

Hemovigilance Module - Annual Facility Survey Acute Care Facility

*Requ	ired for saving	-						
*Fac	cility ID#: *Survey Year:							
For a	all questions, use information	from previous full calendar year.						
NOT	ility Characteristics <u>E:</u> Questions 1 – 7 are comp previous year's survey.	leted automatically (i.e., auto-populated) in the NHSN application with responses from						
*1.	Ownership: (check one)							
	Government Milit For profit Vete	aryNot for profit, including church eran's AffairsPhysician-owned						
*2.	If Yes, check	hospital for physicians and/or physicians-in-training? Yes No						
*3.	Community setting of facility	r: Urban Suburban Rural						
*4.	*4. How is your hospital accredited? (check one)							
	The Joint Commission	American Osteopathic Association (AOA) creditation for Healthcare Organizations (DNV) Other Accrediting Organization						
*6. year	Total beds served by the tr Number of surgeries perfor At what trauma level is you	med per Inpatient: Outpatient:						
	nsfusion Service Charac	,						
		cility areas served by the transfusion service: (check all that apply)						
0.	Cancer center	Orthopedic General medical and surgical						
	Children's cancer cent							
	Chronic disease	Burn center Obstetrics/Gynecology						
	disease	Trauma/Emergency Other (specify)						
*9.	Does your healthcare facili	ty provide all of its own transfusion services, including all laboratory functions?						
	Yes No, we contract with a blood center for some transfusion service functions.							
	No, we contract with a	nother healthcare facility for some transfusion service functions.						
*10.	Is the transfusion service p	art of the facility's core laboratory? Yes 🗌 No						
*11.	2	fusion service staff members are there? (Count full-time equivalents; include supervisors.) dical Technologists: Medical Laboratory Technicians:						
Assurance guarantee	of Confidentiality: The voluntarily provided that it will be held in strict confidence, will be	dedicated position or FTE in a guality or patient safety information obtained in this surveillance system that would permit identification of any individual or institution is collected with a used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the 18(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)). CDC 57.300 Rev. 9, v9.2						
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Public reporting burden of this collection of information is estimated to average 86 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering, and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).

Image: Safety Network Form Approved OMB No. 0920-0666 OMB No. 0920-0666 Exp. Date: 12/31/2026 Exp. Date: 12/31/2026 www.cdc.gov/nhsn www.cdc.gov/nhsn							
function (e.g., TSO) for investigation of transfusion-related adverse reactions?							
*13. Does your hospital have a dedicated position or FTE in a <u>quality or patient safety</u> <u>function (e.g., TSO) for investigation of transfusion errors (i.e., incidents)?</u>							
*14. Is the transfusion service laboratory accredited? Yes No							
If Yes, select all that apply: College of American Pathologists (CAP) AABB TJC							
*15. Does your facility have a committee that reviews blood utilization? Yes No							
*16. Total number of patient samples collected for type and screen or crossmatch:							
*17. Are any of the following issued through the transfusion service? (check all that apply)							
Albumin Factors (VIIa, VIII, IX, ATIII, etc.) Immunoglobulin (IV)							
Immunoglobulin (IM or subcutaneous)							
*18. Does your facility attempt to transfuse only leukocyte-reduced or leuko-poor cellular components? Yes No							
*19. Are all units stored in the transfusion service? Yes No							
If No, indicate the location(s) of satellite storage: (check all that apply)							
Ambulatory Care Cancer Center Cardiac ICU							
Emergency Department Labor and Delivery Medical Flight Facility							
Operating Room Other: (specify)							
*20. To what extent does the transfusion service modify products? (check all that apply)							
Aliquot Deglycerolizing Irradiation Leukoreduction							
reduction Pooling Washing None of these							
*21. Do you collect blood for transfusion at your facility? Yes No							
If Yes, check all that apply: Allogeneic Autologous Directed							
*22. Does your facility perform viral testing on blood for transfusion? Yes No							
*23. Does your facility perform point-of-issue bacterial testing on platelets prior to transfusion? Yes No							
Transfusion Service Computerization							
*24. Is the transfusion service computerized? Yes No (If No, skip to next section)							
If Yes, select system(s) used: (check all that apply) BBCS [®] BloodTrack Tx [®] (Haemonetics)							
Cerner Classic [®] Cerner Millennium [®] HCLL [®] Horizon BB [®] Hemocare [®]							
Lifeline [®] Meditech [®] Misys [®] Safetrace Tx [®] (Haemonetics) Softbank [®]							
Western Star [®] Other (specify)							
*25. Is the system ISBT-128 compliant?							
*26. Does the transfusion service system interface with the patient registration system?							
*27. Are the transfusion service adverse events entered into a hospital-wide electronic reporting system?							
Yes No If Yes, specify system used:							



*28. Does your facility use positive patient ID technology for the transfusion service?

 Yes, hospital wide Yes, certain areas Not used If Yes, select purpose(s): (check all that apply) Specimen collection Product administration If Yes, select system(s) used: (check all that apply) Mechanical barrier system (e.g., Bloodloc[®]) Separate transfusion ID wristband system (e.g., Typenex[®]) 								
Radio frequency identification (RFID) Bedside ID band barcode scanning Other (specify)								
*29. Does your facility have physician online order entry for test requesting? Yes No								
*30. Does your facility have physician online order entry for product requesting? Yes No								
Transfusion Service Specimen Handling and Testing								
*31. Are transfusion service specimens drawn by a dedicated phlebotomy team?								
Always Sometimes, approximately% of the time Never								
*32. What specimen labels are used at your facility? (check all that apply)								
Handwritten Addressograph Computer generated from laboratory test request								
Computer generated by bedside device Other (specify)								
*33. Are phlebotomy staff members allowed to correct patient identification errors on pre-transfusion specimen labels?								
Yes No								
*34. What items can be used to verify patient identification during specimen collection and prior to product administration at your facility? (check all that apply)								
Medical record (or other unique patient ID) number Date of birth								
Gender Gender identity Sex at birth								
Patient first name Patient last name Transfusion specimen ID system (e.g., Typenex®)								
Patient verbal confirmation of name or date of birth Other (specify)								
*35. How is routine type and screen done? (check all that apply and estimate frequency of each)								
Manual technique % Automated technique %								
Both automated and manual technique % Total should equal 100%								
*36. Is the ABO group of a pre-transfusion specimen routinely confirmed?								
If Yes, check one:								
All samples								
If there is no laboratory record of previous determination of patient's ABO group								
If there is no laboratory record of previous determination of patient's ABO group AND the patient is a								
candidate for electronic crossmatching								

If Yes, is the confirmation required on a separately-collected specimen before a unit of Group A, B or AB red $_{\mbox{Page 3 of 4}}$



blood cells is issued for transfusion? Yes No

*37. How many RBC type and screen and crossmatch procedures were performed at your facility by any method?

RBC type and screen: _____ RBC crossmatch

Estimate the % of crossmatch procedures done by each method: (check all that apply)

Electronically	%	Serologically	%	Don't know	Total may be >100%
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