> </ul>	A Y□ N□

<ul> <li>A.3 Measure Steward Agreement: Agreement will be signed and submitted prior to or at the time of measure submission</li> <li>A.4 Measure Steward Agreement attached:</li> </ul>	
<b>B.</b> The measure owner/steward verifies there is an identified responsible entity and process to maintain and update the measure on a schedule that is commensurate with the rate of clinical innovation, but at least every 3 years. Yes, information provided in contact section	B Y N
<ul> <li>C. The intended use of the measure includes <u>both</u> public reporting <u>and</u> quality improvement.</li> <li>▶ Purpose: Public reporting, Internal quality improvement Accreditation</li> </ul>	C Y N
<ul> <li>D. The requested measure submission information is complete. Generally, measures should be fully developed and tested so that all the evaluation criteria have been addressed and information needed to evaluate the measure is provided. Measures that have not been tested are only potentially eligible for a time-limited endorsement and in that case, measure owners must verify that testing will be completed within 12 months of endorsement.</li> <li>D.1Testing: Yes, fully developed and tested</li> <li>D.2 Have NQF-endorsed measures been reviewed to identify if there are similar or related measures? Yes</li> </ul>	D Y N
(for NQF staff use) Have all conditions for consideration been met? Staff Notes to Steward ( <i>if submission returned</i> ):	Met Y N
Staff Notes to Reviewers (issues or questions regarding any criteria):	
Staff Reviewer Name(s):	

TAP/Workgroup Reviewer Name:	l.
Steering Committee Reviewer Name:	
1. IMPORTANCE TO MEASURE AND REPORT	
Extent to which the specific measure focus is important to making significant gains in health care quality (safety, timeliness, effectiveness, efficiency, equity, patient-centeredness) and improving health outcomes for a specific high impact aspect of healthcare where there is variation in or overall poor performance. <i>Measures must be judged to be important to measure and report in order to be evaluated against the remaining criteria</i> . (evaluation criteria) 1a. High Impact	<u>Eval</u> <u>Rating</u>
(for NQF staff use) Specific NPP goal:	
<ul> <li>1a.1 Demonstrated High Impact Aspect of Healthcare: Affects large numbers, Frequently performed procedure, Leading cause of morbidity/mortality, Patient/societal consequences of poor quality</li> <li>1a.2</li> </ul>	
<b>1a.3 Summary of Evidence of High Impact:</b> Since the majority of blood is transfused in hospitals, each patient who receives blood should expect that the correct type will be transfused only when required based on an evidence-based clinical indication. Accurate identification of the patient and monitoring during the transfusion is also vital to ensure patient safety. Transfusion processes need to be monitored and reported because the most serious risk of transfusion could be potentially avoidable human errors due to the complexity of the transfusion process of blood administration within the healthcare organization.	
<b>1a.4 Citations for Evidence of High Impact:</b> Whitsett CF, Robichaux MG. Assessment of blood administration procedures: problems identified by direct observation and administrative incident reporting. Transfusion. 2001;41:581-86. Saxena S, Ramer L, Shulman IA. A comprehensive assessment program to improve blood-administering practices using the FOCUS-PDCA model. Transfusion. 2004; 44:1350-56.	1a C P M N

Novis DA, Miller KA, Howanitz PJ, Renner SW, Walsh MK; College of American Pathologists. Audit of transfusion procedures in 660 hospitals. A College of American Pathologists Q-Probes study of patient identification and vital sign monitoring frequencies in 16494 transfusions. Arch Pathol Lab Med. 2003;127:541-8. Roback JD, ed. Technical manual. 16th ed, Bethseda, MD: AABB, 2008.	
The Joint Commission: Comprehensive Accreditation Manual for Hospitals, 2009. Oakbrook Terrace, IL; Joint Commission Resources, Inc., 2009.	
The Joint Commission, "National Patient Safety Goals (NPSG)", IN: Comprehensive accreditation manual for hospitals, 2009. Oakbrook Terrace, IL; Joint Commission Resources, Inc., 2009, pp. NPSG 1 - NPSG 4. AABB Primer of Blood Administration. Revised August 2008. Bethseda, Maryland. [Available at http://www.aabb.org/Content/Professional_Development/Education_and_Training_Material/edtrain.htm#2 (accessed November 2009).]	
1b. Opportunity for Improvement	
<b>1b.1 Benefits (improvements in quality) envisioned by use of this measure:</b> Variation in the practice of administration of blood is becoming increasingly evident from both local and international reports. Studies have shown that there are opportunities for error at number of crucial points in the transfusion process starting with the decision to transfuse, prescribe and request, patient sampling, pre-transfusion testing and the process of actually administering the blood to the patient. Many errors go unnoticed or are underreported so the actual rate of mistransfusion is unknown, but recent reports from hemovigilance systems indicate that errors from the initial recipient identification to final blood administration occur with a frequency of 1 in 1000 events. About two-thirds of errors are associated with incorrect patient identification at the bedside. This measure is needed to standardize and document the processes of blood administration can be used to audit aspects of the transfusion, and the cause of serious adverse events can be adequately investigated.	
1b.2 Summary of data demonstrating performance gap (variation or overall poor performance) across	
<b>providers:</b> The World Health Organization noted that throughout the health-care industry, the failure to identify patients correctly continues to result in transfusion errors. Patient misidentification was cited in more than 100 individual root case analysis report by the United States Department of Veterans Affairs (VA) National Center for Patient Safety from January 2000 to March 2003. Patient misidentification has also been identified as a root cause for many errors by the Joint Commission and has been recognized as an issue that has been addressed as a National Patient Safety Goal since 2003. Administering the wrong type of blood (ABO incompatibility) is the most serious error resulting from a transfusion. Many of the incidents are due to failure of the final identity check carried out between the patient and the blood to be transfused. A national Japanese study found that 20% of 115 surveyed hospitals experienced ABO mismatched transfusions. The main causes of errors were misidentification of blood bags (42.8%), incorrect blood typing (15.1%) and failure to identify the patient (42.1%). A 2003 College of	
American Pathologists (CAP) Q-probe surveyed documentation practices for transfusion that included patient/unit verification and vital sign recording. Patient/unit identification was completed in only 25.4% of the transfusion events. Vital signs were documented 88.3% at all three required times.	
<b>1b.3 Citations for data on performance gap:</b> Transfusion Today (2006) 60:4-7. Murakami J. Rinsho Byori (2003) Jan;51 (1):43-9.	
Novis DA, Miller KA, Howanitz PJ, et al. Audit of transfusion procedures in 660 hospitals: A college of American Pathologists Q-probe study of patient identification and vital sign monitoring frequencies in 16,494 transfusions Arch Pathol Lab Med 2003;127:541-8.	
Mannos D. NCPS patient misidentification study: a summary of root cause analyses. VA NCPS Topics in Patient Safety. Washington, DC, United States Department of Veteran Affairs, June- July 2003 Available at http://www.va.gov/ncps/TIPS/Docs/_TIPS_Jul03.doc Stainsby D, Russell J, Cohen H, et al. Reducing adverse events in blood transfusions. Br J Haematol	
2008;131(1):8-12. SHOT group analyzed 226 cases if ABO-incompatible transfusions and found that the most frequent error was failure of the pretransfusion verification at the bedside.	1b C P M
ABO-incompatible red blood cell transfusion occurs in 1:27,000 to 1:135,207 transfusions with a fatality rate	N

transmission of a viral infection during transfusion. Inden JV, Wagner K, Voytovich AE, Sheehan J, Transfusion errors in New York State: An analysis of 10 pears' experience. Transfusion 2000;40:1207-13. bible J, Urbanik SJ. Comparing near misses with actual mistransfusion events: A more accurate reflection of transfusion incident reports form 1994 - 1998. Transfusion 2002;42:1356-64. Sodonaugh LT. Risks of blood transfusion. Crit Care Med 2003;31:5678-86. Charanol J, Legrand D, Dettori J, Ferrea V. Analysis of ABO discrepancies occurring in 35 French hospitals. Fransfusion 2004;44:860-4. Williamson LM, Lowe SJ, Love EM, Cohen H, Soldan K, et al. Serious hazards of transfusion (SHOT) initiative: Analysis of the Oficsrepancies occurring in 35 French hospitals. Fransfusion 2004;44:860-4. Williamson LM, Lowe SJ, Love EM, Cohen H, Soldan K, et al. Serious hazards of transfusion (SHOT) initiative: Analysis of the Oficsrepancies occurring in 35 French hospitals. Fransfusion 2004;44:860-4. Williamson LM, Lowe SJ, Love EM, Cohen H, Soldan K, et al. Serious hazards of transfusion (SHOT) initiative: Analysis of the Oficsrepancies occurring in 35 French hospitals. Fransfusion 2004;44:860-4. Williamson LM, Lowe SJ, Love EM, Cohen H, Soldan K, et al. Serious hazards of transfusion SHOT) initiative: Analysis of the Oficsrepancies occurring in 35 French hospitals. Biothy JS, Saxena S, Nelson J, Shulman I. Safe handling and administration of blood components: Review of practical concepts. Arch Pathol Lab Med May 2007;131:690-694. To Autocome or Evidence to Support Measure Focus Ic. Outcome or Evidence to Support Measure Focus Ic. Outcome or Evidence (Sfor non-outcome measures, briefly describe the relationship to desired outcome. For non-outcome measures, briefly describe the relationship to desired outcome. For outcomes, describe why It is relevant to the target population; Blood transfusions can lead to as signif		1 #1J41
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Drik WH, Murphy MF, Andreu G, et al. Biomedical Excellence for Safer Transfusion (BEST) Workpatient sample collection. Yox Sang 2003;85:40-47.         Sazama K: Reports of 355 transfusion associated deaths: 1976 through 1985. Transfusion 1990;30:583-590.         Whitaker BJ, Green J, King MR, et al. The 2007 national blood collection and utilization survey report.         Washington, DC: U.S. Department of Health & Human Services: 2008.         1c.9 Quote the Specific guideline recommendation (including guideline number and/or page number):         Note: Recommendations are not numbered or graded in the on-line guideline.         1. Verify the identity of the patient (p.2)         2. Before starting a transfusion check the patient's vital signs (i.e., blood pressure, pulse and temperature (p.3)         3. Record the start and end time of the blood product transfusion (p.4)         1. C-10 Clinical Practice Guideline Clitation: Guideline below:         Finnish Medical Society Duodecim. Blood transfusion: indications and administration messure, but Infusion Nurses Society. Indusion nursing Standards of Practice. Jnin Wiley & sons; 2008 Jan 10 (Various)         There are no formal US guidelines on which to base the blood administration messure, but Infusion Nurses Society. Indusion nursing standards of Practice. J Infus Nurs 2006. Jan-Feb;29(1 Supp):51-92.         1.1.1 National Guideline Clearinghouse or other URL: http://www.guideline.gov/content.aspXid=12787/Beserch-transfusions         1.5.12 Rating of strength of recommendation ( <i>Id Glifferent from USPSTF system</i> , also describe rating and by whorm):         GRADE (Grading	NQF	<sup>-</sup> #1541
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Measure and Report?       1         Steering Committee: Was the threshold criterion, Importance to Measure and Report, met?       1         Rationale:       Y_         N_       N_	<b>1c.14 Rationale for using this guideline over others:</b> This guideline captures the majority of the criteria evaluated in this measure and the recommendations are	
Rationale: Y		1
2. SCIENTIFIC ACCEPTABILITY OF MEASURE PROPERTIES		1 Y N
	2. SCIENTIFIC ACCEPTABILITY OF MEASURE PROPERTIES	
		<u>Eval</u> <u>Rating</u>

### NQF #1541

	F #1541
2a. MEASURE SPECIFICATIONS	
S.1 Do you have a web page where current detailed measure specifications can be obtained? S.2 If yes, provide web page URL:	
2a. Precisely Specified	
<ul> <li>2a.1 Numerator Statement (Brief, text description of the numerator - what is being measured about the target population, e.g. target condition, event, or outcome):</li> <li>Number of transfusion units or doses with documentation for all of the following:</li> <li>1. Patient identification (ID) and transfusion order (Blood Identification (ID) Number) confirmed prior to the initiation of blood</li> <li>2. Date and time of transfusion</li> <li>3. Blood pressure, pulse and temperature recorded pre, during and post transfusion</li> </ul>	
<b>2a.2 Numerator Time Window (</b> <i>The time period in which cases are eligible for inclusion in the numerator</i> <b>):</b> Episode of care	
<b>2a.3 Numerator Details (</b> <i>All information required to collect/calculate the numerator, including all codes, logic, and definitions</i> <b>):</b> The units in the numerator are a subset of the denominator units. The following data elements are collected for the numerator; Blood ID Number, Patient ID Verification, Plasma ID, Platelet ID, RBC ID, Transfusion Order, Transfusion Start Date, Transfusion Start Time and Vital Sign Monitoring. Detailed descriptions are provided in attachment for Section 2a.30.	
<b>2a.4 Denominator Statement</b> (Brief, text description of the denominator - target population being measured): Number of transfused red blood cells, plasma and platelet units/doses evaluated	
2a.5 Target population gender: Female, Male 2a.6 Target population age range: All ages	
<b>2a.7 Denominator Time Window (</b> <i>The time period in which cases are eligible for inclusion in the denominator</i> <b>):</b> Episode of care	
<b>2a.8 Denominator Details (</b> <i>All information required to collect/calculate the denominator - the target population being measured - including all codes, logic, and definitions</i> <b>):</b> Admission Date	
Birthdate ICD-9-CM Principal and Other Procedures RBC Transfusion Exclusions Detailed descriptions are provided in the attachment for Section 2a.30.	
<b>2a.9 Denominator Exclusions (Brief text description of exclusions from the target population):</b> Units associated with documentation of massive transfusion protocol (MTP) or hemorrhagic shock Uncrossmatched units of RBCs RBC units used to prime pumps	
<b>2a.10 Denominator Exclusion Details (</b> <i>All information required to collect exclusions to the denominator, including all codes, logic, and definitions</i> <b>):</b> The data element, RBC Transfusion Exclusions, is used to exclude units that are administered in an 'emergency' situation when blood is transfused using different processes (more than one unit is being transfused or administered very rapidly), for the transfusion of any uncrossmatched units administered for an emergency situation or for RBC units used to prime a pump for surgery and not administered directly to the patient via an intravenous line. The data element definition is; Documentation that the transfused red blood cell (RBC) unit was administered for a massive transfusion protocol (MTP), was an uncrossmatched unit administered for an 'emergency' situation or was used to prime a pump.	2a- specs C P M
2a.11 Stratification Details/Variables (All information required to stratify the measure including the	N

stratification variables, all codes, logic, and definitions): This measure could be stratified using the data element Blood Administration Location. The definition is the location where the blood transfusion started. Allowable values are: Intraoperative Surgery or Nonintraoperative Setting.

2a.12-13 Risk Adjustment Type: No risk adjustment necessary

**2a.14 Risk Adjustment Methodology/Variables** (List risk adjustment variables and describe conceptual models, statistical models, or other aspects of model or method):

2a.15-17 Detailed risk model available Web page URL or attachment:

2a.18-19 Type of Score: Rate/proportion

2a.20 Interpretation of Score: Better quality = Higher score

**2a.21 Calculation Algorithm** (Describe the calculation of the measure as a flowchart or series of steps): Algorithms are provided in attachment for Section 2a.30.

**2a.22 Describe the method for discriminating performance** (*e.g.*, *significance testing*)**:** During the six-month pilot, the distribution of the hospital rates was reviewed over time.

**2a.23 Sampling (Survey) Methodology** *If measure is based on a sample (or survey), provide instructions for obtaining the sample, conducting the survey and guidance on minimum sample size (response rate):* For pilot testing, hospitals were requested to submit 10 cases for each of the three blood products that were discharged from the designated six months. The units submitted for measures PBM-02 - PBM-04 were used for this measure. Post pilot, the sample size will be based on the number of units submitted per discharge month or quarter from the same measures.

Hospitals that choose to sample have the option of sampling quarterly or monthly. A hospital may choose to use a larger sample size than required. Hospitals with an initial population size less than the minimum number of units/doses transfused per quarter/month for the measure, cannot apply sampling to the measure.

**2a.24 Data Source (***Check the source(s) for which the measure is specified and tested***)** Paper medical record/flow-sheet, Electronic administrative data/claims, Lab data

**2a.25** Data source/data collection instrument (*Identify the specific data source/data collection instrument, e.g. name of database, clinical registry, collection instrument, etc.*): The Joint Commission developed a web-based data collection tool that was used by hospitals and for reliability testing during the pilot test. When the measures become part of The Joint Commission's ORYX data collection and reporting program, the data would be collected using contracted Performance Measurement Systems (vendors) that develop data collection tools based on the measure specifications. The tools are verified and tested by Joint Commission staff to confirm the accuracy of the data collection tool with the specifications.

**2a.26-28** Data source/data collection instrument reference web page URL or attachment: Attachment The\_Patient Blood\_Management\_Tool [1]-634279148888089574.pdf

2a.29-31 Data dictionary/code table web page URL or attachment: Attachment PBMSpecifications.pdf

**2a.32-35 Level of Measurement/Analysis** (Check the level(s) for which the measure is specified and tested)

Facility/Agency, Can be measured at all levels

**2a.36-37 Care Settings (***Check the setting(s) for which the measure is specified and tested)* Hospital

**2a.38-41 Clinical Services** (Healthcare services being measured, check all that apply) Clinicians: Nurses, Clinicians: PA/NP/Advanced Practice Nurse, Clinicians: Physicians (MD/DO)

TESTING/ANALYSIS

### 2b. Reliability testing

**2b.1 Data/sample** (description of data/sample and size): A sample of 194 medical records were reabstracted at 12 randomly selected pilot hospitals July through September 2010.

#### **2b.2** Analytic Method (type of reliability & rationale, method for testing):

Hospitals for reliability testing were randomly selected based on multiple characteristics, including region (west, south, north central, northeast), hospital type (teaching/non-teaching, rural/urban), and bed size (0-99, 100-199, 200-299, 300+). The objectives of the reliability site visits included: evaluation of the reliability of the individual measures and associated data elements, assessment of data collection effort including abstraction time and estimated cost, assessment of measure specifications including definitions, abstraction guidelines, etc. and assessment of sampling strategies. To prepare for the reliability site visits, the data collection tool that was used by the pilot hospitals was enhanced and tested. During the reliability site visit, Joint Commission staff re-abstracted a sub-set of records that had been previously submitted by the hospital into the enhanced data collection tool without knowing the measure specific data values that the hospital had submitted. When reabstraction was completed for each record, the results from the hospital and Joint Commission staff were compared and differences adjudicated in the program. Focus group interviews were conducted at each hospital and findings were discussed with each hospital to understand what aspects could be improved. A comparison of calculated indicator rates using data originally abstracted by hospitals and the data that were reabstracted by The Joint Commission staff was adjudicated on each measure and the individual data elements. Statistical analysis utilized Kappa scores and p values.

**2b.3 Testing Results** (reliability statistics, assessment of adequacy in the context of norms for the test conducted):

The number of originally abstracted denominator units was 274 with a computed original measure rate of 89.4%. The number of re-abstracted denominator units was 433 with a re-abstracted measure rate of 67.7%. The absolute difference was 21.7% with a Kappa score of 0.291. The percent of hospital identified population verified as 65%. The match rate for 369 units for the individual data elements was: Patient ID Verification 98.8%, Transfusion Order 92%, Transfusion Start Date 95%, Transfusion Start Time 85% and Vital Sign Monitoring 89%. Measure specifications have been revised to strengthen and provide additional clarity to the data element definitions and abstraction guidelines.

#### 2c. Validity testing

**2c.1 Data/sample** (description of data/sample and size): Face validity was tested by a total of 63 hospitals of various sizes and geographic locations across the country that represented over 300 individuals during August and May 2009. Measure specifications were sent to the test hospitals for review. In addition, on-site focus interviews were conducted at five hospitals. Criterion validity was evaluated during the reliability site visits mentioned above as well as through an online survey that the participating hospitals completed.

#### **2c.2** Analytic Method (type of validity & rationale, method for testing):

The measure information form and the data dictionary were evaluated for face validity. The following parts of the measure information form were evaluated: numerator statement, numerator inclusions, numerator exclusions, denominator statement, denominator inclusions, denominator exclusions and an overall understanding of the measure information form. Each area was scored utilizing a five-point Likert scale. For each data element, the hospitals were asked to comment on the clarity and understanding of the abstraction guidelines and data definitions. In addition, the data dictionary was reviewed for overall understanding, usefulness and clarity utilizing a five-point Likert scale. Qualitative analysis was performed on measure feedback received during the focus group interviews and from the online surveys.

# **2c.3 Testing Results** (statistical results, assessment of adequacy in the context of norms for the test conducted):

A total of 58 hospitals completed the face validity evaluation and rated the overall understanding of the numerator and denominator statements an average 4.4 % that ranked the measure 1st out of the 10 measures. Modifications to improve the understanding and clarity of the measure specifications were made prior to pilot testing based on feedback received from the hospitals during the face validity evaluation. Analysis of the online survey revealed 98% (57/58) of the alpha hospitals recommended moving the measure forward to the pilot test with suggested modifications.



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d.1 Summary of Evidence supporting exclusion(s);       d.2 Citations for Evidence:       d.3 Data/sample (description of data/sample and size);       d.4 Analytic Method (type analysis & rationale);       d.5 Testing Results (e.g., frequency, variability, sensitivity analyses);       M.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N		F #1541
d.2 Citations for Evidence:       2d         d.3 Data/sample (description of data/sample and size):       2d         d.4 Analytic Method (type analysis & rationale):       9         d.5 Testing Results (e.g., frequency, variability, sensitivity analyses):       M         d.5 Testing Results (e.g., frequency, variability, sensitivity analyses):       M         e. Risk Adjustment for Outcomes/ Resource Use Measures       e.1 Data/sample (description of data/sample and size):         e.1 Data/sample (description of data/sample and size):       e.2 Analytic Method (type of risk adjustment, analysis, & rationale):         e.3 Testing Results (risk model performance metrics):       M         a.4 If outcome or resource use measure is not risk adjusted, provide rationale:       NA         Zf. Identification of Meaningful Differences in Performance       H         f.1 Data/sample from Testing or Current Use (description of data/sample and size): All patients > 4       nonths of age that had been selected for measures PBM-02 -PBM-04 from the eligible measure population of opatient discharges from 7/1/09 - 12/31/09 were abstracted. For each patient, all units or doses of bload room each of the three types of bload products were used to measure in purposes.         f.2 Methods to identify statistically significant and practically/meaningfully differences in performance         scores were used to determine hospital measure rates that were significantly differences in erformance;         f.3 Provide Measure Scores from Testing or Current Use (description of scores,	2d. Exclusions Justified	
d.3 Data/sample (description of data/sample and size):       2d         d.4 Analytic Method (type analysis & rationale):       P         d.5 Testing Results (e.g., frequency, variability, sensitivity analyses):       N         a.5 Testing Results (e.g., frequency, variability, sensitivity analyses):       N         e. Risk Adjustment for Outcomes/ Resource Use Measures       e.1         e.1 Data/sample (description of data/sample and size):       e.2         e.3 Testing Results (risk model performance metrics):       M         M.1       M         Z.1 Identification of Meaningful Differences in Performance       N         f.1 Data/sample from Testing or Current Use (description of data/sample and size): All patients > 4       N         g.1 Identification of Meaningful Differences in Performance       N         f.1 Data/sample from Testing or Current Use (description of data/sample and size): All patients > 4       N         nonths of age that had been selected for measures PBM-02 - PBM-04 from the eligible measure population of patient discharges from 7/109 - 12/31/109 were abstraced. For each patient, all units or doses of blood or on each of the three types of blood products were used for measurement purposes.       f.2         f.2 Methods to identify statistically significant and practically/meaningfully differences in performance;       f.3         f.3 Provide Measure Scores from Testing or Current Use (description of scores, e.g., distribution by uartite, mean, median, 50, e.g.; identification of	2d.1 Summary of Evidence supporting exclusion(s):	
d.3 Data/sample (description of data/sample and size):       2d         d.4 Analytic Method (type analysis & rationale):       P         d.5 Testing Results (e.g., frequency, variability, sensitivity analyses):       N         a.5 Testing Results (e.g., frequency, variability, sensitivity analyses):       N         e. Risk Adjustment for Outcomes/ Resource Use Measures       e.1         e.1 Data/sample (description of data/sample and size):       e.2         e.3 Testing Results (risk model performance metrics):       M         M.1       M         Z.1 Identification of Meaningful Differences in Performance       N         f.1 Data/sample from Testing or Current Use (description of data/sample and size): All patients > 4       N         g.1 Identification of Meaningful Differences in Performance       N         f.1 Data/sample from Testing or Current Use (description of data/sample and size): All patients > 4       N         nonths of age that had been selected for measures PBM-02 - PBM-04 from the eligible measure population of patient discharges from 7/109 - 12/31/109 were abstraced. For each patient, all units or doses of blood or on each of the three types of blood products were used for measurement purposes.       f.2         f.2 Methods to identify statistically significant and practically/meaningfully differences in performance;       f.3         f.3 Provide Measure Scores from Testing or Current Use (description of scores, e.g., distribution by uartite, mean, median, 50, e.g.; identification of		
2d       Analytic Method (type analysis & rationale):       2d         d.4 Analytic Method (type analysis & rationale):       2e         d.5 Testing Results (e.g., frequency, variability, sensitivity analyses):       NA         e. Risk Adjustment for Outcomes/ Resource Use Measures       NA         e.1 Data/sample (description of data/sample and size):       e.2 Analytic Method (type of risk adjustment, analysis, & rationale):         e.2 Analytic Method (type of risk adjustment, analysis, & rationale):       2e         e.3 Testing Results (risk model performance metrics):       M         M.       M         e.4 If outcome or resource use measure is not risk adjusted, provide rationale:       NA         2f. Identification of Meaningful Differences in Performance       NA         f.1 Data/sample from Testing or Current Use (description of data/sample and size): All patients > 4       nonths of age that had been selected for measures PBM-02 + PBM-04 from the eligible measure population of patient, all units or doses of blood orom each of the three types of blood products were used for measurement purposes.       f.2 Methods to identify statistically significant and practically/meaningfully differences in performance:         yreage.       Secores from Testing or Current Use (description of scores, e.g., distribution by uartifie, mean, median, SD, etc.; identification of statistically significant and meaningfully differences in reformance; in erformance;         weralk Rate for All Hospitals = 77.2%       Kandard Devisitals = 76.1%	2d.2 Citations for Evidence:	
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verage.   f.3 Provide Measure Scores from Testing or Current Use (description of scores, e.g., distribution by uartile, mean, median, SD, etc.; identification of statistically significant and meaningfully differences in erformance):   Wean Rate for All Hospitals = 76.1%   Wean Rate for All Hospitals = 77.2%   tandard Deviation = 20.7%   Verall Rate for All Hospitals = 81.2%   tin. = 9.0%   lax. = 100%   ower Quartile = 66%   Ipper Quartile = 95%   < -2* = 0	(type of analysis & rationale):	
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Nin. = 9.0% 2f   Nax. = 100% 2f   ower Quartile = 66% P   Ipper Quartile = 95% P   < -2* = 2	Standard Deviation = 20.7% Median Rate for All Hospitals = 81.2%	
ower Quartile = 66% Ipper Quartile = 95% < -2* = 2 < 2** = 0	Min. = 9.0%	26
X-2* = 2 M   X 2** = 0 N   Is Comparability of Multiple Data Sources/Methods   g.1 Data/sample (description of data/sample and size): P   g.2 Analytic Method (type of analysis & rationale): N	Max. = 100% Lower Quartile = 66%	
< 2** = 0 g. Comparability of Multiple Data Sources/Methods g.1 Data/sample (description of data/sample and size): g.2 Analytic Method (type of analysis & rationale): N	Upper Quartile = 95%	P
g.1 Data/sample (description of data/sample and size):       P	Z < -Z = Z Z < 2** = 0	N
g.1 Data/sample (description of data/sample and size):       P	2g. Comparability of Multiple Data Sources/Methods	2g
g.2 Analytic Method (type of analysis & rationale):	<b>2g.1 Data/sample</b> (description of data/sample and size):	P 🗌
	2g.2 Analytic Method (type of analysis & rationale):	N

#### NQF #1541

2g.3 Testing Results (e.g., correlation statistics, comparison of rankings):	
2h. Disparities in Care	
2h.1 If measure is stratified, provide stratified results (scores by stratified categories/cohorts):	2h C□ P□
2h.2 If disparities have been reported/identified, but measure is not specified to detect disparities, provide follow-up plans:	M N NA
TAP/Workgroup: What are the strengths and weaknesses in relation to the subcriteria for Scientific Acceptability of Measure Properties?	2
Steering Committee: Overall, to what extent was the criterion, Scientific Acceptability of Measure Properties, met? Rationale:	2 C P M N
3. USABILITY	
Extent to which intended audiences (e.g., consumers, purchasers, providers, policy makers) can understand the results of the measure and are likely to find them useful for decision making. (evaluation criteria)	<u>Eval</u> <u>Rating</u>
3a. Meaningful, Understandable, and Useful Information	
3a.1 Current Use: Not in use but testing completed	
<b>3a.2 Use in a public reporting initiative (disclosure of performance results to the public at large)</b> ( <i>If used in a public reporting initiative, provide name of initiative(s), locations, Web page URL(s). <u>If not publicly reported</u>, state the plans to achieve public reporting within 3 years): We intend to incorporate these Patient Blood Management measures into our ORYX initiative with associated public reporting on Quality Check when there is a national call for these measures.</i>	
<b>3a.3 If used in other programs/initiatives (</b> <i>If used in quality improvement or other programs/initiatives, name of initiative(s), locations, Web page URL(s). <u>If not used for QI</u>, state the plans to achieve use for QI within 3 years): The specifications will be posted on the Joint Commission website for public use in 2011.</i>	
Testing of Interpretability(Testing that demonstrates the results are understood by the potential users for public reporting and quality improvement)3a.4 Data/sample (description of data/sample and size):	
<b>3a.5 Methods</b> (e.g., focus group, survey, QI project):	3a C□ P□
<b>3a.6 Results</b> (qualitative and/or quantitative results and conclusions):	MN
3b/3c. Relation to other NQF-endorsed measures	
3b.1 NQF # and Title of similar or related measures:	
(for NQF staff use) Notes on similar/related <u>endorsed</u> or submitted measures:	
<ul> <li>3b. Harmonization</li> <li>If this measure is related to measure(s) already <u>endorsed by NQF</u> (e.g., same topic, but different target population/setting/data source <u>or</u> different topic but same target population):</li> <li>3b.2 Are the measure specifications harmonized? If not, why?</li> </ul>	3b C P M M N N NA

### NQF #1541

<ul> <li>3c. Distinctive or Additive Value</li> <li>3c.1 Describe the distinctive, improved, or additive value this measure provides to existing NQF-endorsed measures:</li> <li>5.1 If this measure is similar to measure(s) already endorsed by NQF (i.e., on the same topic and the same target population), Describe why it is a more valid or efficient way to measure quality:</li> </ul>	3c C P M N NA
TAP/Workgroup: What are the strengths and weaknesses in relation to the subcriteria for <i>Usability</i> ?	3
Steering Committee: Overall, to what extent was the criterion, <i>Usability</i> , met? Rationale:	3 C P M N
4. FEASIBILITY	
Extent to which the required data are readily available, retrievable without undue burden, and can be implemented for performance measurement. ( <u>evaluation criteria</u> )	<u>Eval</u> <u>Rating</u>
4a. Data Generated as a Byproduct of Care Processes	
<b>4a.1-2 How are the data elements that are needed to compute measure scores generated?</b> Data generated as byproduct of care processes during care delivery (Data are generated and used by healthcare personnel during the provision of care, e.g., blood pressure, lab value, medical condition), Coding/abstraction performed by someone other than person obtaining original information (E.g., DRG, ICD-9 codes on claims, chart abstraction for quality measure or registry)	4a C P M N
4b. Electronic Sources	
<ul> <li>4b.1 Are all the data elements available electronically? (elements that are needed to compute measure scores are in defined, computer-readable fields, e.g., electronic health record, electronic claims) No</li> <li>4b.2 If not, specify the near-term path to achieve electronic capture by most providers. The project will begin Phase III in January 2011 to retool the specifications for retrieval from an electronic health record.</li> </ul>	4b C M N
4c. Exclusions	
<ul> <li>4c.1 Do the specified exclusions require additional data sources beyond what is required for the numerator and denominator specifications?</li> <li>No</li> <li>4c.2 If yes, provide justification.</li> </ul>	4c C    P    M    NA
4d. Susceptibility to Inaccuracies, Errors, or Unintended Consequences	
4d.1 Identify susceptibility to inaccuracies, errors, or unintended consequences of the measure and describe how these potential problems could be audited. If audited, provide results. None noted during testing	4d C P M N
4e. Data Collection Strategy/Implementation	
4e.1 Describe what you have learned/modified as a result of testing and/or operational use of the measure regarding data collection, availability of data/missing data, timing/frequency of data collection, patient confidentiality, time/cost of data collection, other feasibility/ implementation issues: Abstraction time for PBM-05 varied based on the number of units transfused and the location of the	4e C P M N

transfusion. While not difficult to abstract the information in a non-surgical setting, it was almost impossible to abstract the information intraoperatively with any reliability due to the lack of documentation or illegibility of the paper record. For example, abstracting the data element Transfusion Order was a challenge for the intraoperative setting because many hospitals stated that transfusion orders are not routinely documented in the anesthesia/surgical record. According to many hospital staff, "the doctor that orders the blood does not need to write an order to transfuse the blood during the operation". To address this issue, documentation of a transfusion order or the blood unit identification (ID) number would be acceptable for units transfused intraoperatively. Requiring the blood ID number would provide a way to document the amount of blood a patient received during surgery as many times it was not clear when and how many units/doses of blood were transfused which is essential to track blood use and link to adverse events for national hemovigilence rates. These differences in processes that were noted between the intraoperative and non-intraoperative settings have been addressed by adding the option to stratify the units by intraoperative and non-intraoperative settings so hospitals can determine where to invest their improvement efforts.

This measure, even though it was developed for abstraction at the unit level, was indirectly affected by the difficulty in determining the associated 'event'. Eliminating the abstraction level of an 'event' will also improve the reliability of this measure.

Documentation of pulse, in addition to temperature and blood pressure was added to the vital sign monitoring data element since most hospitals routinely document it during transfusion.

Pilot hospitals were requested to estimate the time to abstract one unit of blood red blood cells (RBCs), platelets or plasma for the six-month pilot which includes the abstraction time for PBM-05. Twenty hospitals estimated an average time of 30 minutes to abstract a unit of blood with an average cost of \$21-25 per hour. However, these costs do not include the time or cost involved in identifying the patient population, staff training or data collection tool instruction. It should also be noted that the learning curve varied widely due to the staff experience and expertise that were utilized for a 'time-limited' project. Due to the amount of time needed to manually abstract the volume of blood transfusions, we believe that these measures are most suitable for abstraction from an electronic medical record (EHR). Retrieval from an EHR could capture 100% of all units that were transfused and would decrease or eliminate the associated abstraction burden. This method would also improve the identification of patients who received blood since procedure codes to document blood use are not standardized across the country. In the meantime, patients can be identified using blood bank records or procedure codes.

During the 12 reliability site visits, two Joint Commission staff also found that the abstraction time varied widely based on the method of record retrieval (e.g., paper record, scanned record or electronic information) at each hospital and the amount of blood transfused per case. Based on hospital feedback, measure specifications have been revised to strengthen and provide additional clarity to data element definitions and abstraction guidelines. The timing and frequency of data collection will remain monthly or quarterly as it does for the other Joint Commission measure sets. Maintaining patient confidentially was not an issue during the pilot test, since blinded hospital and patient identifiers are used on all data received by The Joint Commission staff for data quality reviews.

# **4e.2 Costs to implement the measure** (costs of data collection, fees associated with proprietary measures):

The majority of hospitals already have processes in place to abstract measures if the patients are identified using procedure codes. However, some hospitals document total hospital blood use using blood bank records that would have to be cross-referenced by the patient medical record number to determine how much and the type of blood product each patient received which adds to the abstraction burden. After identifying the patients, the time to collect the data elements for this measure from the operative section of the record would be increased, if available, using manual abstraction.

There are no Joint Commission fees to abstract the measures, but the abstraction cost in addition to the issues mentioned above would depend on the amount of blood products transfused at each hospital since administration documentation is reviewed for all units included in the transfusion measures PBM-02 - PBM-04. Hospitals with Blood Management or conservation programs may have fewer units to review and those with efficient or electronic processes to document blood may have lower abstraction costs.

4e.3 Evidence for costs:

<b>4e.4 Business case documentation:</b> Even though many hospital staff thought that all of the Patient Blood Measures were important, the Blood Administration Documentation measure has been one of the highest ranked measures in all of the testing phases. The lack of clearly written blood transfusion documentation noted in patients who received blood intraoperatively raises the question of how overuse can be determined and addressed if the number of units transfused is not even mentioned in the post-procedure note. Documenting blood use during surgery is essential to tracking transfusion-related adverse events. Improving patient identification during transfusion has been a Joint Commission National Patient Safety Goal #1 for many years, and this measure would be an excellent vehicle to determine if the goal to improve the accuracy of patient identification to eliminate transfusion errors related to misidentification is being achieved. This measure is needed to monitor and evaluate Patient Safety practices, although manual abstraction is very time-consuming and only abstracts a set number of blood products transfused.	
TAP/Workgroup: What are the strengths and weaknesses in relation to the subcriteria for <i>Feasibility</i> ?	4
Steering Committee: Overall, to what extent was the criterion, <i>Feasibility</i> , met? Rationale:	4 C P M N
RECOMMENDATION	
(for NQF staff use) Check if measure is untested and only eligible for time-limited endorsement.	Time- limited
Steering Committee: Do you recommend for endorsement? Comments:	Y N A
CONTACT INFORMATION	
Co.1 Measure Steward (Intellectual Property Owner) Co.1 <u>Organization</u> The Joint Commission, One Renaissance Boulevard, Oakbrook Terrace, Illinois, 60181	
Co.2 Point of Contact Jerod M., Loeb, PhD, jloeb@jointcommission.org, 630-792-5920-	
Measure Developer If different from Measure Steward Co.3 <u>Organization</u> The Joint Commission, One Renaissance Boulevard, Oakbrook Terrace, Illinois, 60181	
Co.4 <u>Point of Contact</u> Harriet, Gammon, MSN, RN, CPHQ, hgammon@jointcommission.org, 630-792-5926-	
Co.5 Submitter If different from Measure Steward POC Harriet, Gammon, MSN, RN, CPHQ, hgammon@jointcommission.org, 630-792-5926-, The Joint Commission	
Co.6 Additional organizations that sponsored/participated in measure development	
ADDITIONAL INFORMATION	
Workgroup/Expert Panel involved in measure development Ad.1 Provide a list of sponsoring organizations and workgroup/panel members' names and organizations. Describe the members' role in measure development. The technical advisory panel determined priority areas in blood management for measure development. They reviewed public comments and were actively involved in all phases of the project to identify and develop the specifications. Measure recommendations for National Quality Forum endorsement were made after careful r of the pilot results and site feedback.	

Ad.2 If adapted, provide name of original measure: Ad.3-5 If adapted, provide original specifications URL or attachment

Measure Developer/Steward Updates and Ongoing Maintenance

Ad.6 Year the measure was first released:

Ad.7 Month and Year of most recent revision: 12, 2010

Ad.8 What is your frequency for review/update of this measure? Biannually

Ad.9 When is the next scheduled review/update for this measure? 06, 2011

Ad.10 Copyright statement/disclaimers: No royalty or use fee is required for copying or reprinting this manual, but the following are required as a condition of usage: 1) disclosure that the Specifications Manual is periodically updated, and that the version being copied or reprinted may not be up-to-date when used unless the copier or printer has verified the version to be up-to-date and affirms that, and 2) users participating in Joint Commission accreditation, including performance measures systems, are required to update their software and associated documentation based on the published manual production timelines.

Example Acknowledgement: The Specifications Manual for National Hospital Inpatient Quality Measures Patient Blood Management Performance Measure Set is periodically updated by The Joint Commission. Users of the Specifications Manual for National Hospital Inpatient Quality Measures Patient Blood Management Performance Measure Set must update their software and associated documentation based on the published manual production timelines.

Ad.11 -13 Additional Information web page URL or attachment: Attachment TAPLISTWEBc-634276558279803714.doc

Date of Submission (MM/DD/YY): 12/29/2010

# **Patient Blood Management (PBM)**

## **Set Measures**

Set Measure ID	Measure Short Name
PBM-01	Transfusion Consent
PBM-02	RBC Transfusion Indication
PBM-03	Plasma Transfusion Indication
PBM-04	Platelet Transfusion Indication
PBM-05	Blood Administration Documentation
PBM-06	Preoperative Anemia Screening
PBM-07	Preoperative Blood Type Testing and Antibody Screening

## **Measure Set Specific Data Elements**

Element Name	Collected For
Admission From Home	<u>PBM-06,</u>
Anesthesia Start Date	<u>PBM-06,</u>
Blood Administration Location	<u>PBM-02, PBM-03, PBM-04, PBM-05,</u>
Blood Bank Records	<u>PBM-01, PBM-02, PBM-03, PBM-04,</u>
	<u>PBM-05</u> ,
Blood ID Number	<u>PBM-05</u> ,
Blood Type Testing Ordered	<u>PBM-07</u> ,
Clinical Indication for Plasma	<u>PBM-03</u> ,
Clinical Indication for Platelets	<u>PBM-04</u> ,
Clinical Indication for RBCs	<u>PBM-02</u> ,
Education Addressed Risks, Benefits and Alternatives to	<u>PBM-01</u> ,
Transfusion	
Patient ID Verification	<u>PBM-05</u> ,
Plasma ID	<u>PBM-03, PBM-05,</u>
Platelet ID	<u>PBM-04,</u> <u>PBM-05,</u>
Pre-transfusion Hematocrit	<u>PBM-02</u> ,
Pre-transfusion Hemoglobin	<u>PBM-02,</u>
Pre-transfusion PT/INR Result	<u>PBM-03</u> ,
Pre-transfusion Platelet Count	<u>PBM-04</u> ,
Preoperative Anemia Screening Date	<u>PBM-06</u> ,
Preoperative Blood Type Testing	<u>PBM-07</u> ,
RBC ID	<u>PBM-02</u> , <u>PBM-05</u> ,
RBC Unit Exclusions	<u>PBM-02</u> , <u>PBM-05</u> ,
Surgery Scheduled Timeframe	<u>PBM-06</u> ,
Transfusion Consent	<u>PBM-01</u> ,
Transfusion Order	<u>PBM-05,</u>
Transfusion Start Date	<u>PBM-05</u> ,
Transfusion Start Time	<u>PBM-05,</u>
Vital Sign Monitoring	<u>PBM-05,</u>

# **Related Materials**

Document Name z. Appendix E - Miscellaneous Tables

# **Measure Information Form**

Measure Set: Patient Blood Management(PBM)

Set Measure ID: PBM-01

### Performance Measure Name: Transfusion Consent

**Description:** Patients with a signed consent who received information about the risks, benefits and alternatives of transfusion prior to the initial blood transfusion or the initial transfusion was deemed a medical emergency.

**Rationale:** Planning a discussion with a licensed practitioner regarding the risks, benefits and alternatives of transfusion is an opportunity for the patient to participate in decisions about his or her care. It is a process that takes into consideration, each patient's preferences, clinical needs and provides information in compliance with the regulations and policies of the state and facility. Even though policies related to informed consent may vary among hospitals, all hospitals require some type of consent prior to treatment unless emergency care is needed. The elements of performance for the Joint Commission Standard RI.01.03.01 related to the informed consent process include a discussion about the risks, benefits and alternatives, and a discussion about the risk, if care is not received. This measure is also supported by the Joint Commission's National Patient Safety Goal (NPSG) 13 that encourages patients' active involvement in their own care as a patient safety strategy.

For many years, the American Association of Blood Banks (AABB) organization has supported the consent process for transfusion and has developed several standards such as AABB Standard 5.19.1. AABB requires that at a minimum, a recipient consent for transfusion and that should include; a description of the risks, benefits and treatment alternatives, the opportunity to ask questions and the right to accept or refuse transfusion.

### Type of Measure: Process

### Improvement Noted As: Increase in the rate

**Numerator Statement:** Patients with a signed consent who received information about the risks, benefits and alternatives prior to the initial blood transfusion or the initial transfusion was deemed a medical emergency

### Included Populations: Not applicable

### Excluded Populations: None

### Data Elements:

- Education Addressed Risks, Benefits and Alternatives to Transfusion
- Transfusion Consent

### Denominator Statement: Patients who received red blood cell, plasma or platelet transfusions

**Included Populations:** Discharges with an ICD-9-CM Principal or Other Procedure Codes for transfusion as defined in Appendix A, Table 9.3-9.6 or a transfusion documented from Blood Bank Records.

### Excluded Populations: None

### **Data Elements:**

- <u>Admission Date</u>
- Blood Bank Records
- Discharge Date
- ICD-9-CM Other Procedure Codes
- ICD-9-CM Principal Procedure Code

### Risk Adjustment: No.

**Data Collection Approach:** Retrospective data collection sources for required data elements include administrative data and medical records. Hospitals that do not use ICD-9-CM procedure codes to document transfusions may use blood bank records to identify the population.

**Data Accuracy:** Variation may exist in the assignment of ICD-9-CM codes and blood bank records; therefore, coding practices and transfusion documentation may require evaluation to ensure consistency.

**Measure Analysis Suggestions:** Hospitals may want to evaluate the cases according to medical or surgical designation that were not included in the numerator in order to determine if the consent was signed and/or if all or only part of the educational components were given or if documentation was insufficient. Based on this information, hospitals may assess the barriers impacting this measure that could be improved.

Sampling: Yes. For additional information see the Population and Sampling Specifications Section.

Data Reported As: Aggregate rate generated from count data reported as a proportion.

### **Selected References:**

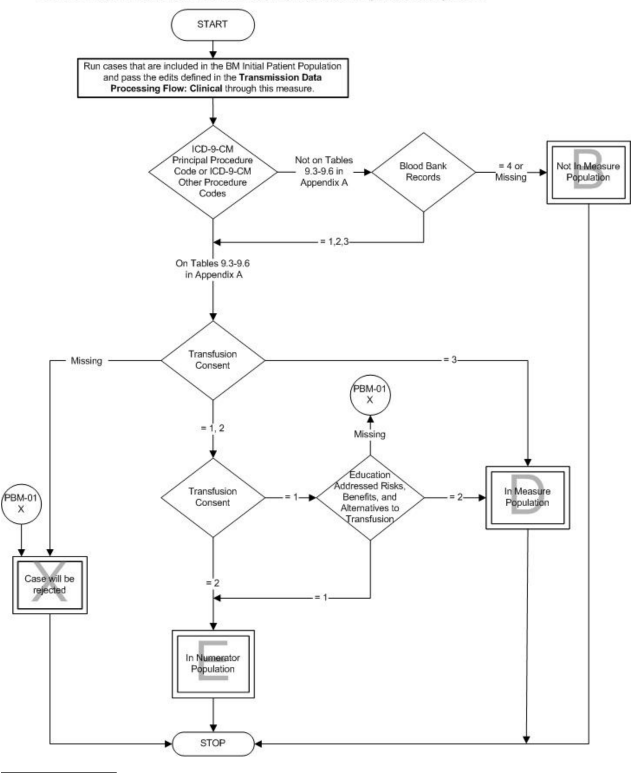
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- Standards for Blood Banks and Transfusion Services, 25th ed. Bethseda, MD: AABB 2008.
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## Measure Algorithm:

### PBM-01: Transfusion Consent

Numerator: Patients with a signed consent who received information about the risks, benefits and alternatives prior to the initial blood transfusion or the initial transfusion was deemed a medical emergency

Denominator: Patients who received red blood cells, platelets or plasma



# **Measure Information Form**

Measure Set: Patient Blood Management(PBM)

Set Measure ID: PBM-02

### Performance Measure Name: RBC Transfusion Indication

**Description:** The number of transfused red blood cell (RBC) units with a pre-transfusion hemoglobin (hgb) or hematocrit (hct) result and clinical indication documented from patients of all ages who received RBCs.

**Rationale:** Improvement of the safety and quality of care that a hospital provides includes the review of the use of blood and blood products. Despite current evidence and best practice guidelines, clinical practice regarding when to transfuse varies among physicians and institutions even though most would agree that blood products should only be given when the benefits outweigh the harm. Many advocate that transfusion decisions should be based on a clinical assessment and not on laboratory values alone to avoid inappropriate over-or-under transfusion. Measuring whether an "indication for transfusion" and a pre-transfusion laboratory value was documented may improve the utilization of blood components. In addition, implementing such a process may simplify the hospital's review for appropriateness of the transfusion when auditing records for accreditation and regulatory agencies. In a study by Friedman and Ebrahim, there was a significant correlation between red blood cell transfusions that lacked documentation of the clinical necessity for transfusion and justification of the transfusion.

### Type of Measure: Process

Improvement Noted As: Increase in the rate

**Numerator Statement:** Number of RBC units with pre-transfusion hemoglobin or hematocrit result and clinical indication documented

Included Populations: Not applicable

Excluded Populations: None

### **Data Elements:**

- Clinical Indication for RBCs
- Pre-transfusion Hematocrit
- Pre-transfusion Hemoglobin
- <u>RBC ID</u>

Denominator Statement: Number of transfused red blood cell units evaluated

### **Included Populations:**

- Discharges with an ICD-9-CM Principal or Other Procedure Codes for transfusion as defined in Appendix A, Tables 9.3 or 9.4 or a RBC transfusion documented from Blood Bank Records.
- The first six RBCs units transfused after hospital arrival

### Excluded Populations: None

### **Data Elements:**

- <u>Admission Date</u>
- Birthdate
- Blood Administration Location
- <u>Blood Bank Records</u>
- <u>Discharge Date</u>
- ICD-9-CM Other Procedure Codes
- ICD-9-CM Principal Procedure Code
- <u>RBC Unit Exclusions</u>

### Risk Adjustment: No.

**Data Collection Approach:** Retrospective data sources for required data elements include administrative/billing data and medical records. Hospitals that do not use ICD-9-CM procedure codes to document transfusions may use blood bank records to identify the population of patients who received RBCs.

**Data Accuracy:** Variation may exist in the assignment of ICD-9-CM codes and blood bank records; therefore, coding practices and transfusion documentation may require evaluation to ensure consistency.

**Measure Analysis Suggestions:** Hospitals may want to use the data to further evaluate the process for determining the need for blood products based on the clinical indications and correlating it with the pre-transfusion value that was documented. This information may assist hospitals to determine if the patients were transfused appropriately or if efforts should be directed toward additional documentation efforts for monitoring blood product usage. Data may be grouped by service designation or by blood products to identify specific areas for staff review.

**Sampling:** Yes. For additional information see the Population and Sampling Specifications Section.

Data Reported As: Aggregate rate generated from count data reported as a proportion.

### Selected References:

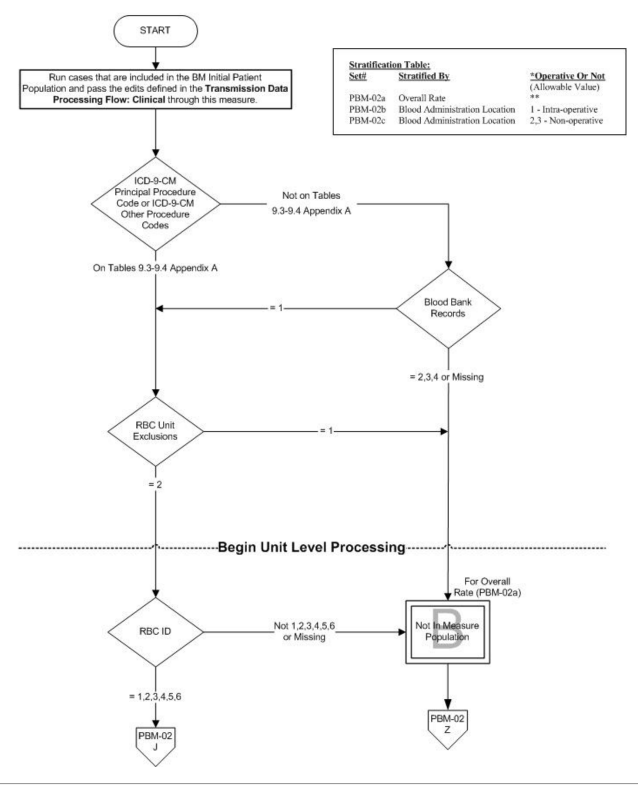
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- Corwin HL, Parsonnet KC, Gettinger A. RBC transfusion in the ICU: is there a reason? Chest. 1995;108: 767-771.
- Tobin SN, Campbell DA, Boyce NW. Durability of response to a targeted intervention to modify clinician transfusion practices in a major teaching hospital. MJA. 2001;174:445-448.
- Clinical practice guideline: Red blood cell transfusion in adult trauma and critical care. Crit Care Med 2009 Vol.37, No.12.

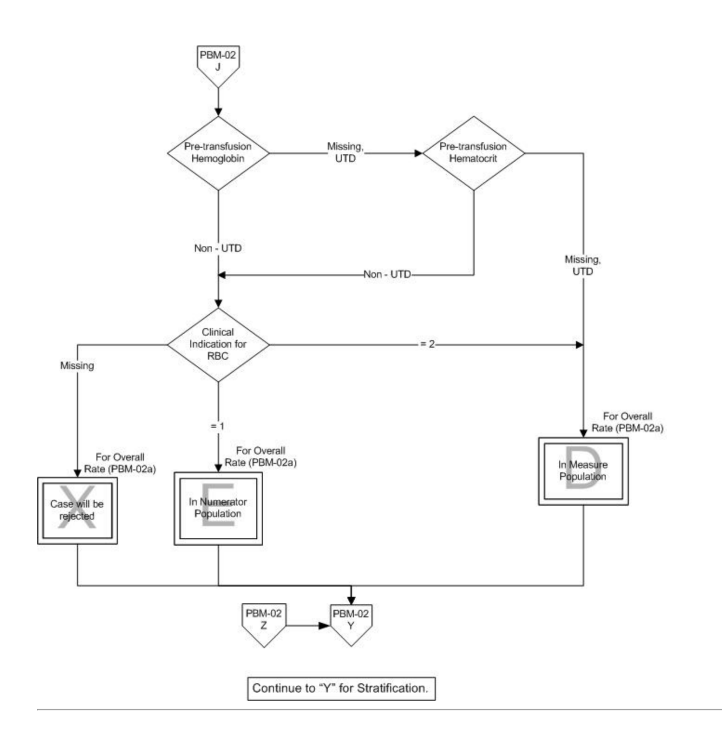
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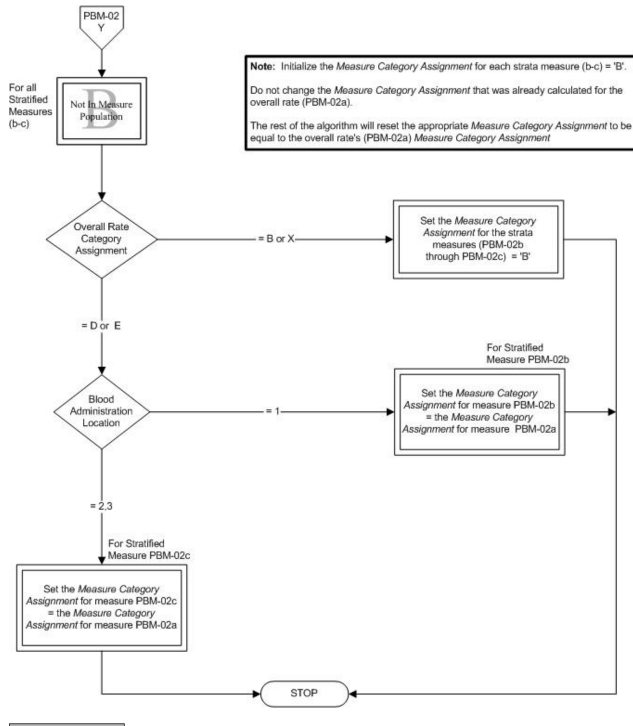
### PBM-02: RBC Transfusion Indication

Numerator: Number of RBC units (bags) with pre-transfusion hemoglobin or hematocrit result and clinical indication documented

Denominator: Number of transfused red blood cell units evaluated







**Related Topics** 

# **Measure Information Form**

Measure Set: Patient Blood Management(PBM)

Set Measure ID: PBM-03

Performance Measure Name: Plasma Transfusion Indication

**Description:** The number of transfused plasma units with a pre-transfusion PT/INR result and clinical indication documented from patients of all ages who received plasma.

**Rationale:** The use of plasma has increased and is disproportionally high compared to other countries with similar levels of health care. Indications for transfusing plasma are very limited, and as a result, published studies often show unjustifiable use of plasma. According to the National Heart Lung and Blood Institute, plasma should be administered only to increase the level of clotting factors in patients with a demonstrated deficiency. If the prothrombin time (PT) and partial thromboplastin time (PTT) are < 1.5 times normal, a plasma transfusion is rarely needed. However, plasma is frequently transfused to patients with mild-to moderate elevations in PT despite numerous studies that have not shown a correlation between the risk of bleeding and mild-to moderate test results. In a study by Wahab et al, transfusion of plasma for mild abnormalities of coagulation values resulted in a partial normalization in a minority of patients, and failed to correct the PT in 99% of the patients. In a 2004 study by Hui, the need to correct prolonged international normalized ratios (INRs) for patients on warfarin emerged as the primary indication for plasma followed by massive transfusions.

## Type of Measure: Process

Improvement Noted As: Increase in the rate

**Numerator Statement:** Number of plasma units with pre-transfusion PT/INR result and clinical indication documented

Included Populations: Not applicable

Excluded Populations: None

### **Data Elements:**

- Clinical Indication for Plasma
- Plasma ID
- Pre-transfusion PT/INR Result

### Denominator Statement: Number of transfused plasma units evaluated

### **Included Populations:**

- Discharges with an ICD-9-CM Principal or Other Procedure Codes for transfusion as defined in Appendix A, Table 9.6 or a plasma transfusion documented from Blood Bank Records
- The first three plasma units transfused from hospital arrival

### **Excluded Populations:**

• Discharges with an ICD-9-CM Principal Diagnosis Code of trauma as defined in Appendix A, Table 9.7.

### **Data Elements:**

- <u>Admission Date</u>
- <u>Birthdate</u>
- Blood Administration Location
- Blood Bank Records
- Discharge Date
- ICD-9-CM Other Procedure Codes
- ICD-9-CM Principal Diagnosis Code
- ICD-9-CM Principal Procedure Code

### Risk Adjustment: No.

**Data Collection Approach:** Retrospective data sources for required data elements include administrative/billing data and medical records. Hospitals that do not use ICD-9-CM procedure codes to document transfusions may use blood bank records to identify the population of patients who received plasma.

**Data Accuracy:** Variation may exist in the assignment of ICD-9-CM codes and blood bank records; therefore, coding practices and transfusion documentation may require evaluation to ensure consistency.

**Measure Analysis Suggestions:** Data from this measure may be used to review the type of invasive procedures or surgeries that use plasma in order to further evaluate appropriateness of use.

Sampling: Yes. For additional information see the Population and Sampling Specifications Section.

Data Reported As: Aggregate rate generated from count data reported as a proportion.

### **Selected References:**

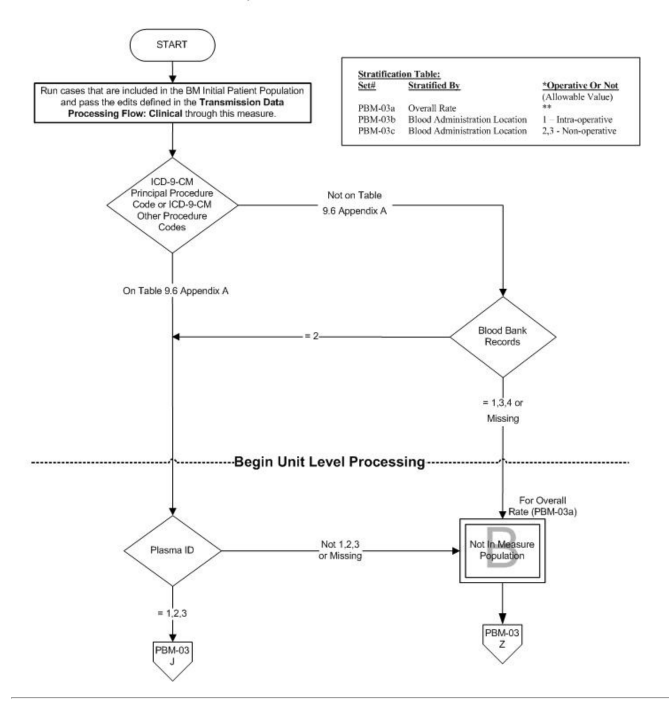
- Hui C, Williams I, Davis K. Clinical audit of the use of fresh-frozen plasma and platelets in a tertiary teaching hospital and the impact of a new transfusion request form. Int Med J. 2005;35:283-288.
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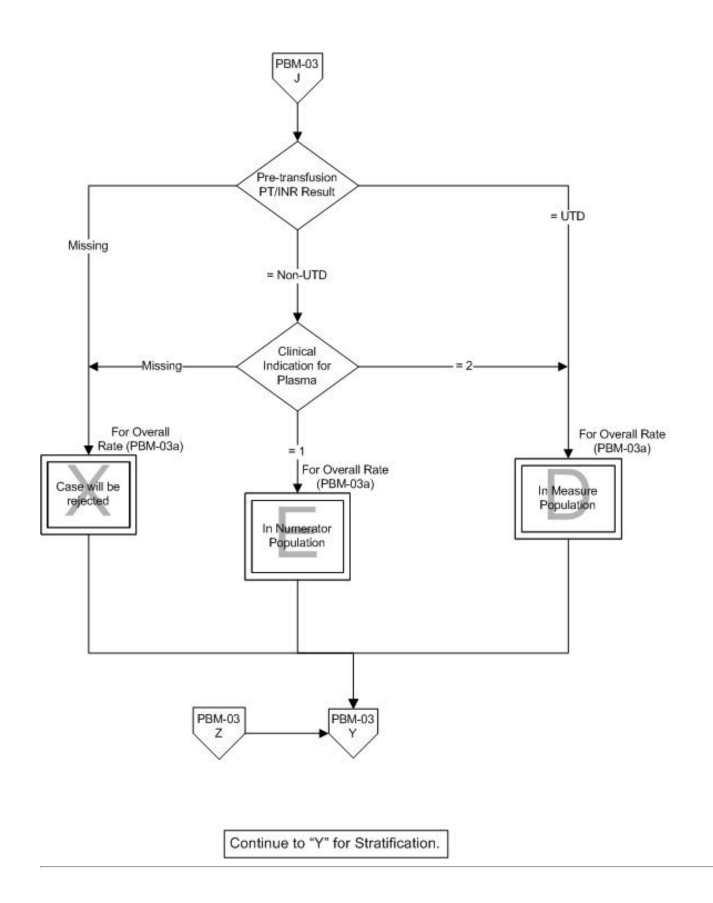
## Measure Algorithm:

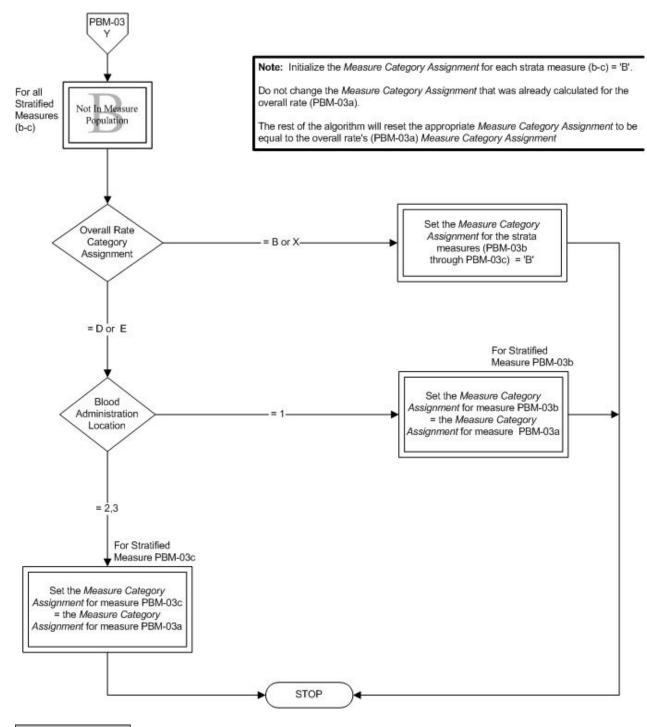
### PBM-03: Plasma Transfusion Indication

Numerator: Number of plasma units with pre-transfusion PT/INR result and clinical indication documented

Denominator: Number of transfused plasma units evaluated







**Related Topics** 

# **Measure Information Form**

Measure Set: Patient Blood Management(PBM)

Set Measure ID: PBM-04

Performance Measure Name: Platelet Transfusion Indication

**Description:** The number of transfused platelet units with pre-transfusion platelet count and clinical indication documented from patients of all ages who received platelets.

**Rationale:** Platelets are transfused to treat or prevent bleeding associated with thrombocytopenia and/or platelet dysfunction. Platelets given therapeutically should help stop the bleeding, and if given prophylactically, post transfusion platelet counts should be obtained to monitor the response to determine the effectiveness of the transfusion. Repeated platelet transfusions can cause alloimmunization and cause platelet refractoriness to future transfusions. Multiple infectious risks are associated with platelet transfusions so patients should only be exposed to the least amount needed.

### Type of Measure: Process

Improvement Noted As: Increase in the rate

**Numerator Statement:** Number of platelet units with pre-transfusion platelet count result and clinical indication documented

Included Populations: Not applicable

Excluded Populations: None

## **Data Elements:**

- Clinical Indication for Platelets
- Platelet ID
- Pre-transfusion Platelet Count

## Denominator Statement: Number of transfused platelet units evaluated

### **Included Populations:**

- Discharges with an ICD-9-CM Principal or Other Procedure Codes for transfusion as defined in Appendix A, Table 9.5 or a platelet transfusion documented from Blood Bank Records
- The first three platelet units transfused after hospital arrival

## Excluded Populations: None

## Data Elements:

- <u>Admission Date</u>
- Blood Administration Location
- Blood Bank Records

- Discharge Date
- ICD-9-CM Other Procedure Codes
- ICD-9-CM Principal Procedure Code

### Risk Adjustment: No.

**Data Collection Approach:** Retrospective data sources for required data elements include administrative/billing data and medical records. Hospitals that do not use ICD-9-CM procedure codes to document transfusions may use blood bank records to identify the population of patients who received platelets.

**Data Accuracy:** Variation may exist in the assignment of ICD-9-CM codes and blood bank records; therefore, coding practices and transfusion documentation may require evaluation to ensure consistency.

**Measure Analysis Suggestions:** Data from this measure may be used to evaluate the utilization and approriateness of platelets used by an organization.

Sampling: Yes. For additional information see the Population and Sampling Specifications.

Data Reported As: Aggregate rate generated from count data reported as a proportion.

### Selected References:

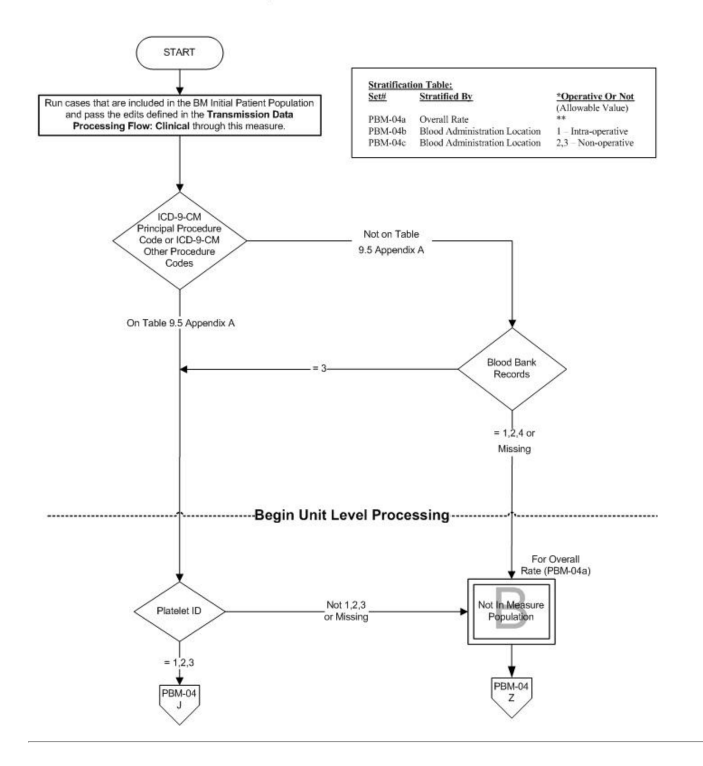
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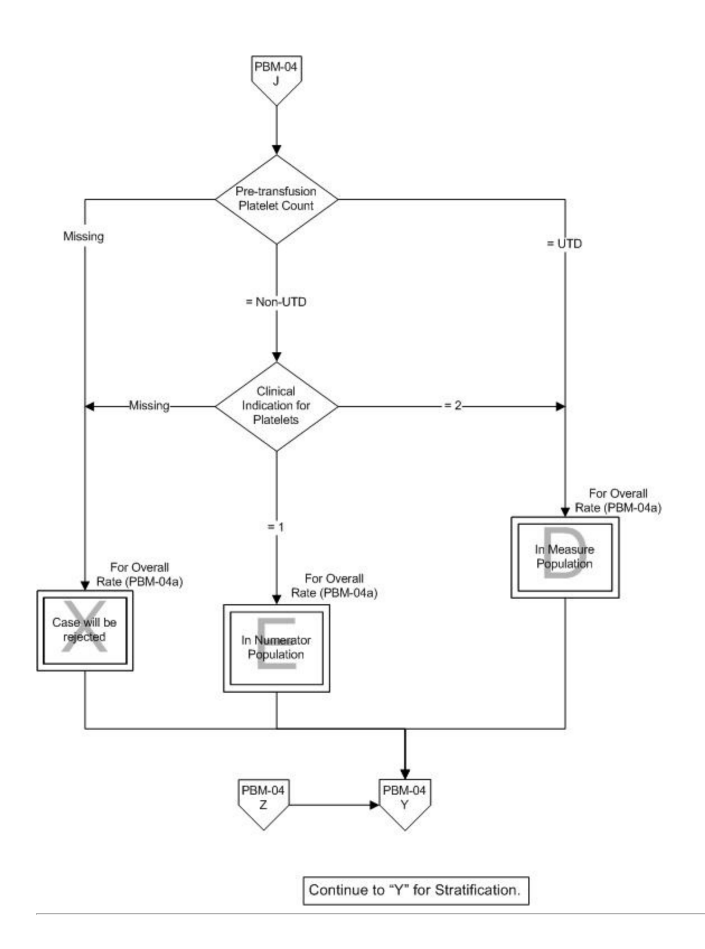
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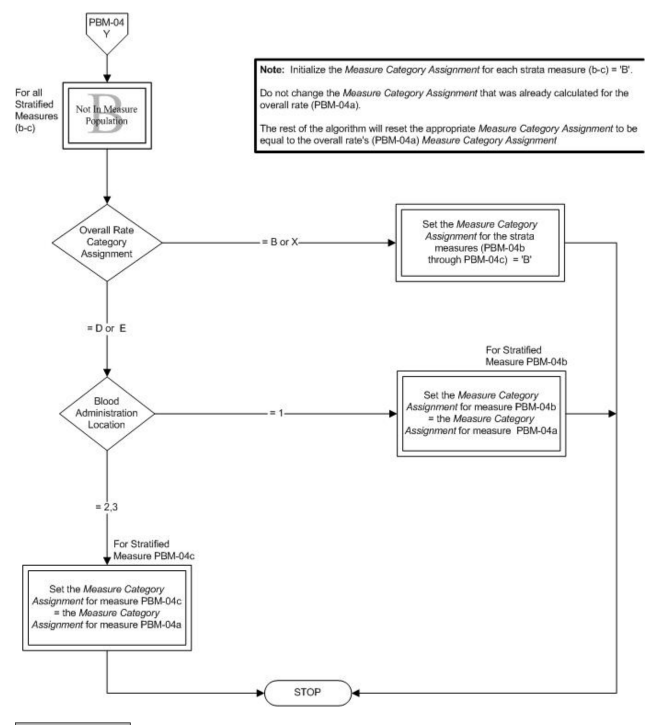
## PBM-04: Platelet Transfusion Indication

Numerator: Number of platelet doses with pre-transfusion platelet count result and clinical indication documented

Denominator: Number of transfused platelet units evaluated







Related Topics

# **Measure Information Form**

Measure Set: Patient Blood Management(PBM)

### Set Measure ID: PBM-05

### Performance Measure Name: Blood Administration Documentation

**Description:** The number of transfused red blood cells, plasma or platelet transfusion units/doses (bags) that had documentation of the following: patient identification and an order to transfuse (Blood ID Number) confirmed prior to the initiation of transfusion, transfusion start date and time, and blood pressure, pulse and temperature recorded at specific intervals.

**Rationale:** Since the majority of blood units are transfused in hospitals, specific policies and procedures have been developed by each hospital to address documentation of blood administration standards in accordance with their state and federal regulations. Though documentation components vary among organizations, identification of the patient and confirmation of the order to transfuse are common indicators used for all blood products since incomplete patient identification could result in an adverse outcome. Prior to administering blood or blood products, patient identification by two identifiers is required by numerous organizations including the AABB Standard 5.19.3, and the Joint Commission National Patient Safety Goal (NPSG) 1. In addition, numerous organizations require or advise that the licensed staff confirm that there is a transfusion order as directed by the AABB Standard 5.19.6 and the elements of performance for the Joint Commission NPSG.01.01.01.

Patient monitoring during the transfusion is an important component related to patient safety. The first 10 to 15 minutes of the transfusion are considered the most critical to assess for a potential transfusion reaction and close observation during this time is recommended in the AABB Primer. Monitoring of vital signs at baseline, during and at the completion of the transfusion in addition to observation are used to assess the patient's condition for any changes.

### Type of Measure: Process

### Improvement Noted As: Increase in the rate

Numerator Statement: Number of units/doses (bags) with documentation for all of the following:

- patient identification and transfusion order (Blood ID Number) confirmed prior to the initiation of transfusion
- transfusion start date and time
- blood pressure, pulse and temperature recorded pre, during and post transfusion

### Included Populations: Not applicable

### Excluded Populations: None

### **Data Elements:**

- Blood ID Number
- Patient ID Verification
- Plasma ID

- Platelet ID
- RBC ID
- Transfusion Order
- <u>Transfusion Start Date</u>
- <u>Transfusion Start Time</u>
- <u>Vital Sign Monitoring</u>

**Denominator Statement:** Number of transfused red blood cells, plasma or platelet units/doses (bags) evaluated

### **Included Populations:**

 Discharges with an ICD-9-CM Principal or Other Procedure Codes for transfusion as defined in Appendix A, Table 9.3-9.6 or a transfusion documented from Blood Bank Records

### **Excluded Populations:**

- Units used in massive transfusion protocols
- Uncrossmatched units
- Units used to prime equipment

### **Data Elements:**

- Admission Date
- Birthdate
- Blood Administration Location
- Blood Bank Records
- Discharge Date
- ICD-9-CM Other Procedure Codes
- ICD-9-CM Principal Procedure Code
- <u>RBC Unit Exclusions</u>

## Risk Adjustment: No.

**Data Collection Approach:** Retrospective data sources for required data elements include administrative/billing data and medical records. Hospitals that do not use ICD-9-CM procedure codes to document transfusions may use blood bank records to identify the population.

**Data Accuracy:** Variation may exist in the assignment of ICD-9-CM codes and blood bank records; therefore, coding practices and transfusion documentation may require evaluation to ensure consistency.

**Measure Analysis Suggestions:** The data from this measure may be used to evaluate the adherence to organizational policies and procedures for blood administration for each of the blood products. Data could be evaluated by unit or service in order to identify areas for staff education. The data could also be used during accreditation surveys to document the hospital's efforts to improve the accuracy of patient identification when administering blood related to the Joint Commission National Patient Safety Goal #1.

Sampling: Yes. For additional information see the Population and Sampling Specifications.

Data Reported As: Aggregate rate generated from count data reported as a proportion.

### **Selected References:**

Patient Blood Management NQF - Do NOT Distribute

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- AABB Primer of Blood Administration. Revised August 2008. Bethseda, Maryland. [Available at

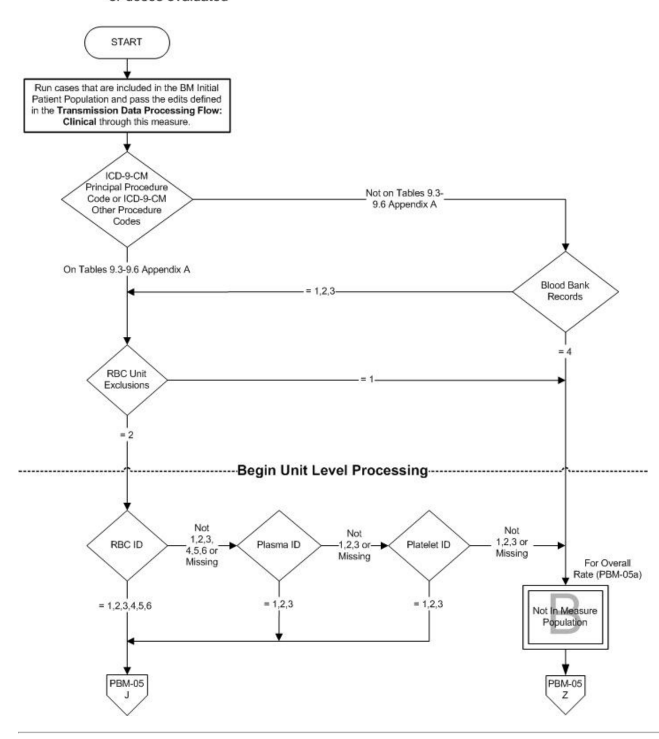
http://www.aabb.org/Content/Professional\_Development/Education\_and\_Training\_Material/edtr (accessed November 2009).]

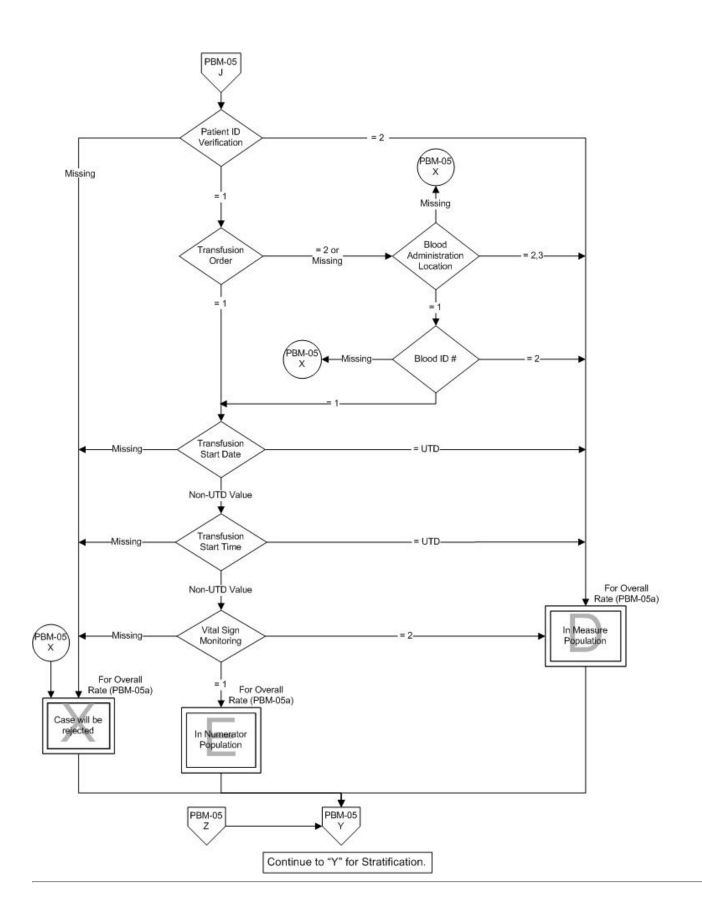
#### Measure Algorithm:

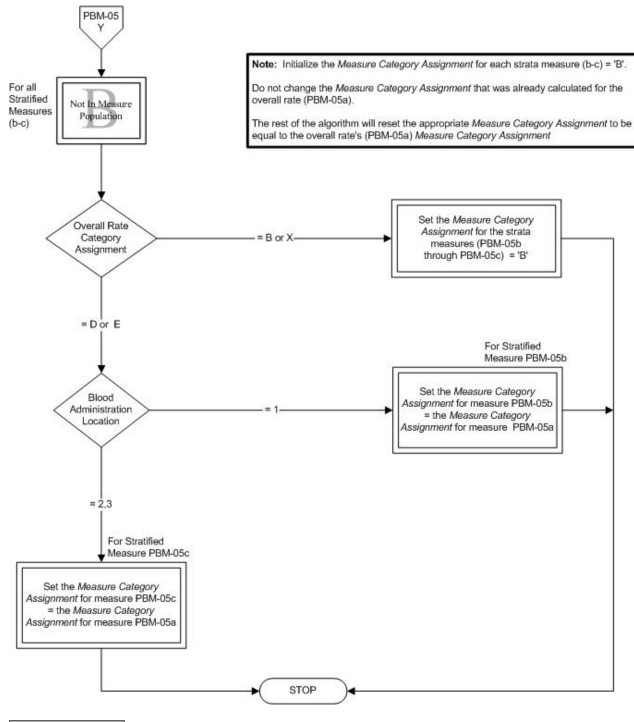
## PBM-05: Blood Administration Documentation

Numerator: Number of blood transfusion units (bags) or doses with documentation for all of the following:

- patient identification (ID) and transfusion order (blood ID number) confirmed prior to the initiation of blood
- · date and time of transfusion
- · blood pressure, pulse and temperature recorded pre, during and post transfusion
- Denominator: Number of transfused red blood cells, plasma and platelet units (bags) or doses evaluated







**Related Topics** 

# **Measure Information Form**

Measure Set: Patient Blood Management(PBM)

### Set Measure ID: PBM-06

### Performance Measure Name: Preoperative Anemia Screening

**Description:** Selected elective orthopedic, cardiac and hysterectomy surgical patients with documentation of preoperative anemia screening date 14 - 45 days before surgery start date for procedures scheduled 14 or more days before surgery.

**Rationale:** Development of formal protocols for preoperative testing of hemoglobin (hgb) for potential high-blood loss elective surgeries could be used to identify and intervene for optimal management of blood resources. Preoperative anemia often goes unrecognized and untreated unless tests are ordered in advance of a planned surgery. Early recognition of anemia offers patients an opportunity to receive the most appropriate transfusion-sparing strategy, and avoid the risk of a potential transfusion. Researchers have shown that preoperative hgb and hematocrit can be used as predictors of outcome for specific types of patients such as cardiac artery bypass graft or orthopedic surgery. In a study by Salido, orthopedic patients with a preoperative hemoglobin <13 g/dL had four times the risk of transfusion than those with a hemoglobin level between 13 g/dL and 15 g/dL.

#### Type of Measure: Process

#### Improvement Noted As: Increase in the rate

**Numerator Statement:** Patients with preoperative anemia screening 14 - 45 days before Anesthesia Start Date

Included Populations: Not applicable

Excluded Populations: None

**Data Elements:** 

• Preoperative Anemia Screening Date

Denominator Statement: Selected elective surgical patients

#### **Included Populations:**

• Discharges with an ICD-9-CM Principal Procedure Codes of selected surgeries as defined in Appendix A, Tables 2.2, 5.01, 5.02, 5.08, 5.11, 5.22, 5.23, 9.1 or 9.2.

## **Excluded Populations:**

- Patients less than 18 years of age
- Patients with surgery scheduled less than 14 days before Anesthesia Start Date
- · Patients not admitted from home

## **Data Elements:**

- Admission Date
- Admission From Home
- Birthdate
- Discharge Date
- ICD-9-CM Principal Procedure Code
- ICD-9-CM Principal Procedure Date
- Surgery Scheduled Timeframe

#### Risk Adjustment: No.

**Data Collection Approach:** Retrospective data sources for required data elements include administrative data and medical records.

**Data Accuracy:** Variation may exist in the assignment of ICD-9-CM codes; therefore, coding practices may require evaluation to ensure consistency.

**Measure Analysis Suggestions:** These data may be used to evaluate specific patient groups at high risk for a blood transfusion that did not have their pre-operative hemoglobin and/or transfusion testing completed and/or documented prior to surgery. The data could be further analyzed based on physician or type of procedure. Patients who are not included in the numerator could be tracked to see if there were any adverse outcomes due to the lack of preoperative anemia screening.

**Sampling:** Yes. For additional information see the Population and Sampling Specifications Section.

Data Reported As: Aggregate rate generated from count data reported as a proportion.

Selected References: \* Roback JD, ed. Technical manual. 16th ed, Bethseda, MD: AABB, 2008.

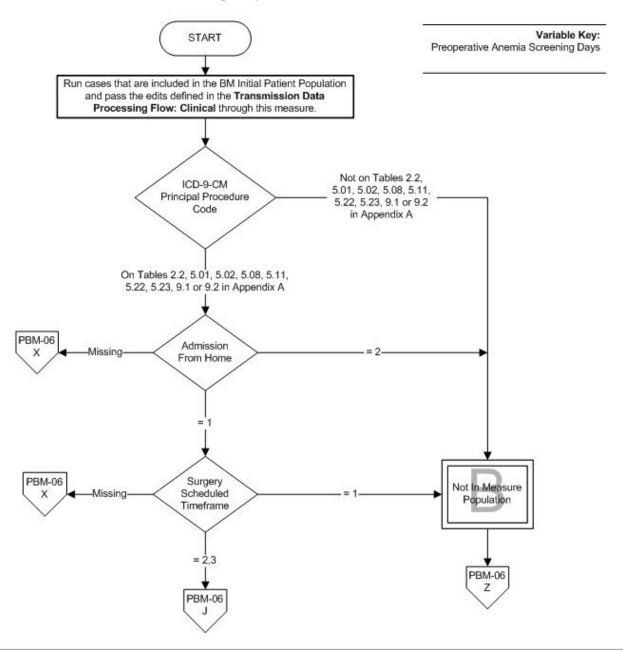
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- Rady MY, Ryan T, Starr NJ. Perioperative determinants of morbidity and mortality in elderly patients undergoing cardiac surgery. Crit Care Med. 1998;26: 225-235.
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- Campbell DA, Henderson WG, Englesbe, MJ, Hall BL, O'Reilly M, Bratzler D et al. Surgical site infection prevention: the importance of operative duration and blood transfusion-results of the first american college of surgeons –national surgical quality improvement program best practices initiative. J AM Coll Surg 2008;207:810-820.

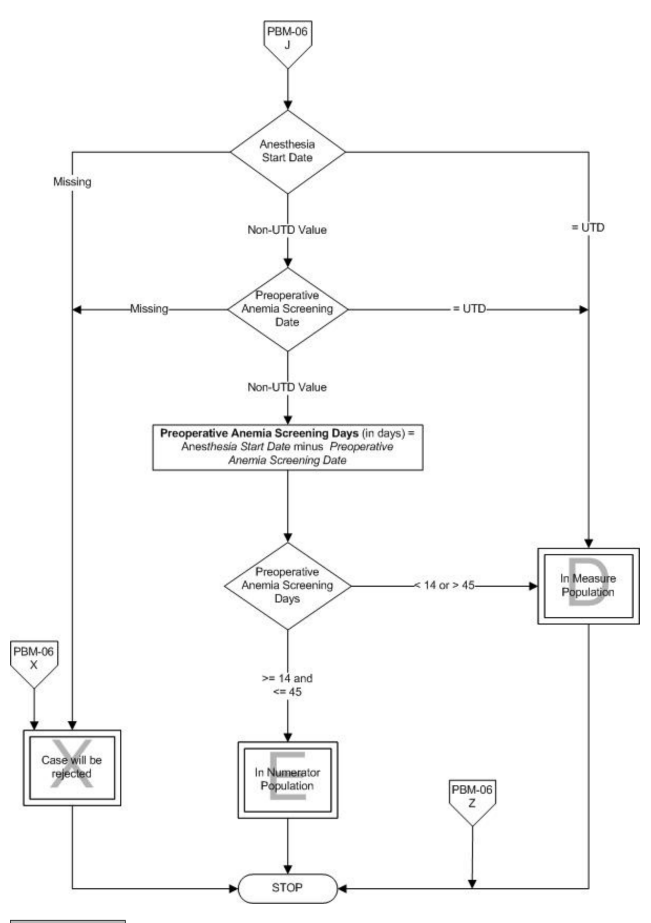
#### Measure Algorithm:

# PBM-06: Preoperative Anemia Screening

Numerator: Patients with documentation of preoperative anemia screening 14 - 45 days before Anesthesia Start Date

Denominator: Selected elective surgical patients





Related Topics

# **Measure Information Form**

Measure Set: Patient Blood Management(PBM)

#### Set Measure ID: PBM-07

Performance Measure Name: Preoperative Blood Type Testing and Antibody Screening

**Description:** Selected elective orthopedic, cardiac and hysterectomy surgical patients who had preoperative blood type testing and antibody screening (type and screen or type and crossmatch) completed prior to surgery start time if ordered preoperatively.

**Rationale:** Hospitals need to ensure that sufficient compatible blood is available for each scheduled procedure. Since about 3% of specimens have a serologic finding that requires further investigation that may cause a delay in the availability of the blood, patient screening of ABO group and Rh type should be collected in sufficient time to complete all pretransfusion testing before surgery begins. According to the Joint Commission's Pre-publication National Patient Safety Goal UP.01.01.01 for 2010, a preprocedure verification process should be conducted to identify items that must be available for the procedure and use a standardized list to verify their availability. Documentation of any required blood products for the procedure is required. Development of formal protocols to ensure that patients have blood testing completed prior to surgery start time for potential high-blood loss elective surgeries may optimize management of blood resources and maximize patient safety.

#### Type of Measure: Process

Improvement Noted As: Increase in the rate

**Numerator Statement:** Patients with preoperative type and crossmatch or type and screen completed prior to surgery start time

Included Populations: Not applicable

Excluded Populations: None

**Data Elements:** 

• Preoperative Blood Type Testing

Denominator Statement: Selected elective surgical patients

#### **Included Populations:**

• Discharges with an ICD-9-CM Principal Procedure Code of selected surgeries as defined in Appendix A, Tables 2.2, 5.01, 5.02, 5.08, 5.11, 5.22, 5.23, 9.1 or 9.2.

#### **Excluded Populations:**

- Patients less than 18 years of age
- Patients with type and screen or type and crossmatch ordered preoperatively

#### **Data Elements:**

- Admission Date
- Birthdate
- Blood Type Testing Ordered
- Discharge Date
- ICD-9-CM Principal Procedure Code

## Risk Adjustment: No.

**Data Collection Approach:** Retrospective data collection sources for required data elements include administrative data and medical records.

**Data Accuracy:** Variation may exist in the assignment of ICD-9-CM codes; therefore, coding practices may require evaluation to ensure consistency.

**Measure Analysis Suggestions:** These data may be used to evaluate specific patient groups at high risk for a blood transfusion that did not have pre-operative transfusion testing completed and/or documented prior to surgery start time. The data could be further analyzed based on physician or type of procedure. Patients who are not included in the numerator could be tracked to see if there were any adverse outcomes due to the lack of preoperative testing.

Sampling: Yes. For additional information see the Population and Sampling Specifications.

Data Reported As: Aggregate rate generated from count data reported as a proportion.

**Selected References:** \* Saxena S, Nelson JM, Osby M, Shah M, Kempf R, Shulman IA. Ensuring timely completion of type and screen testing and the verification of ABO/Rh status for elective surgical patients. Arch Pathol Lab Med. 2007;131:576-81.

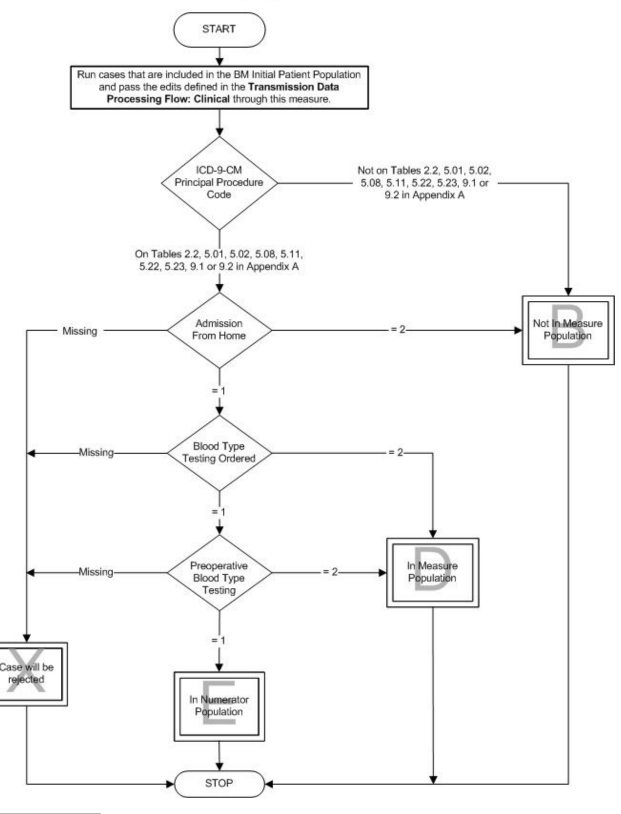
- Friedberg RC, Jones BA, Walsh MK. Type and screen completion for scheduled surgical procedures. A College of American Pathologists Q-Probes study of 8941 type and screen tests in 108 institutions. Arch Pathol Lab Med. 2003;127:533-40.
- Roback JD, ed. Technical manual. 16th ed, Bethseda, MD: AABB, 2008.
- Magovern JA, Sakert T, Magovern GJ et al. A model that predicts morbidity and mortality after coronary artery bypass graft surgery. J Am Coll Cardiol. 1996;28: 1147-1153.
- The Joint Commission 2010 National Patient Safety Goals, Oakbrook Terrace, IL [Available at <a href="http://www.jointcommission.org/NR/rdonlyres/868C9E07-037F-433D-8858-0D5FAA4322F2/0/RevisedChapter\_HAP\_NPSG\_20090924.pdf">http://www.jointcommission.org/NR/rdonlyres/868C9E07-037F-433D-8858-0D5FAA4322F2/0/RevisedChapter\_HAP\_NPSG\_20090924.pdf</a> (accessed January 27, 2010).]

## Measure Algorithm:

# PBM-07: Preoperative Blood Type Testing and Antibody Screening

Numerator: Patients with documentation of preoperative type and crossmatch or type and screen completed prior to Anesthesia Start Time

Denominator: Selected elective surgical patients



**Related Topics** 

Data Element Name:	Admission From Home
Collected For:	<u>PBM-06,</u>
Definition:	Patient was admitted for the pre-scheduled elective surgery procedure from home.
Suggested Data Collection Question:	Was the patient admitted from home?
Format:	Length: 1 Type: Alphanumeric Occurs: 1
Allowable Values:	<ol> <li>Patient was admitted from home.</li> <li>Patient was not admitted from home or unable to determine from medical record documentation.</li> </ol>
Notes for Abstraction:	<ul> <li>Patients who have to stay overnight at a location other than their primary residence due to long distance travel for procedure are considered admitted from home.</li> </ul>
Suggested Data Sources:	<ul> <li>Face sheet</li> <li>Nursing admission assessment</li> <li>Physician's notes</li> <li>Preop checklist</li> </ul>
Additional Notes:	
	Guidelines for Abstraction:

Inclusion	Exclusion
None	None

Data Element Name:	Anesthesia Start Date	
Collected For:	<u>PBM-06,</u>	
Definition:	The date the anesthesia for the procedure started.	
Suggested Data Collection Question:	On what date did the anesthesia for the procedure start?	
Format:	<ul> <li>Length: 10 – MM-DD-YYYY (includes dashes)</li> <li>Type: Date</li> <li>Occurs: 1</li> </ul>	
Allowable Values:	MM-DD-YYYY	
	MM = Month (01-12) DD = Day (01-31) YYYY = Year (2001-Current Year) Leave Blank if Unable to Determine	
Notes for Abstraction:	If the Anesthesia Start Date cannot be determined from medical record documentation, enter UTD. When the date documented is obviously invalid (not a valid format/range [12-39-20xx] or after the Discharge Date or Anesthesia End Date) and no other documentation can be found that provides the correct information, the abstractor should select "UTD."	
	Example: Patient expires on 02-12-20xx and documentation indicates the Anesthesia Start Date was 03-12-20xx. Other documentation in the medical record supports the date of death as being accurate, but no other documentation of the Anesthesia Start Date can be found. Since the Anesthesia Start Date is outside of the parameter for care (after the Discharge Date [death]) and no other documentation is found, the abstractor should leave blank.	
	If the Anesthesia Start Date is incorrect (in error) but it is a valid date and the correct date can be supported with other documentation in the medical record, the correct date may be entered. If supporting documentation of the correct date cannot be found, the medical record must be abstracted as documented or at "face value."	
	Examples: The anesthesia form is dated 12-10-2007, but other documentation in the medical record supports that the correct date was 12-10-2009. Enter the correct date of 12-10-2009 as the Anesthesia Start Date.	
	An Anesthesia End Date of 11-20-20xx is documented but the Anesthesia Start Date is documented as 11-10-20xx. If no other documentation can be found to support another Anesthesia Start Date, then it must be abstracted as 11-10-20xx because the date is not considered invalid or outside the parameter of care.	

## Suggested Data

Sources:

Other Suggested Sources:

- Intraoperative record
- Circulator record
- Post-anesthesia evaluation record
- · Operating room notes

Additional Notes: Suggested Data Sources:

Note: The anesthesia record is the priority data source for this data element, if a valid Anesthesia Start Date is found on the anesthesia record, use that date. If a valid date is not on the anesthesia record, other suggested data sources may be used in no particular order to determine the Anesthesia Start Date.

Priority Source:

· Anesthesia record

Inclusion	Exclusion
None	None

Data Element Name:	Blood Administration Location
<b>Collected For:</b>	<u>PBM-02, PBM-03, PBM-04, PBM-05,</u>
Definition:	The hospital setting (intraoperative or non-intraoperative) where the blood product began infusing.
Suggested Data Collection Question:	In what setting did the blood product begin infusing?
Format:	Length: 1 Type: Alphanumeric Occurs: 1-12
Allowable Values:	1 Intraoperative setting
	2 Non-introperative setting
	3 Unable to determine
Notes for Abstraction:	<ul> <li>Select setting for each unit transfused based on the physical location of the patient.</li> <li>Intraoperative setting is anytime during the operation.</li> </ul>
	<ul> <li>Non-intraoperative setting is any area outside of the operating room. For example, setting such as the intensive care unit, surgical floor or emergency room.</li> </ul>
Sources:	<ul> <li>Anesthesia record</li> <li>Emergency department record</li> <li>Nursing notes</li> <li>Nursing flow sheet</li> <li>Nursing admission assessment</li> <li>Progress notes</li> <li>Physician's notes</li> <li>Operative notes</li> <li>Operative report</li> <li>Procedure notes</li> <li>ICU notes</li> <li>PACU/recovery room record</li> </ul>
	Blood Administration Documentation Sheet
Additional Notae	

Inclusion	Exclusion
None	None

Data Element Name:	Blood Bank Records
Collected For:	<u>PBM-01, PBM-02, PBM-03, PBM-04, PBM-05,</u>
Definition:	Documentation that the patient received red blood cells (RBCs), plasma or platelets after hospital arrival.
Suggested Data Collection Question:	Was there documentation that the patient received RBCs, plasma or platelets after hospital arrival?
Format:	Length: 1 Type: Alphanumeric Occurs: 1-12
Allowable Values:	<ul> <li>Select all that apply: 1 RBCs</li> <li>2 Plasma</li> <li>3 Platelets</li> <li>4 None of the above or unable to determine from medical record documentation</li> </ul>
Notes for Abstraction:	<ul> <li>Include transfusions given in the emergency room or observation area.</li> </ul>
Suggested Data Sources:	Blood Bank Records
Additional Notes:	

Inclusion	Exclusion

Data Element Name:	Blood ID Number
Collected For:	<u>PBM-05,</u>
Definition:	Documentation of the actual blood bank identification number in the intraoperative record for the unit that was transfused.
Suggested Data Collection Question:	Was there documentation of a blood bank identification number for the unit or dose of blood transfused during surgery?
Format:	Length: 1 Type: Alphanumeric Occurs: 1
Allowable Values:	1 There is documentation of a blood bank identification number for the unit that was transfused.
	2 There is no documentation of a blood bank identification number for the unit that was transfused or unable to determine from medical record documentation.
Notes for Abstraction:	
Suggested Data Sources:	<ul><li>Anesthesia record</li><li>Operative report</li></ul>
	Blood administration record
Additional Notes:	
	Guidelines for Abstraction:

Inclusion	Exclusion
None	None

Data Element Name:	Blood Type Testing Ordered	
Collected For:	<u>PBM-07</u> ,	
Definition:	A type and screen and/or type and crossmatch was ordered preoperatively for the elective surgery.	
Suggested Data Collection Question:	Was a type and screen and/or type and crossmatch ordered preoperatively?	
Format:	Length: 1 Type: Alphanumeric Occurs: 1	
Allowable Values:	<ol> <li>A type and screen and/or type and crossmatch was ordered preoperatively.</li> <li>A type and screen and/or type and crossmatch was not ordered preoperatively or unable to determine</li> </ol>	
Notes for Abstraction:		
Suggested Data Sources:	<ul><li>Physician orders</li><li>Preop checklist</li></ul>	
Additional Notes:		

Inclusion	Exclusion
None	None

Data Element Name:	Clinical Indication for Plasma	
Collected For:	<u>PBM-03,</u>	
Definition:	Documentation by the physician/advance practice nurse/physician assistant or (physician/APN/PA) of the clinical indication for the plasma transfusion unit.	
Suggested Data Collection Question:	Was there a clinical indication documented by the physician/APN/PA for the transfused plasma unit?	
Format:	Length: 1 Type: Numeric Occurs: 1 - 3	
Allowable Values:	1 There was a clinical indication documented by the physician/APN/PA for the transfused plasma unit.	
	2 There was no documentation of a clinical indication for the transfusion or unable to determine from the medical record.	
Notes for Abstraction:	<ul> <li>The clinical indication for the transfusion must be documented within 24 hours after the start of the transfusion.</li> <li>Select the first four plasma transfusion units closest to hospital arrival for abstraction.</li> </ul>	
Suggested Data Sources:	ONLY PHYSICIAN/APN/PA DOCUMENTATION OF THE CLINICAL INDICATION FOR ADMINISTERING BLOOD:	
	<ul> <li>Anesthesia record</li> <li>Consultation notes</li> <li>Emergency department record</li> <li>Physician orders</li> <li>Progress notes</li> </ul>	
Additional Notes		

Inclusion	Exclusion
None	None

Data Element Name:	Clinical Indication for Platelets	
<b>Collected For:</b>	<u>PBM-04,</u>	
Definition:	Documentation by the physician/advance practice nurse/physician assistant (physician/APN/PA) of the clinical indication for the transfused platelet unit.	
Suggested Data Collection Question:	Was there a clinical indication documented by the physician/APN/PA for the transfused platelet unit?	
Format:	Length: 1 Type: Numeric Occurs: 1 - 3	
Allowable Values:	1 There was a clinical indication documented by the physician/APN/PA for the transfused platelet unit.	
	2 There was no documentation of clinical indication for the platelet transfusion or unable to determine from the medical record	
Notes for Abstraction:	<ul> <li>The clinical indication for the transfusion must be documented within 24 hours after the start of the transfusion.</li> </ul>	
	<ul> <li>Select the first three units transfused after hospital arrival for abstraction.</li> </ul>	
Suggested Data Sources:	ONLY PHYSICIAN/APN/PA DOCUMENTATION OF THE CLINICAL INDICATION FOR ADMINISTERING PLASMA:	
	<ul> <li>Anesthesia record</li> <li>Consultation notes</li> <li>Emergency department record</li> <li>Physician orders</li> <li>Progress notes</li> </ul>	
Additional Notes:		

Inclusion	Exclusion
None	None

Data Element Name:	Clinical Indication for RBCs	
<b>Collected For:</b>	<u>PBM-02,</u>	
Definition:	Documentation by the physician/advance practice nurse/physician assistant (physician/APN/PA) of the clinical indication for the tranfused red blood cell (RBCs) unit.	
Suggested Data Collection Question:	Was there a clinical indication documented by the physician/APN/PA for the transfused RBC unit?	
Format:	Length: 1 Type: Numeric Occurs: 1 - 6	
Allowable Values:	1 There was a clinical indication documented by the physician/APN/PA for the transfused RBC unit.	
	2 There was no clinical indication documented by the physician/APN/PA for the transfused RBC unit or unable to determine from medical record documentation.	
Notes for Abstraction:	<ul> <li>The clinical indication for the transfusion must be documented within 24 hours after the start of the transfusion.</li> <li>Select the first six RBC transfusion units after hospital arrival for abstraction.</li> </ul>	
Suggested Data Sources:	ONLY PHYSICIAN/APN/PA DOCUMENTATION OF THE CLINICAL INDICATION FOR ADMINISTERING RBCs:	
	<ul> <li>Anesthesia record</li> <li>Consultation notes</li> <li>Emergency department record</li> <li>Operative notes</li> <li>Physician orders</li> <li>Progress notes</li> </ul>	
Additional Notes:		
	Guidelines for Abstraction:	

Inclusion	Exclusion
None	None

Data Element Name:	Education Addressed Risks, Benefits and Alternatives to Transfusion	
Collected For:	<u>PBM-01,</u>	
Definition:	Documentation that information addressing risks, benefits and alternatives to transfusion was given to the patient/caregiver prior to the initial transfusion or the initial transfusion was deemed a medical emergency after hospital arrival.	
Suggested Data Collection Question:	Was there documentation that information regarding risks, benefits and alternatives to transfusion was given to the patient/caregiver prior to the initial transfusion event or was the initial transfusion deemed a medical emergency after hospital arrival?	
Format:	Length: 1 Type: Numeric Occurs: 1	
Allowable Values:	1 Information addressing the risks, benefits and alternatives to transfusion was given to the patient/caregiver prior to the initial transfusion after hospital arrival.	
	2 Information addressing the risks, benefits and alternatives to transfusion was not given to the patient/caregiver prior to the initial transfusion after hospital arrival or unable to determine from medical record documentation.	
Notes for Abstraction:	<ul> <li>Use only documentation provided in the medical record.</li> <li>If the patient refused information about risks, benefits and alternatives to transfusion, select "1."</li> <li>The caregiver is defined as the patient's family or any other person (e.g., guardian) who will be responsible for care of the patient.</li> </ul>	
Suggested Data Sources:	<ul> <li>Consultation notes</li> <li>Emergency department record</li> <li>History and physical</li> <li>Nursing notes</li> <li>Progress notes</li> <li>Operative notes</li> <li>Admission forms</li> <li>Consent form</li> <li>Emergency department record</li> <li>Progress notes</li> <li>Nursing notes</li> </ul>	

Inclusion	Exclusion
None	None

Data Element Name:	Patient ID Verification
<b>Collected For:</b>	<u>PBM-05</u> ,
Definition:	Documentation that two unique patient identifiers were checked during a two-person verification process (or the use of automated identification technology may be used in place of one of the individuals) prior to the administration of the transfusion unit/dose (bag).
Suggested Data Collection Question:	Was there documentation that two unique patient identifiers were checked or automated identification was used in place of one person during the verification process prior to the administration of the blood transfusion unit/dose (bag)?
Format:	Length: 1 Type: Numeric Occurs: 1 - 12
Allowable Values:	1 There was documentation that two unique patient identifiers were checked during the two person verification process or an automated identification system was used in place of one of the individuals prior to the administration of the transfusion unit/dose (bag).
	2 There was no documentation that two unique patient identifiers or automated identification were used during the two-person identification check prior to the administration of the transfusion unit/dose (bag) or unable to determine from medical record documentation.
Notes for Abstraction:	<ul> <li>Patient ID Verification must be associated with the blood product and RBC ID that was selected for abstraction.</li> <li>Patient ID Verification can be documented by the signature of two persons that attest that two unique patient identifiers were checked to verify the identification of the patient prior to the transfusion or the signature of one person and an automated identification device.</li> <li>Patient identification number or unique identifier given at the time the crossmatch was drawn.</li> <li>The patient room number should not be used to identify the patient.</li> </ul>
•	<ul> <li>Anesthesia record</li> <li>Emergency department record</li> <li>Nursing notes</li> <li>Progress notes</li> <li>Physician's notes</li> <li>Operative notes</li> <li>Operative report</li> <li>Procedure notes</li> <li>PACU/recovery room record</li> </ul>

• Blood administration form

## **Additional Notes:**

Inclusion	Exclusion
None	None

Data Element Name:	Plasma ID
Collected For:	<u>PBM-03, PBM-05,</u>
Definition:	The number assigned to designate whether the plasma unit was the first, second or third unit transfused after hospital arrival.
Suggested Data Collection Question:	What number was assigned to the plasma unit selected for abstraction?
Format:	Length: 1 Type: Numeric Occurs: 1 - 3
Allowable Values:	1 First Plasma Unit
	2 Second Plasma Unit
	3 Third Plasma Unit
Notes for Abstraction:	<ul> <li>The abstractor assigns a plasma identification (ID) number for each unit evaluated.</li> <li>Each allowable value is only used one time and is determined by the order in which it was administered.</li> <li>Abstract up to three plasma transfusion units per patient.</li> <li>Include plasma transfusions administered after hospital arrival.</li> </ul>
Suggested Data Sources:	<ul> <li>Anesthesia record</li> <li>Emergency department record</li> <li>Progress notes</li> <li>Operative notes</li> <li>Blood administration form</li> <li>Blood bank records</li> </ul>
Additional Notes:	

Inclusion	Exclusion
None	None

Data Element Name:	Platelet ID
<b>Collected For:</b>	<u>PBM-04, PBM-05,</u>
Definition:	The number assigned to designate whether the platelet unit was the first, second or third unit that was transfused after hospital arrival.
Suggested Data Collection Question:	What number was assigned to the platelet unit selected for abstraction?
Format:	Length: 2 Type: Numeric Occurs: 1 - 3
Allowable Values:	1 First Platelet Unit
	2 Second Platelet Unit
	3 Third Platelet Unit
Notes for Abstraction:	<ul> <li>The abstractor assigns a platelet identification (ID) number for each unit evaluated.</li> <li>Each allowable value is only used one time and is determined by the order in which it was administered.</li> <li>Abstract up to three platelet units per patient</li> <li>Include platelet transfusions administered after hospital arrival.</li> </ul>
Sources:	<ul> <li>Anesthesia record</li> <li>Emergency department record</li> <li>Progress notes</li> <li>Operative notes</li> <li>Blood administration form</li> <li>Blood bank records</li> </ul>
Additional Notes:	
	Guidelines for Abstraction:

Inclusion	Exclusion
None	None

Data Element Name:	Pre-transfusion Hematocrit
Collected For:	<u>PBM-02,</u>
Definition:	Documentation of the closest hematocrit (hct) completed prior to the RBC transfusion.
Suggested Data Collection Question:	What was documented as the closest pre-transfusion hct prior to the RBC transfusion?
Format:	Length: 4 Type: Alphanumeric Occurs: 1 - 6
Allowable Values:	Enter the patient's closest hematocrit result (number only, reported in percent) performed prior to each RBC transfusion.
	UTD = Unable to Determine
	<ul> <li>For abstraction, select either the pre-transfusion hematocrit or the hemoglobin result; both are not required.</li> <li>Select the result associated with the RBC ID selected for abstraction.</li> <li>When recording the allowable value for hematocrit, input 23.00 if the patient's hematocrit is 23%.</li> </ul>
Notes for Abstraction:	
Suggested Data Sources:	<ul> <li>Consultation notes</li> <li>Emergency department record</li> <li>History and physical</li> <li>Laboratory report</li> <li>Progress notes</li> <li>Operative report</li> <li>Blood administration form</li> </ul>

Inclusion	Exclusion
None	None

Data Element Name:	Pre-transfusion Hemoglobin
Collected For:	<u>PBM-02</u> ,
Definition:	Documentation of the closest hemoglobin (hgb) completed prior to the RBC transfusion.
Suggested Data Collection Question:	What was documented as the closest pre-transfusion hgb prior to the RBC transfusion?
Format:	Length: 4 Type: Alphanumeric Occurs: 1 - 6
Allowable Values:	Enter the patient's closest hemoglobin result reported in g/dL performed prior to transfusion.
	<b>UTD</b> = Unable to Determine
	<ul> <li>For abstraction, select either the pre-transfusion hematocrit or the hemoglobin result; both are not required.</li> <li>Select the hemoglobin result that is associated with the RBC ID selected for abstraction.</li> <li>If the hemoglobin result is 9.9 g/dL, enter 9.9.</li> </ul>
Notes for Abstraction:	
Suggested Data Sources:	<ul> <li>Consultation notes</li> <li>Emergency department record</li> <li>History and physical</li> <li>Laboratory report</li> <li>Progress notes</li> <li>Operative report</li> <li>Blood administration form</li> </ul>

Inclusion	Exclusion
None	None

Data Element Name:	Pre-transfusion PT/INR Result
<b>Collected For:</b>	<u>PBM-03,</u>
Definition:	Documentation of PT/INR result completed prior to the plasma transfusion.
Suggested Data Collection Question:	What was the PT/INR result completed prior to the plasma transfusion.
Format:	Length: 1 - 5 Type: Alphanumeric Occurs: 1 - 3
Allowable Values:	Enter the closest PT/INR result to the plasma transfusion. UTD = Unable to determine
Notes for Abstraction:	<ul> <li>Enter the PT/INR result that is associated with the plasma ID selected for abstaction.</li> </ul>
	<ul> <li>An allowable value should be entered with one decimal. For example, a PT/INR of 1.5 should be entered as written. INR values over 10 should be entered as 10.00.</li> </ul>
Suggested Data Sources:	
Additional Notes:	
	Guidelines for Abstraction:

Inclusion	Exclusion
None	None

Data Element Name:	Pre-transfusion Platelet Count	
Collected For:	<u>PBM-04,</u>	
Definition:	Documentation of the closest platelet count completed prior to the platelet transfusion.	
Suggested Data Collection Question:	What was the closest platelet count documented prior to the platelet transfusion?	
Format:	Length: 1 - 5 Type: Alphanumeric Occurs: 1 - 3	
Allowable Values:	Enter the patient's closest platelet count result, in 10 <sup>9</sup> /µL performed prior to the platelet transfusion selected for abstraction.	
	UTD = Unable to Determine	
	Note:	
	<ul> <li>Select the platelet count result that is associated with the Platelet ID selected for abstraction.</li> <li>An allowable value for a platelet count result should be entered as '11.00' for a platelet count of 11,000.</li> </ul>	
Notes for Abstraction:		
Suggested Data Sources:	<ul> <li>Anesthesia record</li> <li>Consultation notes</li> <li>Emergency department record</li> <li>History and physical</li> <li>Laboratory report</li> <li>Progress notes</li> <li>Operative report</li> <li>Blood administration form</li> </ul>	
Additional Notes:		
	Guidelines for Abstraction:	
	nclusion Exclusion	

Inclusion	Exclusion
None	None

Data Element Name:	Preoperative Anemia Screening Date	
Collected For:	<u>PBM-06,</u>	
Definition:	The date that preoperative anemia screening or a hemoglobin (hgb)or hematocrit (hct) result was completed.	
Suggested Data Collection Question:	What date was preoperative anemia screening or a hgb or hct result completed?	
Format:	<ul><li>Length: 10 - MM-DD-YYYY (includes dashes)</li><li>Type: Date</li><li>Occurs: 1</li></ul>	
Allowable Values:	MM-DD-YYYY	
	MM = Month (01-12) DD = Day (01-31) YYYY = Year (2001-Current Year) UTD	
Notes for Abstraction:	<ul> <li>Select the <i>Preoperative Anemia Screening Date</i> associated with the elective surgical procedure selected for abstraction. <i>Preoperative Transfusion Testing</i>.</li> <li>The medical record must be abstracted as documented (taken at "face value"). When the date documented is obviously in error (not a valid date/format) and no other documentation is found that provides this information, the abstractor should select UTD.</li> <li>Example: Documentation indicates the Preoperative Anemia Screening Date was 03-42-2008. No other documentation in the medical record provides a valid date. Since the Preoperative Anemia Screening Date is outside of the range listed in the Allowable Values for "Day," it is not a valid date, and the abstractor should select UTD.</li> </ul>	
•	Nursing notes Progress notes Preop checklist Pre-arrival laboratory reports	
Additional Notes:		
	Guidelines for Abstraction:	

Inclusion	Exclusion
None	None

Data Element Name:	Preoperative Blood Type Testing
<b>Collected For:</b>	<u>PBM-07,</u>
Definition:	Documentation that a type and screen or type and crossmatch was completed prior to anesthesia start time.
Suggested Data Collection Question:	Was there documentation of a type and screen or type and crossmatch completed prior to anesthesia start time?
Format:	Length: 1 Type: Numeric Occurs: 1
Allowable Values:	1 There is documentation that a type and screen or type and crossmatch was completed prior to anesthesia start time.
	2 There is no documentation that a type and screen or type and crossmatch was completed prior to anesthesia start time or unable to determine from medical record documentation.
Notes for Abstraction:	<ul> <li>If type and screen and type and crossmatch were completed prior to the surgical procedure, select "1".</li> </ul>
	<ul> <li>Anesthesia Start Time is the same as surgery start time.</li> </ul>
Suggested Data Sources:	<ul> <li>Consultation notes</li> <li>History and physical</li> <li>Progress notes</li> <li>Preop checklist</li> <li>Pre-arrival laboratory reports</li> </ul>
Additional Notes:	

Inclusion	Exclusion
None	None

Data Element Name:	RBC ID
Collected For:	<u>PBM-02, PBM-05,</u>
Definition:	The number assigned to designate whether the RBC transfusion was the first through the sixth RBC transfusion unit that was transfused after hospital arrival.
Suggested Data Collection Question:	What RBC unit was selected for abstraction?
Format:	Length: 1 Type: Numeric Occurs: 1 - 6
Allowable Values:	1 First RBC Unit
	2 Second RBC Unit
	3 Third RBC Unit
	4 Fourth RBC Unit
	5 Fifth RBC Unit
	6 Sixth RBC Unit
Notes for Abstraction:	<ul> <li>The abstractor assigns a RBC identification (ID) number for each unit evaluated.</li> <li>Each allowable value is used only one time and is determined by the order in which it was administered.</li> <li>Abstract up to six RBC transfusion units per patient.</li> <li>Include RBC transfusions administered after hospital arrival.</li> </ul>
Suggested Data Sources:	<ul> <li>Anesthesia record</li> <li>Emergency department record</li> <li>Progress notes</li> <li>Operative notes</li> <li>Operative report</li> <li>Medication administration record (MAR)</li> <li>Blood administration form</li> <li>Blood bank records</li> </ul>

Inclusion Exclusion
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Data Element Name:	RBC Unit Exclusions	
Collected For:	<u>PBM-02, PBM-05,</u>	
Definition:	Red blood cell (RBC) units that are excluded from abstraction. The following RBC units excluded from abstraction are; units used for a massive transfusion protocol or documentation of hemorrhagic shock, uncrossmatched units given during an emergency situation and units used to prime equipment for treatment.	
Suggested Data Collection Question:	Was this unit transfused for a massive transfusion protocol, hemorrhagic shock, uncrossmatched or used to prime equipment?	
Format:	Length: 1 Type: Alphanumeric Occurs: 1-6	
Allowable Values:	<ol> <li>There was documentation that this unit was transfused for a massive transfusion protocol, hemorrhagic shock, uncrossmatched or used to prime equipment</li> </ol>	
	<ol> <li>There was no documentation that this unit was transfused for a massive transfusion protocol, hemorrhagic shock, uncrossmatched or used to prime equipment or unable to determine from medical record documentation.</li> </ol>	
Notes for Abstraction:	<ul> <li>If the initial six units transfused are excluded due to the exclusion criteria, abstract the next six units that were tranfused. If the patient only received RBC units that are excluded, then no RBC units should be abstracted.</li> </ul>	
Suggested Data Sources:	<ul> <li>Anesthesia record</li> <li>Circulation record</li> <li>Emergency department record</li> <li>Laboratory report</li> <li>Nursing notes</li> <li>Nursing flow sheet</li> <li>Progress notes</li> <li>Physician orders</li> <li>Physician's notes</li> <li>Operative notes</li> <li>Operative report</li> <li>Procedure notes</li> <li>ICU notes</li> </ul>	

Inclusion	Exclusion
None	None

Data Element Name:	Surgery Scheduled Timeframe	
Collected For:	<u>PBM-06</u> ,	
Definition:	The elective surgery was scheduled in less than 14 days from the planned surgery start date.	
Suggested Data Collection Question:	Was the elective surgery scheduled in less than 14 days from the planned surgery?	
Format:	Length: 1 Type: Alphanumeric Occurs: 1	
Allowable Values:	<ol> <li>There was documentation that the elective surgery was scheduled in less than 14 days from the planned surgery.</li> <li>There was no documentation that the elective surgery was scheduled in less than 14 days from the planned surgery or unable to determine from medical record documentation.</li> </ol>	
Notes for Abstraction:		
Suggested Data Sources:	Preop checklist	
	Preoperative paperwork	
Additional Notes:		

Inclusion	Exclusion
None	None

Data Element Name:	Transfusion Consent	
Collected For:	<u>PBM-01,</u>	
Definition:	Documentation of a signed consent <b>prior</b> to the first transfusion of RBCs, platelets or plasma.	
Suggested Data Collection Question:	Was there documentation of a signed consent <b>prior</b> to the first blood transfusion?	
Format:	Length: 1 Type: Numeric Occurs: 1	
Allowable Values:	<b>1</b> There was documentation of a signed consent prior to the first blood transfusion.	
	2 The first blood transfusion was deemed a medical emergency.	
	<b>3</b> There was no documentation of a blood transfusion consent prior to the first blood transfusion or unable to determine from medical record documentation.	
Notes for Abstraction:	<ul> <li>The consent may be signed by the patient or caregiver.</li> <li>If organizations require a consent prior to every transfusion, then review the record for the first transfusion to answer this data element.</li> <li>For hospitals that use a general consent for treatment that includes transfusions, select "Yes".</li> <li>If a patient receives chronic transfusions and a previous consent is acceptable for a defined timeframe within the institution, select "1" if the consent is valid.</li> </ul>	
Suggested Data Sources:	<ul> <li>Emergency department record</li> <li>History and physical</li> <li>Nursing notes</li> <li>Progress notes</li> <li>Operative notes</li> <li>Consent form</li> </ul>	
Additional Notes:		
	Guidelines for Abstraction:	

Inclusion	Exclusion
None	None

Data Element Name:	Transfusion Order	
Collected For:	<u>PBM-05,</u>	
Definition:	An order to transfuse was written by the physician/advance practice nurse/physician assistant (physician/APN/PA) <b>prior</b> to the initiation of the transfusion.	
Suggested Data Collection Question:	Was there documentation of an order to transfuse <b>prior</b> to the transfusion?	
Format:	Length: 1 Type: Numeric Occurs: 1 - 12	
Allowable Values:	<ol> <li>There was documentation of an order to transfuse prior to transfusion.</li> </ol>	
	2 There was no documentation of an order to transfuse prior to transfusion or unable to determine from medical record documentation.	
Notes for Abstraction:	<ul> <li>A verbal or telephone order that was written prior to the transfusion is acceptable.</li> <li>The Transfusion Order must be associated with the blood product unit ID that was selected for abstraction.</li> <li>Note: Transfusion Order may apply to more than one unit/dose (bag). For example: An order written to "Transfuse two doses of platelets" would apply to both bags that were administered.</li> </ul>	
Suggested Data Sources:	ONLY PHYSICIAN/APN/PA DOCUMENTATION OF THE ORDER TO TRANSFUSE:	
	<ul> <li>Anesthesia record</li> <li>Consultation notes</li> <li>Emergency department record</li> <li>Operative notes</li> <li>Physician orders</li> <li>Progress notes</li> </ul>	
Additional Notes:		

#### **Additional Notes:**

Inclusion	Exclusion
None	None

Data Element Name:	Transfusion Start Date	
Collected For:	<u>PBM-05,</u>	
Definition:	The date that the blood transfusion unit/dose (bag) was administered.	
Suggested Data Collection Question:	What is the date that the blood transfusion unit/dose (bag) was administered?	
Format:	<ul> <li>Length: 10 – MM-DD-YYYY (includes dashes)</li> <li>Type: Date</li> <li>Occurs: 1 - 12</li> </ul>	
Allowable Values:	MM-DD-YYYY	
	MM = Month (01-12) DD = Day (01-31) YYYY = Year (2001-Current Year) UTD	
Notes for Abstraction:	<ul> <li>Abstract the Transfusion Date associated with the Transfusion Start Time of the unit/dose (bag) from the blood product ID selected for abstraction.</li> <li>Some of the dates of the transfusion units may be the same date. Record a transfusion date for each unit abstracted up to three units for plasma or platelets or up to six units for RBCs.</li> <li>The medical record must be abstracted as documented (taken at "face value"). When the date documented is obviously in error (not a valid date/format) and no other documentation is found that provides this information, the abstractor should select UTD. Example: Documentation indicates the Transfusion Start Date was 03-42-2008. No other documentation in the medical record provides a valid date. Since the Transfusion Start Date is outside of the range listed in the Allowable Values for "Day," it is not a valid date and the abstractor should select UTD.</li> </ul>	
Suggested Data Sources:	<ul> <li>Anesthesia record</li> <li>Emergency department record</li> <li>Nursing notes</li> <li>Progress notes</li> <li>Operative notes</li> <li>Blood administration record</li> </ul>	
Additional Notes:		

Inclusion	Exclusion
None	None

Data Element Name:	Transfusion Start Time	
Collected For:	<u>PBM-05,</u>	
Definition:	The start time (military time) of the unit/dose (bag) of RBCs, plasma or platelets that was administered.	
Suggested Data Collection Question:	What was the start time of the blood unit/dose (bag) administration?	
Format:	<ul> <li>Length: 5 - HH:MM (with or without colon) or UTD</li> <li>Type: Time</li> <li>Occurs: 1 - 12</li> </ul>	
Allowable Values:	Select the Transfusion Start Time associated with the Transfusion Start Date of the unit/dose (bag) from the associated blood product ID being abstracted.	
	HH = Hour (00-23) MM = Minutes (00-59) UTD = Unable to Determine	
Notes for Abstraction:	Time must be recorded in military time format. With the exception of Midnight and Noon:	
	<ul> <li>If the time is in the a.m., conversion is not required</li> <li>If the time is in the p.m., add 12 to the clock time hour</li> </ul>	
	Examples: Midnight - 00:00 Noon - 12:00 5:31 am - 05:31 5:31pm - 17:31 11:59 am - 11:59 11:59pm - 23:59	
	<ul> <li>For times that include "seconds," remove the seconds and record the time as is. Example: 15:00:35 would be recorded as 15:00</li> <li>If more than one Transfusion Start Time is documented, use the earliest time documented.</li> <li>The medical record must be abstracted as documented (taken at "face value"). When the time documented is obviously in error (not a valid format/range) and no other documentation is found that provides this information, the abstractor should select "UTD."</li> <li>Example: Documentation indicates the Transfusion Start Time was 3300. Since the Transfusion Start Time is outside of the range in the Allowable Values for "Hour," it is not a valid time and the abstractor should select "UTD."</li> </ul>	
Suggested Data Sources:	Anesthesia record	

- Emergency department record
- Nursing notes
- Operative notes
- Operative report
- Blood administration form

#### Additional Notes:

Select the Transfusion Start Time associated with the Transfusion Start Date of the unit/dose (bag) from the blood product ID identified for abstraction.

Time must be recorded in military time format. With the exception of Midnight and Noon:

- If the time is in the a.m., conversion is not required
- If the time is in the p.m., add 12 to the clock time hour.

The medical record must be abstracted as documented (taken at "face value"). When the time documented is obviously in error (not a valid format/range) and no other documentation is found that provides this information, the abstractor should select "UTD."

#### Example:

Documentation indicates the Transfusion Start Time was 3300. Since the Transfusion Start Time is outside of the range in the Allowable Values for "Hour," it is not a valid time and the abstractor should select "UTD."

Inclusion	Exclusion
None	None

Data Element Name:	Vital Sign Monitoring
<b>Collected For:</b>	<u>PBM-05,</u>
Definition:	Documentation of blood pressure (BP), pulse and temperature monitored at specific intervals for the transfusion. The intervals are:
	<ul> <li>Pre-transfusion, within 15 minutes of the initiation of the transfusion and within one hour of transfusion completion</li> </ul>
Suggested Data Collection Question:	Was there documentation of BP and temperature monitored for all of the specified intervals for the transfusion?
Format:	Length: 2 Type: Numeric Occurs: 1 -12
Allowable Values:	1 There was documentation for all of the BP, pulse and temperature monitoring intervals for the transfusion.
	2 There was no documentation for all of the blood pressure, pulse and temperature monitoring intervals for the transfusion or unable to determine from medical record documentation.
Notes for Abstraction:	<ul> <li>All vital signs must be recorded at the following times: pre-transfusion, within 15 minutes of the initiation of the transfusion and within one hour of transfusion completion. To select "1", all recordings must be documented.</li> <li>The pre-transfusion BP, pulse and temperature must be within one hour of the Transfusion Start Time. Vitals documented at the start of the transfusion are considered "within one hour of transfusion initiation".</li> <li>For blood that may be transfused within 15 minutes, select "1" if the pre-transfusion and the within one hour of transfusion completion vitals are documented.</li> <li>Vitals documented at the completion of the transfusion are considered "within one hour of transfusion are considered the transfusion are selected for abstraction.</li> </ul>
	<ul> <li>Anesthesia record</li> <li>Consultation notes</li> <li>Emergency department record</li> <li>Nursing notes</li> <li>Progress notes</li> <li>Operative notes</li> </ul>

### **Additional Notes:**

Inclusion	Exclusion
None	None

Index		
Number	Name	Page
Table 2.2	Left Ventricular Assistive Device (LVAD) and Heart	
	Transplant	
Table 5.01	Coronary Artery Bypass Graft (CABG)	
Table 5.02	Other Cardiac Surgery	
Table 5.08	Vascular Surgery	
Table 5.11	Cardiac Surgery	
Table 5.22	Elective Hip Replacement	
Table 5.23	Elective Total Knee Replacement	
Table 9.1	Elective Cardiac Surgery	
Table 9.2	Elective Hysterectomy	
Table 9.3	Previously Donated Autologous Transfusion	
Table 9.4	Packed Red Blood Cell Transfusion	
Table 9.5	Platelet Transfusion	
Table 9.6	Plasma (Serum) Transfusion	
Table 9.7	Trauma	

Table 2	Table 2.2         Left Ventricular Assistive Device (LVAD) and Heart Transplant		
Code	ICD-9-CM Description	Shortened Description	
33.6	Combined heart-lung transplantation	COMB HEART/LUNG	
		TRANSPLA	
37.51	Heart transplantation	HEART TRANSPLANTATION	
37.52	Implantation of total replacement heart system	IMPLANT TOT REP HRT SYS	
37.53	Replacement or repair of thoracic unit of total	REPL/REP THORAC UNIT HRT	
	replacement heart system		
37.54	Replacement or repair of other implantable	REPL/REP OTH TOT HRT SYS	
	component of total replacement heart system		
37.62	Insertion of non-implantable heart assist system	INS NON-IMPL HRT ASSIST	
37.63	Repair of heart assist system	REPAIR HEART ASSIST SYS	
37.64	Removal of heart assist system	REMOVE HEART ASSIST SYS	
37.65	Implant of external heart assist system	IMP EXT HRT ASSIST SYST	
37.66	Insertion of implantable heart assist system	IMPLANTABLE HRT ASSIST	
37.68	Insertion of percutaneous external heart assist	PERCUTAN HRT ASSIST SYST	
	device		

Table 5	Table 5.01         Coronary Artery Bypass Graft (CABG)		
Code	ICD-9-CM Description	Shortened Description	
36.10	Aortocoronary bypass for heart revascularization,	AORTOCORONARY BYPASS	
	not otherwise specified	NOS	
36.11	(Aorto)coronary bypass of one coronary artery	(AORTO)COR BYPAS-1 COR ART	
36.12	(Aorto)coronary bypass of two coronary arteries	(AORTO)COR BYPAS-2 COR ART	
36.13	(Aorto)coronary bypass of three coronary arteries	(AORTO)COR BYPAS-3 COR	
		ART	
36.14	(Aorto)coronary bypass of four coronary arteries	(AORT)COR BYPAS-4+ COR	
		ART	
36.15	Single internal mammary-coronary artery bypass	1 INT MAM-COR ART BYPASS	
36.16	Double internal mammary-coronary artery bypass	2 INT MAM-COR ART BYPASS	
36.17	Abdominal-coronary artery bypass	ABD-CORON ARTERY	
		BYPASS	
36.19	Other bypass anastomosis for heart	HRT REVAS BYPS ANAS NEC	
	revascularization		

Table 5	Table 5.02   Other Cardiac Surgery		
Code	ICD-9-CM Description	Shortened Description	
35.10	Open heart valvuloplasty, without replacement, unspecified valve	OPEN VALVULOPLASTY NOS	
35.11	Open heart valvuloplasty of aortic valve without replacement	OPN AORTIC VALVULOPLASTY	
35.12	Open heart valvuloplasty of mitral valve without replacement	OPN MITRAL VALVULOPLASTY	
35.13	Open heart valvuloplasty of pulmonary valve without replacement	OPN PULMON VALVULOPLASTY	
35.14	Open heart valvuloplasty of tricuspid valve without	OPN TRICUS	

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	replacement	VALVULOPLASTY
35.20	Replacement of unspecified heart valve	REPLACE HEART VALVE NOS
35.20	Replacement of aortic valve with tissue graft	REPLACE AORT VALVE NOS
35.22	Other replacement of aortic valve	
05.00		
35.23	Replacement of mitral valve with tissue graft	REPLACE MITR VALV-TISSUE
35.24	Other replacement of mitral valve	REPLACE MITRAL VALVE NEC
35.25	Replacement of pulmonary valve with tissue graft	REPLACE PULM VALV-TISSUE
35.26	Other replacement of pulmonary valve	REPLACE PULMON VALVE
35.27	Replacement of tricuspid valve with tissue graft	REPLACE TRIC VALV-TISSUE
35.28	Other replacement of tricuspid valve	REPLACE TRICUSP VALV NEC
35.31	Operations on papillary muscle	PAPILLARY MUSCLE OPS
35.32	Operations on chordae tendineae	CHORDAE TENDINEAE OPS
35.33	Annuloplasty	ANNULOPLASTY
35.34	Infundibulectomy	INFUNDIBULECTOMY
35.35	Operations on trabeculae carneae cordis	TRABECUL CARNEAE CORD
		OP
35.39	Operations on other structures adjacent to valves of heart	TISS ADJ TO VALV OPS NEC
35.42	Creation of septal defect in heart	CREATE SEPTAL DEFECT
35.50	Repair of unspecified septal defect of heart with	PROSTH REP HRT SEPTA
	prosthesis	NOS
35.51	Repair of atrial septal defect with prosthesis, open	PROS REP ATRIAL DEF-OPN
	technique	
35.53	Repair of ventricular septal defect with prosthesis, open technique	PROS REP VENTRIC DEF- OPN
35.54	Repair of endocardial defect with prosthesis	PROS REP ENDOCAR CUSHION
35.60	Repair of unspecified septal defect with tissue graft	GRFT REPAIR HRT SEPT NOS
35.61	Repair of atrial septal defect with tissue graft	GRAFT REPAIR ATRIAL DEF
35.62	Repair of ventricular septal defect with tissue graft	GRAFT REPAIR VENTRIC DEF
35.63	Repair of endocardial cushion defect with tissue	GRFT REP ENDOCAR
	graft	CUSHION
35.70	Other and unspecified repair of unspecified septal defect of heart	HEART SEPTA REPAIR NOS
35.72	Other and unspecified repair of ventricular septal defect	VENTR SEPTA DEF REP NEC
35.73	Other and unspecified repair of endocardial cushion defect	ENDOCAR CUSHION REP
35.81	Total repair of tetralogy of Fallot	TOT REPAIR TETRAL FALLOT
35.82	Total repair of total anomalous pulmonary venous connection	TOTAL REPAIR OF TAPVC
35.83	Total repair of truncus arteriosus	TOT REP TRUNCUS ARTERIOS
35.84	Total correction of transposition of great vessels, not elsewhere classified	TOT COR TRANSPOS GRT VES
35.91	Interatrial transposition of venous return	INTERAT VEN RETRN TRANSP

35.92	Creation of conduit between right ventricle and pulmonary artery	CONDUIT RT VENT-PUL ART
35.93	Creation of conduit between left ventricle and aorta	CONDUIT LEFT VENTR-
		AORTA
35.94	Creation of conduit between atrium and pulmonary	CONDUIT ARTIUM-PULM ART
	artery	
35.98	Other operations on septa of heart	OTHER HEART SEPTA OPS
35.99	Other operations on valves of heart	OTHER HEART VALVE OPS

Table 5	Table 5.08   Vascular Surgery		
Code	ICD-9-CM Description	Shortened Description	
38.14	Endarterectomy, aorta	ENDARTERECTOMY OF	
		AORTA	
38.16	Endarterectomy, abdominal arteries	ABDOMINAL	
		ENDARTERECTOMY	
38.18	Endarterectomy, lower limb arteries	LOWER LIMB ENDARTERECT	
38.34	Resection of vessel with anastomosis, aorta	<b>AORTA RESECTION &amp; ANAST</b>	
38.36	Resection of vessel with anastomosis, abdominal	ABD VESSEL RESECT/ANAST	
	arteries		
38.37	Resection of vessel with anastomosis, abdominal	ABD VEIN RESECT & ANAST	
	veins		
38.44	Resection of vessel with replacement, aorta,	RESECT ABDM	
	abdominal		
38.48	Resection of vessel with replacement, lower limb	LEG ARTERY RESEC W	
	arteries	REPLA	
38.49	Resection of vessel with replacement, lower limb	LEG VEIN RESECT W REPLAC	
	veins		
38.64	Other excision of vessels, aorta, abdominal	EXCISION OF AORTA	
39.25	Aorta-iliac-femoral bypass	AORTA-ILIAC-FEMOR BYPASS	
39.26	Other intra-abdominal vascular shunt or bypass	INTRA-ABDOMIN SHUNT NEC	
39.29	Other (peripheral) vascular shunt or bypass	VASC SHUNT & BYPASS NEC	

Table 5	Table 5.11 Cardiac Surgery		
Code	ICD-9-CM Description	Shortened Description	
35.10	Open heart valvuloplasty without replacement,	OPEN VALVULOPLASTY NOS	
	unspecified valve		
35.11	Open heart valvuloplasty of aortic valve without	OPN AORTIC	
	replacement	VALVULOPLASTY	
35.12	Open heart valvuloplasty of mitral valve without	OPNMITRAL VALVULOPLASTY	
	replacement		
35.13	Open heart valvuloplasty of pulmonary valve	OPN PULMON	
	without replacement	VALVULOPLASTY	
35.14	Open heart valvuloplasty of tricuspid valve without	OPN TRICUS	
	replacement	VALVULOPLASTY	
35.20	Replacement of unspecified heart valve	REPLACE HEART VALVE NOS	
35.21	Replacement of aortic valve with tissue graft	REPLACE AORT VALVE-	
		TISSUE	
35.22	Other replacement of aortic valve	REPLACE AORT VALVE NEC	

35.23	Replacement of mitral valve with tissue graft	REPLACE MITR VALVE-
00.20		TISSUE
35.24	Other replacement of mitral valve	REPLACE MITRAL VALVE NEC
35.25	Replacement of pulmonary valve with tissue graft	REPLACE PULM VALV-TISSUE
35.26	Other replacement of pulmonary valve	REPLACE PULMON VALVE
		NEC
35.27	Replacement of tricuspid valve with tissue graft	REPLACE TRICUSP VALV NEC
35.28	Other replacement of tricuspid valve	REPLACE TRICUSP VALV NEC
35.31	Operations on papillary muscle	PAPILLARY MUSCLE OPS
35.32	Operations on chordae tendineae	CHORDAE TENDINEAE OPS
35.33	Annuloplasty	ANNULOPLASTY
35.34	Infundibulectomy	INFUNDIBULECTOMY
35.35	Operations of trabeculae carneae cordis	TRABECUL CARNEAE CORD
35.39	Operations on other structures adjacent to valves of heart	TISS ADJ TO VALV OPS NEC
35.42	Creation of septal defect in heart	CREATE SEPTAL DEFECT
35.50	Repair of unspecified septal defect of heart with	PROSTH REP HRT SEPTA
	prosthesis	NOS
35.51	Repair of atrial septal defect with prosthesis, open technique	PROS REP ATRIAL DEF-OPN
35.53	Repair of ventricular septal defect with prosthesis, open technique	PROS REP VENTRIC DEF-
35.54	Repair of endocardial cushion defect with	PROS REP ENDOCAR
	prosthesis	CUSHION
35.60	Repair of unspecified septal defect of heart with tissue graft	GRFT REPAIR HRT SEPT NOS
35.61	Repair of atrial septal defect with tissue graft	GRAFT REPAIR ATRIAL DEF
35.62	Repair of ventricular septal defect with tissue graft	GRAFT REPAIR VENTRIC DEF
35.63	Repair of endocardial cushion defect with tissue graft	GRFT REP ENDOCAR CUSHION
35.70	Other and unspecified repair of unspecified septal defect of heart	HEART SEPTA REPAIR NOS
35.71	Other and unspecified repair of atrial septal defect	ATRIA SEPTA DEF REP NEC
35.72	Other and unspecified repair of ventricular septal defect	VENTR SEPTA DEF REP NEC
35.73	Other and unspecified repair of endocardial cushion defect	ENDOCAR CUSHION REP
35.81	Total repair of tetralogy of Fallot	TOT REPAIR TETRAL FALLOT
35.82	Total repair of total anomalous pulmonary venous connection	TOTAL REPAIR OF TAPVC
35.83	Total repair of truncus arteriosus	TOT REP TRUNCUS ARTERIOS

Table 5.	11 Cardiac Surgery (cont.)	
Code	ICD-9-CM Description	Shortened Description

35.84	Total connection of transposition of great vessels, not elsewhere classified	TOT COR TRANSPOS GRT VES
35.91	Interatrial transposition of venous return	INTERAT VEN RETRN TRANSP
35.92	Creation of conduit between right ventricle and pulmonary artery	CONDUIT RT VENT-PUL ART
35.93	Creation of conduit between left ventricle and aorta	CONDUIT LEFT VENTR- AORTA
35.94	Creation of conduit between atrium and pulmonary artery	CONDUIT ARTIUM-PULM ART
35.98	Other operations on septa of heart	OTHER HEART SEPTA OPS
35.99	Other operations on valves of heart	OTHER HEART VALVE OPS
36.03	Open chest coronary artery angioplasty	OPEN CORONRY ANGIOPLASTY
36.10	Aortocoronary bypass for heart revascularization, not otherwise specified	AORTOCORONARY BYPASS NOS
36.11	Aortocoronary bypass of one coronary artery	AORTOCOR BYPASS-1 COR ART
36.12	Aortocoronary bypass of two coronary arteries	AORTOCOR BYPASS-2 COR ART
36.13	Aortocoronary bypass of three coronary arteries	AORTOCOR BYPASS-3 COR ART
36.14	Aortocoronary bypass of four or more coronary arteries	AORTOCOR BYPASS-4+ COR ART
36.15	Single internal mammary-coronary artery bypass	1 INT MAM-COR ART BYPASS
36.16	Double internal mammary-coronary artery bypass	2 INT MAM-COR ART BYPASS
36.17	Abdominal-coronary artery bypass	ABD-CORON ARTERY BYPASS
36.19	Other bypass anastomosis for heart revascularization	HRT REVAS BYPS ANAS NEC
36.31	Open chest transmyocardial revascularization	OPEN CHEST TRANS REVASC
36.32	Other transmyocardial revascularization	OTH TRANSMYO REVASCULAR
36.39	Other heart revascularization	OTH REVASCULAR
36.91	Repair of aneurysm of coronary vessel	CORON VESS ANEURYSM REP
36.99	Other operations on vessels of heart	HEART VESSEL OP NEC
37.10	Incision of heart, not otherwise specified	INCISION OF HEART NOS
37.11	Cardiotomy	CARDIOTOMY
37.31	Pericardiectomy	PERICARDIECTOMY
37.32	Excision of aneurysm of heart	HEART ANEURYSM EXCISION
37.33	Excision or destruction of other lesion or tissue of heart, open approach	EXC/DEST HRT LESION OPEN
37.35	Partial ventriculectomy	PARTIAL VENTRICULECTOMY
37.41	Implantation of prosthetic cardiac support device around the heart	IMPL CARDIAC SUPPORT DEV
37.49	Other repair of heart and pericardium	HEART/PERICARD REPR NEC
37.51	Heart transplantation	HEART TRANSPLANTATION

37.52	Implantation of total replacement heart system	IMPLANT TOT REP HRT SYS
37.53	Replacement or repair of thoracic unit of total replacement heart system	REPL/REP THORAC UNIT HRT
37.54	Replacement or repair of other implants component of total replacement heart system	REPL/REP OTH TOT HRT SYS
37.62	Insertion of non-implantable heart assist system	INS NON-IMPL HRT ASSIST
37.63	Repair of heart assist system	REPAIR HEART ASSIST SYS
37.64	Removal of heart assist system	REMOVE HEART ASSIST SYS
37.66	Insertion of implantable heart assist system	IMPLANTABLE HRT ASSIST
37.67	Implantation of cardiomyostimulation system	IMP CARDIOMYOSTIMUL SYS

Table 5	Table 5.22 Elective Hip Replacement		
Code	ICD-9-CM Description	Shortened Description	
00.70	Revision of hip replacement, both acetabular and femoral components	REV HIP REPL-ACETAB/FEM	
00.71	Revision of hip replacement, acetabular component	REV HIP REPL-ACETAB COMP	
00.72	Revision of hip replacement, femoral component	REV HIP REPL-FEM COMP	
00.73	Revision of hip replacement, acetabular liner and/or femoral head only	REV HIP REPL-LINER/HEAD	
00.77	Hip bearing surface, ceramic-on-polyethylene	HIP SURFACE, CERMC/POLY	
00.85	Resurfacing hip, total, acetabulum and femoral head	RESRF HIP, TOTAL-ACET/FEM	
00.86	Resurfacing hip, partial, femoral head	RESRF HIP, PART-FEM HEAD	
00.87	Resurfacing hip, partial, acetabulum	RESRF HIP, PART-ACETABLUM	
81.51	Total hip replacement	TOTAL HIP REPLACEMENT	
81.52	Partial hip replacement	PARTIAL HIP REPLACEMENT	
81.53	Revision of hip replacement	REVISE HIP REPLACEMENT	

Table 5	Table 5.23 Elective Total Knee Replacement		
Code	ICD-9-CM Description	Shortened Description	
00.80	Revision of knee replacement, total (all components)	REV KNEE REPLACEMT-TOTAL	
00.81	Revision of knee replacement, tibial component	REV KNEE REPL-TIBIA COMP	
00.82	Revision of knee replacement, femoral component	REV KNEE REPL-FEMUR COMP	
00.83	Revision of knee replacement, patellar component	REV KNEE REPLACE-PATELLA	
00.84	Revision of total knee replacement, tibial insert (liner)	REV KNEE REPL-TIBIA LIN	
81.54	Total knee replacement	TOTAL KNEE REPLACEMENT	
81.55	Revision of knee replacement	REVISE KNEE REPLACEMENT	

Table 9.1         Elective Cardiac Surgery (Selected Codes from Table 5.25)		
Code	ICD-9-CM Description	Shortened Description
35.71	Other and unspecified repair of atrial septal defect	ATRIA SEPTA DEF REP NEC
36.03	Open chest coronary artery angioplasty	OPEN CORONRY ANGIOPLASTY
36.31	Open chest transmyocardial revascularization	OPEN CHEST TRANS REVASC
36.32	Other transmyocardial revascularization	OTH TRANSMYO REVASCULAR
36.39	Other heart revascularization	OTH HEART REVASCULAR
36.91	Repair of aneurysm of coronary vessel	CORON VESS ANEURYSM REP
36.99	Other operations on vessels of heart	HEART VESSEL OP NEC
37.10	Incision of heart, not otherwise specified	INCISION OF HEART NOS
37.11	Cardiotomy	CARDIOTOMY
37.32	Excision of aneurysm of heart	HEART ANEURYSM EXCISION
37.33	Excision or destruction of other lesion or tissue of heart, open approach	EXC/DEST HRT LESION OPEN
37.35	Partial ventriculectomy	PARTIAL VENTRICULECTOMY
37.36	Excision or destruction of left atrial appendage (LAA)	
37.41	Implantation of prosthetic cardiac support device around the heart	IMPL CARDIAC SUPPORT DEV
37.49	Other repair of heart and pericardium	HEART/PERICARD REPR NEC
37.51	Heart transplantation	HEART TRANSPLANTATION
37.52	Implantation of total internal biventricular heart replacement system	
37.53	Replacement or repair of thoracic unit of (total) replacement heart system	REPL/REP THR UNT TOT HRT
37.54	Replacement or repair of other implantable component of (total) replacement heart system	REPL/REP OTH TOT HRT SYS
37.55	Removal of internal biventricular heart replacement system	REM INT BIVENT HRT SYS
37.60	Implantation or insertion of biventricular external heart assist system	IMP BIVN EXT HRT AST SYS
37.62	Insertion of temporary non-implantable extracorporeal circulatory assist device	INSRT NON-IMPL CIRC DEV
37.63	Repair of heart assist system	REPAIR HEART ASSIST SYS
37.64	Removal of external heart assist system(s) or device(s)	REMVE EXT HRT ASSIST SYS
37.66	Insertion of implantable heart assist system	IMPLANTABLE HRT ASSIST
37.67	Implantation of cardiomyostimulation system	IMP CARDIOMYOSTIMUL SYS

Table 9.2 Elective Gynecological		
Code	ICD-9-CM Description	Shortened Description
68.31	Other incision and excision of uterus, subtotal abdominal hysterectomy, other incision and excision of uterus, laparoscopic supracervical hysterectomy [LSH]	Lap scervic hysterectomy
68.39	Other incision and excision of uterus, subtotal abdominal hysterectomy, other incision and excision of uterus, other and unspecified subtotal	Subtotl abd hyst NEC/NOS

	abdominal hysterectomy	
68.41	Other incision and excision of uterus, total abdominal hysterectomy, laparoscopic total	Lap total abdominal hyst
	abdominal hysterectomy	
68.49	Other incision and excision of uterus, total	Total abd hyst NEC/NOS
	abdominal hysterectomy, other and unspecified total abdominal hysterectomy	
68.51	Vaginal hysterectomy, laparoscopically assisted vaginal hysterectomy [LAVH]	Lap ast vag hysterectomy
68.59	Vaginal hysterectomy, other and unspecified	Vag hysterectomy NEC/NOS
	vaginal hysterectomy	
68.61	Radical abdominal hysterectomy, laparoscopic radical abdominal hysterectomy	Lap radical abdomnl hyst
68.69	Radical abdominal hysterectomy, other and unspecified radical abdominal hysterectomy	Radical abd hyst NEC/NOS
68.71	Radical vaginal hysterectomy, laparoscopic radical vaginal hysterectomy [LRVH]	Lap radical vaginal hyst
68.79	Radical vaginal hysterectomy, other and unspecified radical vaginal hysterectomy	Radical vag hyst NEC/NOS
68.9	Other and unspecified hysterectomy	Hysterectomy NEC/NOS

Table 9	Table 9.3         Previously Donated Autologous Transfusion		
Code	ICD-9-CM Description	Shortened Description	
99.02	Other nonoperative procedures, transfusion of	TRANSFUS PREV AUTO	
	blood and blood components, transfusion of	BLOOD	
	previously collected autologous blood		

Table 9	4 Packed Red Blood Cell Transfusion	
Code	ICD-9-CM Description	Shortened Description
99.04	Other nonoperative procedures, transfusion of blood and blood components, transfusion of packed cells	PACKED CELL TRANSFUSION

Table 9.5 Platelet Transfusion		
Code	ICD-9-CM Description	Shortened Description
99.05	Other nonoperative procedures, transfusion of blood and blood components, transfusion of platelets	PLATELET TRANSFUSION

Table 9	6 Plasma Transfusion	
Code	ICD-9-CM Description	Shortened Description
99.07	Other nonoperative procedures, transfusion of blood and blood components, transfusion of other serum	SERUM TRANSFUSION NEC

Table 9	.7 Trauma	
Code	ICD-9-CM Description	Shortened Description
800	Fracture of vault of skull	CLOSED SKULL VAULT FX
801	Fracture of base of skull	CLOS SKULL BASE
		FRACTURE
802	Fracture of face bones	NASAL BONE FX-CLOSED
803	Other and unqualified skull fractures	CLOSE SKULL FRACTURE
		NEC
804	Multiple fractures involving skull or face with other bones	CL SKUL FX W OTH BONE FX
805	Fracture of vertebral column without mention of spinal cord injury	FX CERVICAL VERT NOS-CL
806	Fracture of vertebral column with spinal cord injury	C1-C4 FX-CL/CORD INJ NOS
807	Fracture of rib(s), sternum, larynx, and trachea	FRACTURE RIB NOS-CLOSED
808	Fracture of pelvis	FRACTURE ACETABULUM-
		CLOS
809	III-defined fractures of bones of trunk	FRACTURE TRUNK BONE- CLOS
810	Fracture of clavicle	FX CLAVICLE NOS-CLOSED
811	Fracture of scapula	FX SCAPULA NOS-CLOSED
812	Fracture of humerus	FX UP END HUMERUS NOS-
		CL
813	Fracture of radius and ulna	FX UPPER FOREARM NOS-CL
814	Fracture of carpal bones(s)	FX CARPAL BONE NOS-
		CLOSE
815	Fracture of metacarpal bones(s)	FX METACARPAL NOS- CLOSED
816	Fracture of one or more phalanges of hands	FX PHALANX, HAND NOS-CL
817	Multiple fractures of hand bones	MULTIPLE FX HAND-CLOSED
818	III-defined fractures of upper limb	FX ARM MULT/NOS-CLOSED
819	Multiple fractures involving both upper limbs, and	FX ARMS W RIB/STERNUM-CL
	upper limb with rib(s) and sternum	
820	Fracture of neck of femur	FX FEMUR INTRCAPS NOS-CL
821	Fracture of other and unspecified parts of femur	FX FEMUR NOS-CLOSED
822	Fracture of patella	FRACTURE PATELLA-CLOSED
823	Fracture of tibia and fibula	FX UPPER END TIBIA-CLOSE
824	Fracture of ankle	FX MEDIAL MALLEOLUS-
		CLOS
825	Fracture of one or more tarsal and metatarsal	FRACTURE CALCANEUS-
-	bones	CLOSE
826	Fracture of one or more phalanges of foot	FX PHALANX, FOOT-CLOSED
827	Other, multiple, and ill-defined fractures of lower	FX LOWER LIMB NEC-
	limb	CLOSED
828	Multiple fractures involving both lower limbs, lower	FX LEGS W ARM/RIB-CLOSED
-	with upper limb, and lower limb(s) with rib(s) and	
	sternum	
829	Fracture of unspecified bones	FRACTURE NOS-CLOSED
830	Dislocation of jaw	DISLOCATION JAW-CLOSED
831	Dislocation of shoulder	DISLOC SHOULDER NOS-

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		CLOS
832	Dislocation of elbow	DISLOCAT ELBOW NOS-
002		CLOSE
833	Dislocation of wrist	DISLOC WRIST NOS-CLOSED
834	Dislocation of finger	DISL FINGER NOS-CLOSED
835	Dislocation of hip	DISLOCAT HIP NOS-CLOSED
836	Dislocation of knee	TEAR MED MENISC KNEE-
		CUR
837	Dislocation of ankle	DISLOCATION ANKLE- CLOSED
838	Dislocation of foot	DISLOCAT FOOT NOS- CLOSED
839	Other, multiple, and ill-defined dislocations	DISLOC CERV VERT NOS-CL
840	Sprains and strains of shoulder and upper arm	SPRAIN
010		ACROMIOCLAVICULAR
841	Sprains and strains of elbow and forearm	SPRAIN RADIAL COLLAT LIG
842	Sprains and strains of wrist and hand	SPRAIN OF WRIST NOS
843	Sprains and strains of hip and thigh	SPRAIN ILIOFEMORAL
844	Sprains and strains of knee and leg	SPRAIN LATERAL COLL LIG
845	Sprains and strains of ankle and foot	SPRAIN OF ANKLE NOS
846	Sprains and strains of sacroiliac region	SPRAIN LUMBOSACRAL
847	Sprains and strains of other and unspecified parts	SPRAIN OF NECK
011	of back	
848	Other and ill-defined sprains and strains	SPRAIN OF NASAL SEPTUM
850	Concussion	CONCUSSION W/O COMA
851	Cerebral laceration and contusion	CEREBRAL CORTX
		CONTUSION
852	Subarachnoid, subdural, and extradural	TRAUM SUBARACHNOID HEM
	hemorrhage, following injury	
853	Other and unspecified intracranial hemorrhage following injury	TRAUMATIC BRAIN HEM NEC
854	Intracranial injury of other and unspecified nature	BRAIN INJURY NEC
860	Traumatic pneumothorax and hemothorax	TRAUM PNEUMOTHORAX-
000		CLOSE
861	Injury to heart and lung	HEART INJURY NOS-CLOSED
862	Injury to other and unspecified intrathoracic organs	DIAPHRAGM INJURY-CLOSED
863	Injury to gastrointestinal tract	STOMACH INJURY-CLOSED
864	Injury to liver	LIVER INJURY NOS-CLOSED
865	Injury to spleen	SPLEEN INJURY NOS-
000		CLOSED
866	Injury to kidney	KIDNEY INJURY NOS-CLOSED
867	Injury to pelvic organs	BLADDER/URETHRA INJ-
007		CLOS
868	Injury to other intra-abdominal organs	INTRA-ABDOM INJ NOS-CLOS
869	Internal injury to unspecified or ill-defined organs	INTERNAL INJ NOS-CLOSED
870	Open wound of ocular adnexa	LAC EYELID SKN/PERIOCULR
870	Open wound of eyeball	OCULAR LAC W/O PROLAPSE
872	Open wound of ear	OPN WOUND EXTERN EAR
012		

		NOS
873	Other open wound of head	OPEN WOUND OF SCALP
874	Open wound of neck	OPN WND LARYNX W
0.1		TRACHEA
875	Open wound of chest (wall)	OPEN WOUND OF CHEST
876	Open wound of back	OPEN WOUND OF BACK
877	Open wound of buttock	OPEN WOUND OF BUTTOCK
878	Open wound of genital organs (external), including traumatic amputation	OPEN WOUND OF PENIS
879	Open wound of other and unspecified sites, except limbs	OPEN WOUND OF BREAST
880	Open wound of shoulder and upper arm	OPEN WOUND OF SHOULDER
881	Open would of elbow, forearm, and wrist	OPEN WOUND OF FOREARM
882	Open wound of hand except finger(s) alone	OPEN WOUND OF HAND
883	Open wound of finger(s)	OPEN WOUND OF FINGER
884	Multiple and unspecified open wound of upper limb	OPEN WOUND ARM
		MULT/NOS
885	Traumatic amputation of thumb (complete) (partial)	AMPUTATION THUMB
886	Traumatic amputation of other finger(s) (complete) (partial)	AMPUTATION FINGER
887	Traumatic amputation of arm and hand (complete) (partial)	AMPUT BELOW ELB, UNILAT
890	Öpen wound of hip and thigh	OPEN WOUND OF HIP/THIGH
891	Open wound of knee, leg [except thigh], and ankle	OPEN WND KNEE/LEG/ANKLE
892	Open wound of foot except toe(s) alone	OPEN WOUND OF FOOT
893	Open wound of toe(s)	OPEN WOUND OF TOE
894	Multiple and unspecified open wound of lower limb	OPEN WOUND OF LEG NEC
895	Traumatic amputation of toe(s) (complete) (partial)	AMPUTATION TOE
896	Traumatic amputation of foot (complete) (partial)	AMPUTATION FOOT, UNILAT
897	Traumatic amputation of leg(s) (complete) (partial)	AMPUT BELOW KNEE, UNILAT
900	Injury to blood vessels of head and neck	INJUR CAROTID ARTERY NOS
901	Injury to blood vessels of thorax	INJURY THORACIC AORTA
902	Injury to blood vessels of abdomen and pelvis	INJURY ABDOMINAL AORTA
903	Injury to blood vessels of upper extremity	INJ AXILLARY VESSEL NOS
904	Injury to blood vessels of lower extremity and unspecified sites	INJ COMMON FEMORAL ARTER
905	Late effects of musculoskeletal and connective tissue injuries	LATE EFFEC SKULL/FACE FX
906	Late effects of injuries to skin and subcutaneous tissues	LT EFF OPN WND HEAD/TRNK
907	Late effects of injuries to the nervous system	LT EFF INTRACRANIAL INJ
908	Late effects of other and unspecified injuries	LATE EFF INT INJUR CHEST
909	Late effects of other and unspecified external causes	LATE EFF DRUG POISONING
910	Superficial injury of face, neck, and scalp except eye	ABRASION HEAD
911	Superficial injury of trunk	ABRASION TRUNK
912	Superficial injury of shoulder and upper arm	ABRASION SHOULDER/ARM

913	Superficial injury of elbow, forearm, and wrist	ABRASION FOREARM
913	Superficial injury of hand(s) except finger(s) alone	ABRASION HAND
915	Superficial injury of finger(s)	ABRASION FINGER
916	Superficial injury of hip, thigh, leg, and ankle	ABRASION HIP & LEG
917	Superficial injury of foot and toe(s)	ABRASION FOOT & TOE
917	Superficial injury of eye and adnexa	SUPERFIC INJ PERIOCULAR
919	Superficial injury of other, multiple, and unspecified	ABRASION NEC
	sites	
920	Contusion of face, scalp, and neck except eye(s)	CONTUSION
		FACE/SCALP/NCK
921	Contusion of eye and adnexa	BLACK EYE NOS
922	Contusion of trunk	CONTUSION OF BREAST
923	Contusion of upper limb	CONTUSION SHOULDER REG
924	Contusion of lower limb and of other and	CONTUSION OF THIGH
	unspecified sites	
925	Crushing injury of face, scalp, and neck	
926	Crushing injury of trunk	CRUSH INJ EXT GENITALIA
927	Crushing injury of upper limb	CRUSH INJ SHOULDER REG
928	Crushing injury of lower limb	CRUSHING INJURY THIGH
929	Crushing injury of multiple and unspecified sites	CRUSH INJ MULT SITE NEC
930	Foreign body on external eye	CORNEAL FOREIGN BODY
931	Foreign body in ear	FOREIGN BODY IN EAR
932	Foreign body in nose	FOREIGN BODY IN NOSE
933	Foreign body in pharynx and larynx	FOREIGN BODY IN PHARYNX
934	Foreign body in trachea, bronchus, and lung	FOREIGN BODY IN TRACHEA
935	Foreign body in mouth, esophagus, and stomach	FOREIGN BODY IN MOUTH
936	Foreign body in intestine and colon	FB IN INTESTINE & COLON
937	Foreign body in anus and rectum	FOREIGN BODY
		ANUS/RECTUM
938	Foreign body in digestive system, unspecified	FOREIGN BODY GI NOS
939	Foreign body in genitourinary tract	FB BLADDER & URETHRA
940	Burn confined to eye and adnexa	CHEMICAL BURN
		PERIOCULAR
941	Burn of face, head, and neck	BURN NOS HEAD-UNSPEC
942	Burn of trunk	BURN NOS TRUNK-UNSPEC
943	Burn of upper limb, except wrist and hand	BURN NOS ARM-UNSPEC
944	Burn of wrist(s) and hand(s)	BURN NOS HAND-UNSPEC
945	Burn of lower limb(s)	BURN NOS LEG-UNSPEC
946	Burns of multiple specified sites	BURN NOS MULTIPLE SITE
947	Burn of internal organs	BURN OF MOUTH & PHARYNX
948	Burns classified according to extent of body surface involved	BDY BRN < 10%/3D DEG NOS
949	Burn, unspecified	BURN NOS
950	Injury to optic nerve and pathways	OPTIC NERVE INJURY
951	Injury to other cranial nerve(s)	INJURY OCULOMOTOR NERVE
952	Spinal cord injury without evidence of spinal bone injury	C1-C4 SPIN CORD INJ NOS

953	Injury to nerve roots and spinal plexus	CERVICAL ROOT INJURY
954	Injury to other nerve(s) of trunk, excluding shoulder	INJ CERV SYMPATH NERVE
554	and pelvic girdles	
955	Injury to peripheral nerve(s) of shoulder girdle and	INJURY AXILLARY NERVE
000	upper limb	
956	Injury to peripheral nerve(s), of pelvic girdle and	INJURY SCIATIC NERVE
000	lower limb	
957	Injury to other and unspecified nerves	INJ SUPERF NERV HEAD/NCK
958	Certain early complications of trauma	AIR EMBOLISM
959	Injury, other and unspecified	
960	Poisoning by antibiotics	POISONING-PENICILLINS
961	Poisoning by other anti-infectives	POISONING-SULFONAMIDES
962	Poisoning by hormones and synthetic substitutes	POIS-CORTICOSTEROIDS
963	Poisoning by primarily systemic agents	POIS-ANTIALLRG/ANTIEMET
964	Poisoning by agents primarily affecting blood	POISONING-
	constituents	IRON/COMPOUNDS
965	Poisoning by analgesics, antipyretics, and	POISONING-OPIUM NOS
	antirheumatics	
966	Poisoning by anticonvulsants and anti-	POISON-OXAZOLIDINE DERIV
	Parkinsonism drugs	
967	Poisoning by sedatives and hypnotics	POISONING-BARBITURATES
968	Poisoning by other central nervous system	POIS-CNS MUSCLE DEPRESS
	depressants and anesthetics	
969	Poisoning by psychotropic agents	POISON-ANTIDEPRESNT NOS
970	Poisoning by central nervous system stimulants	POISONING-ANALEPTICS
971	Poisoning by drugs primarily affecting the	POIS-
	autonomic nervous system	PARASYMPATHOMIMETIC
972	Poisoning by agents primarily affecting the	POIS-CARD RHYTHM
	cardiovascular system	REGULAT
973	Poisoning by agents primarily affecting the	POIS-ANTACID/ANTIGASTRIC
	gastrointestinal system	
974	Poisoning by water, mineral, and uric acid	POIS-MERCURIAL DIURETICS
	metabolism drugs	
975	Poisoning by agents primarily acting on the smooth	POISONING-OXYTOCIC
	and skeletal muscles and respiratory system	AGENT
976	Poisoning by agents primarily affecting skin and	POIS-LOCAL ANTI-INFECT
	mucous membrane, ophthalmological,	
	otorhinolaryngological, and dental drugs	
977	Poisoning by other and unspecified drugs and	POISONING-DIETETICS
070	medicinal substances	
978	Poisoning by bacterial vaccines	POISONING-BCG VACCINE
979	Poisoning by other vaccines and biological	POISON-SMALLPOX VACCINE
080	substances Toxic effect of alcohol	TOXIC EFF ETHYL ALCOHOL
980		TOXIC EFF ETHYL ALCOHOL
981	Toxic effect of petroleum products	PROD
982	Toxic effect of solvents other than petroleum-based	TOXIC EFFECT BENZENE
983	Toxic effect of corrosive aromatics, acids, and	TOX EFF CORROSIVE
	caustic alkalis	AROMAT

984	Toxic effect of lead and its compounds (including	TX EFF INORG LEAD	
	fumes)	COMPND	
985	Toxic effect of other metals	TOXIC EFFECT MERCURY	
986	Toxic effect of carbon monoxide	TOX EFF CARBON MONOXIDE	
987	Toxic effect of other gases, fumes, or vapors	TOXIC EFF LIQ PETROL GAS	
988	Toxic effect of noxious substances eaten as food	TOXIC EFF FISH/SHELLFISH	
989	Toxic effect of other substances, chiefly	TOXIC EFFECT CYANIDES	
	nonmedicinal as to source		
990	Effects of radiation, unspecified	EFFECTS RADIATION NOS	
991	Effects of reduced temperature	FROSTBITE OF FACE	
992	Effects of heat and light	HEAT STROKE & SUNSTROKE	
993	Effects of air pressure	BAROTRAUMA, OTITIC	
994	Effects of other external causes	EFFECTS OF LIGHTNING	
995	Certain adverse effects not elsewhere classified	ANAPHYLACTIC SHOCK	
996	Complications peculiar to certain specified	MALFUNC CARD DEV/GRF	
	procedures	NOS	
997	Complications affecting specified body systems,	NERVOUS SYST COMPLC	
	not elsewhere classified	NOS	
998	Other complications of procedures, not elsewhere	POSTOPERATIVE SHOCK	
	classified		
999	Complications of medical care, not elsewhere	GENERALIZED VACCINIA	
	classified		

#### How to Log In and Get Started

- Once you have registered and received your confirmation to submit data for the Blood Management Project, you may access the project website at: <u>http://manual.jointcommission.org</u>
- 2) Click on "Login" in the upper right hand corner.

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H O M E	Welcome to the Performance Measurement Network Q&A Forum Published Manuals	
	Joint Commission Only Measures  UPDATED Hospital Based Psychiatric Inpatient Services (HBIPS) and Perinatal Care (PC) Measures (version 2010A2) Original release (version 2010A) Original release (version 2010A1)	CMS and Joint Commission Aligned Measures   • Current Specification Manual for National Hospital Quality Measures  • Future Specification Manual for National Hospital Quality Measures  • Historical Specification Manuals for National Hospital Quality Measures
	Important publications: Dr. Mark Chassin, President of The Joint Commission, recently con <u>Postindustrial Care — The Revolution in Health Care Delivery (<i>New E)</i> <u>January 20, 2010, at NEJM.org)</u>. The article provides a perspective on care that may be of interest to you.</u>	ngland Journal of Medicine, published on

3) Enter your Login and Password and click "ok".

Welcome to the Performance Measurement Network Please enter your username and password.					
Login: Password	Login: testuser50 ** Password: •••••••• ** OK Clear Cancel				
See also: <u>Create Login/Register</u> , <u>Forgot password?</u> Contact <u>SWilliams@jointcommission.org</u> if you have any questions.					

4) Welcome to the Performance Measurement Network. Select the "Blood Mgmt Project" link from the left hand navigation bar.



5) You are now on the Blood Management Project Page. You will see your hospitals(s) listed here. In the Project Help section, you will find a link to the measure specifications, an example of the import file template, and other material intended to assist you with your participation in this project. Please click on the hospital name to enter blood management data.



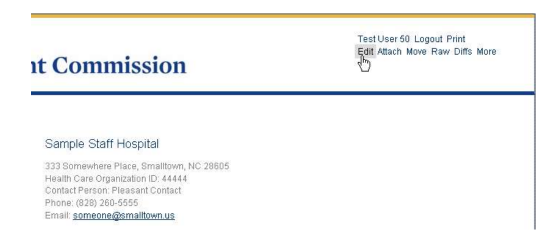
- 6) You are now on your hospital page. From this page, you can:
  - update your hospital demographic information
  - enter new records
  - import new records
  - view and update existing records
  - add RBC, Plasma and Platelet events
  - mark records as "complete"
  - review records that have been completed
  - view import attachments

Each function will be discussed in detail below.



### Navigating the Blood Management Project Data Collection Tool <u>Updating your Hospital Demographic Information</u>

a) To update your hospital's demographic information, click the "Edit" link, Fill out the form that appears, and click the "Save" button at the bottom of the form.



You will be directed to the Edit form, and you can change your hospital's contact details here. Click "Save" to save your changes, or "Cancel" to exit without saving.

Address:	333 Somewhere Place
City:	Smalltown
State:	NC
Zip Code:	28605
Contact Person:	Pleasant Contact
Contact Phone:	(828) 260-5555
Contact Email:	someone@smalltown.us
Save Save and Contin	ue Preview Change form Cancel
d'h	and washeeder washeeden washeeden

#### **Importing Records**

a) To import data, click on the "Import" link on your hospital home page. The template for this import file can be found on the project home page.

#### Import Data

Steps for importing base data set using a properly formatted Excel spreadsheet:

1. Save the file that is to be imported with the EXACT Name: "import.xls".

Click the link planet.x1s" file.

3. Once you have uploaded the file, 👉 Click here to finish the upload process.

a. Once the import has been completed, you will need to click your web browser's "Back" button and then "Refresh" the web page before you will see your new data records.

b) Click on "browse" to find and select your import file (which must be named "import.xls"), and click on "Upload File". You do not need to check the checkboxes, but <u>you may want to add</u> a comment to keep track of your imports (e.g., April 2010 discharges; 51 records)

Attach file to Sample Staff Hospital

File: Comment:	G:11 Web Activities\Wiki\Blood Management Impo
	<ul> <li>Create a link to the attached file at the end of the topic.</li> <li>Hide attachment in normal topic view.</li> </ul>
	Upload file Show all attachments Cancel

c) Once you have uploaded your file, you will need to click on the "Click here" link to finish the upload process. You'll then need to click your browser's "Back" button and "Refresh" your hospital page.

#### Import Data

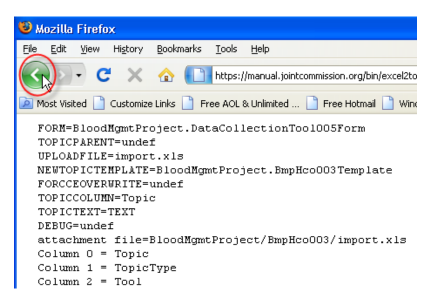
Steps for importing base data set using a properly formatted Excel spreadsheet:

- 1. Save the file that is to be imported with the EXACT Name: "import.xls".
- Click the link: F Import and follow the instructions to select and upload your "import.x1s" file.
- Once you have uploaded the file Click here to finish the upload process.
  - a. Once the import has been completed, you will need to click your web browser's "Back" button and then "Refresh" the web page before you will see your new data records.

d) You may notice a form at the bottom of your hospital page. It displays the most recently imported file. This area will only be used to verify that your import was successful (note the date, time and comments to ensure that it represents the file you imported.

Attachments *					
	Attachment	Action	Size	Date	Who
×	import.xls	props, move	55.0 K	22 Feb 2010 - 08:20	ScottWilliams
	Monday 2/22 tes	st of import			

e) Your uploaded records are shown here (in a rather unappealing format!) and you will need to click on your browser's "Back" button to return to your hospital home page.



f) You are now back on your hospital's home page. Please click on your browser's "Refresh" button to view the records you just imported. Your records have been imported, but you will not be able to see them until the page is refreshed (or you navigate away from it and then back to it).

🕹 Sample Sta	ff Hospital	- BloodMgn	ntProject - Performance Measurement Network - Mozilla Firefox
<u>File E</u> dit <u>V</u> ie	W History	<u>B</u> ookmarks	Tools Help
< <u>&gt;</u> -(	C)X	<b>d</b>	https://manual.jointcommission.org/bin/view/BloodMgmtProject/BmpHco003
🧟 Most Visited	Customize	e Links 📄 Fri	ee AOL & Unlimited 📋 Free Hotmail 📋 Windows Marketplace 📋 Windows I

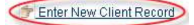
g) Your uploaded files should now viewable in the "Submitted Data" section of your hospital home page.

UBCI	Birthdate	Admitted	Discharged	Completed 🚺
333333	03-03-1983	02-02-2010	02-05-2010	Γ
333331	05-01-2001	01-01-2010	01-10-2010	Γ
555555	04-04-1974	07-04-2009	07-07-2009	Γ
333332	03-03-1983	02-02-2010	02-05-2010	Γ
333335	05-01-2001	01-01-2010	01-10-2010	Γ
1234567	12-30-2008	01-26-2010	02-02-2010	Γ
2223	05/01/01	01/01/10	01/10/10	Γ
333336	03-03-1983	02-02-2010	02-05-2010	Γ
555556	12-09-1970	08-08-2009	08-12-2009	Γ

#### Show all Records (including complete)

#### Navigating the Blood Management Project Data Collection Tool Enter New Records (without using the file import

a) To enter a new record, click on the "Enter New Client Record" link (right below the data record table).



b) You are now viewing the data collection tool for Blood Management. Enter data for the client record. Note: hovering over the green "i" next to a data element will show you the question and allowable values associated with that data element as well as a link to the data element page.

Unique Blinded Case Identifier	12			
Admission Date	and the second			
Bithdate				
Discharge Date	MMOD-YMY			
Discharge Status Selec	a. 💌 🖪			
Sex 🔿 M	0=0+1			
ICD-5-CM Principal Diagnosis Code	11			
ED & CM Other Dagnman Cades				
ICD-9-CM Other Disgnosis Codes				
	Add another respon			
ICD-B-CM Principal Procedure Code				
ICD-9-CM Principal Procedure Code	a			
1	0			
ICD-9-CM Principal Procedure Date	0			
ICD-9-CM Principal Procedure Date				
ICD-9-CM Principal Procedure Date				
ICD-9-CM Principal Procedure Date	T Add anyther response			
ICD-9-CM Principal Procedure Date	Add another response			
ICD-9-CM Principal Procedure Date	Add another response			
ICD-9-CM Principal Procedure Date ICD-9-CM Other Procedure Codes ICD-9-CM Other Procedure Codes ICD-9-CM Other Procedure Dates ICD-9-CM Other Procedure Dates ICD-9-CM Other Procedure Dates ICD-9-CM Other Procedure Dates ICD-9-CM Other Procedure Codes ICD-9-CM Other Procedure Cod	a Add.aoxfbar.rusuear 0 2 0 0 2 0 2 12 0 2 12			
ICD-9-CM Principal Procedure Date ICD 9-CM Other Procedure Codes Date ICD-9-CM Other Procedure Codes ICD-9-CM Other Procedure Codes ICD-9-CM Other Procedure Dates ICD-9-CM Other Procedure Dates ICD-9-CM Other Procedure Codes ICD-9-CM Other Procedur	11 12 13 Add asyther response 0 2 12 0 2 12 0 2 12 0 4 12			
ICD-9-CM Principal Procedure Date ICD-9-CM Other Procedure Codes Date ICD-9-CM Other Procedure Codes ICD-9-CM Other Procedure Dates ICD-9-CM Other Procedure Codes Dates ICD-9-CM Other	11 12 13 Add asyther response 0 2 12 0 2 12 0 2 12 0 4 12			

c) Once you have completed data entry for this record, click on "Save Data Record".

#### Navigating the Blood Management Project Data Collection Tool To View and Update Existing Records

a) There are two ways to view the list of submitted records. The default view is of all incomplete records. If you would like to view all records, including completed (locked) records, click the link "Show all Records (including complete)".

View of the default setting showing a list of only incomplete records:

UBCI	Birthdate	Admitted	Discharged	Completed
333333	03-03-1983	02-02-2010	02-05-2010	F
333331	05-01-2001	01-01-2010	01-10-2010	Г
555555	04-04-1974	07-04-2009	07-07-2009	Г
333332	03-03-1983	02-02-2010	02-05-2010	Г
333335	05-01-2001	01-01-2010	01-10-2010	, F
1234567	12-30-2008	01-26-2010	02-02-2010	Г
2223	05/01/01	01/01/10	01/10/10	Г
333336	03-03-1983	02-02-2010	02-05-2010	Г
555556	12-09-1970	08-08-2009	08-12-2009	Г

View of alternate setting showing list of all records (both incomplete and complete). To return the default setting, click the link "Show Incomplete Records Only"

UBCI	Birthdate	Admitted	Discharged	Completed 🚺
333333	03-03-1983	02-02-2010	02-05-2010	Г
333331	05-01-2001	01-01-2010	01-10-2010	Г
555555	04-04-1974	07-04-2009	07-07-2009	Г
333332	03-03-1983	02-02-2010	02-05-2010	Г
1234567	12-30-2008	01-26-2010	02-02-2010	Г
333335	05-01-2001	01-01-2010	01-10-2010	Г
333336	03-03-1983	02-02-2010	02-05-2010	Г
2223	05/01/01	01/01/10	01/10/10	Г
555558	12-09-1970	08-08-2009	08-12-2009	Г
333334	05-01-2001	01-01-2010	01-10-2010	<b>a</b>
99999999	01-01-1901	11-11-2010	11-15-2010	18
4445	03/03/83	02/02/10	02/05/10	<u> </u>
444555	03/03/83	02/02/10	02/05/10	1
2224	05/01/01	01/01/10	01/10/10	

b) To view or update data in an existing record, click on the UBCI number. This will create a drop down that includes all of the information for that client record. You can contract the drop down by clicking on the "-"or expand by clicking on the "+" before the different sections.

		a Collection I	
234567	12-30-2008	01-26-2010	02-02-2010
🗄 General an	d other patient-level o	lata elements 🥒	
Discharg	ge Status		
Sex			М
ICD-9-C	M Principal Diagnosis	Code	49301
ICD-9-C	M Other Diagnosis Co	des	
ICD-9-C	M Principal Procedure	Code	7301
	M Principal Procedure	D - I -	01-25-2010
ICD-9-C	M Other Procedure Co		
ICD-9-C	M Other Procedure Da	ites	
Transfus	ion Consent		
	in Addressed Risks, B	enefits and Alternat	tives
to Trans			
Elective	Surgery		
	sia Start Date		
Preoper	ative Anemia Screenir	ig Date	
	ative Anemia Screenir		
Preoper	ative Blood Type Testi	ng	
	et Specific Data Elem	ents	
E RBC Eve	1 /		
	d RBC Event record (3	<u>3 left)</u>	
🖻 Plasma	1 1		
	d Plasma Event recor	<u>a (3 len)</u>	
Platelet		1 (0.1-4)	
<u></u>	d Platelet Event record	<u>1 (3 IEII)</u>	

c) To edit the "General and other patient-level data elements", click on the pencil icon.

1234567	12-30-2008	01-26-2010	02-02-2010	
	d other patient-level d	ata element <mark>s 🖉</mark>		
Dischar	ge Status			01
Sex				M
	M Principal Diagnosis	Code		49301
ICD-9-C	M Other Diagnosis Co	des		
ICD-9-C	M Principal Procedure	Code		7301
ICD-9-C	M Principal Procedure	Date		01-25-2010
-ICD-9-C	M Other Procedure Co	des		

d) Make changes to the "General and other patient-level data elements" and click "Save" when you are done.

▼ Form Data	Permissions	
— Draft Data Coll	ection Tool	
	Unique Blinded Case Identifier	1234567
	Admission Date	01-26-2010 MM-DD-YYYY 🚺
	Birthdate	12-30-2008
	Discharge Date	02-02-2010
	Discharge Status	01 🗸 🚺
	Sex	⊙ M 🔿 F 🔿 U 🚺
ICD-9	3-CM Principal Diagnosis Code	49301
- ICD-9-CM Oth	ner Diagnosis Codes ————	
	ICD-9-CM Other Diagnosis Cor	des 🚺
Save Save an	d Continue Preview Cha	nge form Cancel 🔲 New Revision

#### Navigating the Blood Management Project Data Collection Tool <u>Add RBC Events and BM Unit Level Data Elements</u>

a) To add a RBC event (NOTE: you can add up to three RBC events), click on the "Add RBC Event Record" Link.



b) Enter data for RBC Event 1 and click "Save Data Record"

- RBC Event	
	RBC Event ID 🗾 💿 1 🔿 2 🔿 3
	RBC Event Total Doses 🚺
	Clinical Indication For RBCs 🚺 Select 💌
	Pre-transfusion Hemoglobin 🚺
	Pre-transfusion Hematocrit 🚺
	Surgical Procedure 🚺 🔿 1 🔿 2
Save Data Re	cord

c) Data for "RBC Event 1" is now included with this client record. To edit the RBC Event data that you just entered, click on the pencil icon next to the event. To add unit level data for RBC Event 1, click on the "Add BM Unit Level Data Elements Record" link. (NOTE: you can add up to three BM Unit Level Records)

05-01-2001	01-01-2010	01-10-2010	
sure Set Specific Data Elemen			
RBC Event 1 🥒			1
			2
			<u> </u>
Clinical Indication for RBC	s		1
-Pre-transfusion Hemoglob	in		8
-Pre-transfusion Hematocri	t		21
Surgical Procedure			1
🖻 BM Unit Level Data Eleme	nts(s)		
Add BM Unit Level D:	ata Elements re	cord (3 left)	
- 👉 Add RBC Event record (2 le	eft)		
asma Event(s)			
👉 Add Plasma Event record (	<u>3 left)</u>		
atelet Event(s)			
👉 Add Platelet Event record (	<u>3 left)</u>		
	eral and other patient-level dat sure Set Specific Data Elemen BC Event(s) RBC Event 1 2 RBC	eral and other patient-level data elements Sure Set Specific Data Elements BC Event(s) RBC Event 1 RBC Event 1 RBC Event 1 RBC Event Total Doses Clinical Indication for RBCs Pre-transfusion Hemoglobin Pre-transfusion Hematocrit Surgical Procedure BM Unit Level Data Elements(s) CMAdd BM Unit Level Data Elements re Add RBC Event record (2 left) lasma Event(s)	eral and other patient-level data elements sure Set Specific Data Elements BC Event(s) ■ RBC Event 1 ■ RBC Event 1 ■ RBC Event Total Doses ■ Clinical Indication for RBCs ■ Official Indication Hemoglobin ■ Pre-transfusion Hemoglobin ■ Pre-transfusion Hematocrit ■ Surgical Procedure ■ Surgical Procedure ■ Surgical Procedure ■ M Unit Level Data Elements(s) ■ M Unit Level Data Elements record (3 left) ■ Add BM Unit Level Data Elements record (3 left) ■ Add RBC Event record (2 left) ■ Add Plasma Event record (3 left) ■ Add Plasma Event record (3 left)

d) Enter data for the BM Unit Level Record for RBC Event 1 and click "Save Data Record"

BM Unit Level Data Elements	
Transfusion Start Date 🚺	
Transfusion Start Time 🚺	
Transfusion Order 🚺	O Y O N
Patient ID Verification 🚺	○1○2
Vital Sign Monitoring 🚺	0102

Save Data Record

e) Data for "BM Unit 1" for "RBC Event 1" is now included with this client record. To edit the BM unit data that you just entered, click on the pencil icon. To add another BM Unit for RBC Event 1, click on "Add BM Unit Level Data Elements Record" link. To add another RBC Event, click on "Add RBC Event Record".

33331	05-01-2001	01-01-2010	01-10-2010
Measure : RBC Ev B RBC FV B RBC F F - F	nd other patient-level Set Specific Data Elen vent(s) C Event 1 2 RBC Event ID RBC Event Total Doses Clinical Indication for RI Pre-transfusion Hemog	nents BCs	1 2 1 8
	Pre-transfusion Hemato		21
	Surgical Procedure		1
	BM Unit Level Data Eler BM Unit Level Data E Transfusion Start Transfusion Start	ilements 1 🖉 Date	03-03-20 11:00 Y
	Transfusion Orde	[	Y
	-Patient ID Verifica -Vital Sign Monitori		1
	Add BM Unit Level	Data Elements reco	ord (2 left) 🛶 🔤
5 A	dd RBC Event record (	entral production of the	
- TA	a Event(s) Idd Plasma Event reco It Event(s)	rd (3 left)	
the second se	rt Event(s) Idd Platelet Event recor	rd (3 left)	

## Navigating the Blood Management Project Data Collection Tool Add Plasma Events and BM Unit Level Data Elements

a) To add a Plasma event, click on the "Add Plasma Event Record" Link



b) Enter data for Plasma Event 1 and click "Save Data Record"

Plasma Event	
Plasma Event	ID 🚺 🔿 1 🔿 2 🔿 3
Plasma Event Total Dos	es 🚺
Clinical Indication For Plasn	na 🚺 Select 💙
Pre-transfusion Laboratory Testin	ng 🚺 🔿 1 🔿 2 🔿 3 🔿 4 🔿 5

Save Data Record

c) Data for "Plasma Event 1" is now included with this client record. To edit the Plasma Event data that you just entered, click on the pencil icon next to the event. To add unit level data for Plasma Event 1, click on the "Add BM Unit Level Data Elements Record" link. (NOTE: you can add up to three BM Unit Level Records)

333331	05-01-2001	01-01-2010	01-10-2010	
	her patient-level dat pecific Data Elemen ;)			
🖻 Plasma Eve	nt(s)			
⊡ Plasma E Plasm	Event 1 🖉 🛛			1
Plasm	a Event Total Doses			2
Clinica	al Indication for Plasr	ma		1
Pre-tra	Insfusion Laboratory	/ Testing		2
⊟ -BM_Un	it Level Data Elemei	nts(s)		
31	Add BM Unit Level Da	ata Elements record	<u>(3 left)</u>	
👉 Add Pl	<u>asma Event record (</u>	2 left)		
🖻 Platelet Eve	nt(s)			
👉 Add Pl	atelet Event record (3	<u>3 left)</u>		

d) Enter data for the BM Unit Level Record for Plasma Event 1 and click "Save Data Record"

Transfusion Start Date 👔 Transfusion Start Time 👔 Transfusion Order 👔 O Y O N Patient ID Verification 👔 O 1 O 2 Vital Sign Monitoring 👔 O 1 O 2	- BM Unit Level Data Elements	
Transfusion Order 🚺 O Y O N Patient ID Verification 🚺 O 1 O 2	Transfusion Start Date 👔	
Patient ID Verification 1 0 2	Transfusion Start Time 🚺	
	Transfusion Order 🚺	O Y O N
Vital Sign Monitoring 🚺 🔘 1 🔘 2	Patient ID Verification 🚺	○1○2
	Vital Sign Monitoring 🚺	○1○2

e) Data for "BM Unit Level 1" for "Plasma Event 1" is now included with this client record. To edit the BM unit data that you just entered, click on the pencil icon. To add another BM Unit for Plasma Event 1, click on "Add BM Unit Level Data Elements Record" link. To add another Plasma Event, click on "Add Plasma Event Record".

333331	05-01-2001	01-01-2010	01-10-2010	
🖃 Measur	and other patient-level d e Set Specific Data Eleme			
	Event(s)			
	ma Event(s) asma Event 1 🥒			
	Plasma Event ID			1
	-Plasma Event Total Dose	: S		2
		sma		1
	-Pre-transfusion Laborato	ry Testing		2
	BM Unit Level Data Elem			
	🖻 BM Unit Level Data Ele			
	Transfusion Start D	ate		03-03-2010
	Transfusion Start T	ime		11:00
	Transfusion Order			Y
	Patient ID Verificati			1
	Vital Sign Monitorin	a		1
	Add BM Unit Level (	Data Elements rec	ord (2 left) 🔶	
	Add Plasma Event record	l (2 left) 🔶		
🖃 Plate	elet Event(s)			
	Add Platelet Event record	(3 left)		

#### Navigating the Blood Management Project Data Collection Tool <u>Add Platelet Events and BM Unit Level Data Elements</u>

a) To add a Platelet event, click on the "Add Platelet Event Record" Link



b) Enter data for Platelet Event 1 and click "Save Data Record"

Platelet Event	
Platelet Event ID 🚺	010203
District Event Tatal Darras	
Platelet Event Total Doses 🚺	
Clinical Indication For Platelets 🚺	Select 🔽
Pre-transfusion Platelet Count 🚺	
	0400
Pre-transfusion Platelet Testing 🚺	0102



c) Data for "Platelet Event 1" is now included with this client record. To edit the Platelet Event data that you just entered, click on the pencil icon next to the event. To add unit level data for Platelet Event 1, click on the "Add BM Unit Level Data Elements Record" link. (NOTE: you can add up to three BM Unit Level Records)

333331	05-01-2001	01-01-2010	01-10-2010	
⊡ Measur ⊡ RBC I	and other patient-level da e Set Specific Data Eleme Event(s) na Event(s)			
⊡- <b>Plate</b> ⊡-Pl	Ilet Event(s) atelet Event 1 2 Platelet Event ID			1
	-Platelet Event Total Dose -Clinical Indication for Plat			3
	Pre-transfusion Platelet C	ount		100
	BM Unit Level Data Eleme	ents(s)	ord (3 left)	
23	Add Platelet Event record	(2 left)		

d) Enter data for the BM Unit Level Record for Platelet Event 1 and click "Save Data Record"

	BM Unit Level Data Elements
	Transfusion Start Date 🚺
	Transfusion Start Time 🚺
	Transfusion Order 🚺 🔘 Y 🔘 N
	Patient ID Verification 🚺 🔘 1 🔘 2
	Vital Sign Monitoring 🚺  ○ 1 ○ 2
(	Save Data Record

e) Data for "BM Unit Level 1" for "Platelet Event 1" is now included with this client record. To edit the BM unit data that you just entered, click on the pencil icon. To add another BM Unit for Platelet Event 1, click on "Add BM Unit Level Data Elements Record" link. To add another Platelet Event, click on "Add Platelet Event Record".

333331	05-01-2001	01-01-2010	01-10-2010	
🖃 Meası	al and other patient-level da Ire Set Specific Data Eleme			
<b>⊕</b> Pla	C Event(s) sma Event(s) telet Event(e)			
	telet Event(s) Platelet Event 1 🖉 			1
	Platelet Event Total Doses	) 		
	Clinical Indication for Plate Pre-transfusion Platelet C			100
	Pre-transfusion Platelet To	esting		1
	BM Unit Level Data Element     Transfusion Start Data     Transfusion Start Til     Transfusion Order     Patient ID Verificatio     Vital Sign Monitoring     Add BM Unit Level D	ments 1 🖉 ate me n	:ord (2 left)	Ŷ
	FAdd Platelet Event record (	( <u>2 left)</u>		

#### Marking Records As "Complete"

a) Once you are done entering and editing data for a record, you will need to mark the record as complete. **Please note: Once you check the box for a record under "Complete" you are BOTH marking the record as complete AND locking that record for any further editing.** When you click on the checkbox, the record will "disappear" from view. Do not be alarmed. The default view of the table is to only show incomplete records. To view the record you just completed, click on the link to "Show all Records (including complete)"

UBCI	Birthdate	Admitted	Discharged	Completed 🚺
333333	03-03-1983	02-02-2010	02-05-2010	Г
333331	05-01-2001	01-01-2010	01-10-2010	Г
555555	04-04-1974	07-04-2009	07-07-2009	Г
333332	03-03-1983	02-02-2010	02-05-2010	Г
333335	05-01-2001	01-01-2010	01-10-2010	Г
1234567	12-30-2008	01-26-2010	02-02-2010	F
2223	05/01/01	01/01/10	01/10/10	Г
333336	03-03-1983	02-02-2010	02-05-2010	Г
555556	12-09-1970	08-08-2009	08-12-2009	<b>—</b>

# **Reviewing Records That Have Been Completed**

a) To review a record that has been marked complete, switch the view on your hospital home page by clicking on the "Show all Records (including complete)" link.

Submitted Data	
Show all Records (including complete)	

b) In this view you can see all records both complete and incomplete. Completed records are now LOCKED and can not be edited.

UBCI	Birthdate	Admitted	Discharged	Completed 🚺
333333	03-03-1983	02-02-2010	02-05-2010	Г
333331	05-01-2001	01-01-2010	01-10-2010	Г
555555	04-04-1974	07-04-2009	07-07-2009	Г
333332	03-03-1983	02-02-2010	02-05-2010	Г
1234567	12-30-2008	01-26-2010	02-02-2010	Г
333335	05-01-2001	01-01-2010	01-10-2010	Г
333336	03-03-1983	02-02-2010	02-05-2010	Г
2223	05/01/01	01/01/10	01/10/10	Г
555556	12-09-1970	08-08-2009	08-12-2009	Г
333334	05-01-2001	01-01-2010	01-10-2010	> 🔒
99999999	01-01-1901	11-11-2010	11-15-2010	<b>e</b>
4445	03/03/83	02/02/10	02/05/10	<b>e</b>
444555	03/03/83	02/02/10	02/05/10	<b>e</b>
2224	05/01/01	01/01/10	01/10/10	<b>a</b>

Show incomplete Records Only

b) If, for any reason, you need to unlock a record, you will need to send an e-mail to the project leader, Harriet Gammon. To send your e-mail request, click on the "lock" icon, and an e-mail form should appear. It will be addressed to Harriet, and the subject line will contain a reference to the specific record.

🛄 То	Gammon, Harriet
🛄 Cc	
Subject:	Request to unlock record BloodMgmtProject/RecBmpHco003C333334L0D40188

c) In your e-mail, please briefly explain why the record needs to be unlocked (e.g., Accidentally clicked the "Complete" checkbox).

#### PATIENT BLOOD MANAGEMENT PERFORMANCE MEASURES PROJECT - Technical Advisory Panel

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