

Hemovigilance Module Adverse Reaction Post Transfusion Purpura

*Required for saving *Facility ID#: NHSN Adverse Reaction #: **Patient Information** F Other *Date of Birth: / / *Gender: M *Patient ID: Secondary ID: _____ Medicare #: _____ Social Security #: _____ Middle Name: Last Name: First Name: **Ethnicity** Hispanic or Latino Not Hispanic or Not Latino Race American Indian/Alaska Native Asian Black or African American White Native Hawaiian/Other Pacific Islander O-A+ | | B-AB+ O+ Blood type not done *Blood Group: Patient Medical History (Use worksheet on page 4 for additional codes and descriptions.) (part 1) List the patient's admitting diagnosis. (Use ICD-10 Diagnostic codes/descriptions) Description: Description: Code: _____ Description: Code: (part 2) List the patient's underlying indication for transfusion. (Use ICD-10 Diagnostic codes/descriptions) Code: Description: Code: _____ Description: Code: Description: **UNKNOWN** (part 3) List the patient's comorbid conditions at the time of the transfusion related to the adverse reaction. (Use ICD-10 Diagnostic codes/descriptions) NONE Code: _____ Description: _____ Code: _____ Description: Description: Code: Continued >> Assurance of Confidentiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or institution is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)). Public reporting burden of this collection of information is estimated to average 20 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 D-74, Atlanta, GA 30333 ATTN: PRA (0920-0666). CDC 57.314 (Front) Rev 1, v8.8



Patient Medical History (Use worksheet on page 4 for additional codes and descriptions.)
(part 4) List the patient's relevant medical procedure including past procedures and procedures to be performed during the current hospital or outpatient stay. (Use ICD-10 Procedure codes/descriptions)
Code: Description:
Code: Description:
Code: Description:
(part 5) Additional Information
Transfusion History (Use worksheet on page 4 for additional transfusion history.)
*Has the patient received a previous transfusion? YES NO UNKNOWN
**If yes, provide information about the transfusion event. If not, skip to Reaction Details section.
Blood Product: WB RBC Platelet Plasma Cryoprecipitate Granulocyte
Date of Transfusion:// UNKNOWN
Did the patient experience a transfusion adverse reaction?
If yes, provide information about the transfusion adverse reaction.
Type of transfusion adverse reaction: Allergic AHTR DHTR DSTR FNHTR
HTR TTI PTP TACO TAD TA-GVHD TRALI UNKNOWN
OTHER Specify
Reaction Details
*Date reaction occurred:/ *Time reaction occurred:: Time unknown
*Facility location where patient was transfused:
*Is this reaction associated with an incident?
After recognition of the transfusion reaction, was the current transfusion:
Continued Stopped and restarted Stopped indefinitely
Investigation Results
* Post transfusion purpura (PTP)
*Case Definition
Check all that occurred after cessation of transfusion :
Alloantibodies in the patient directed against HPA or other platelet specific antigen detected at or after development of thrombocytopenia.
Thrombocytopenia (i.e., decrease in platelets to less than 20% of pre-transfusion count).
Decrease in platelets to levels between 20% and 80% of pre-transfusion count.
Continued >>



Investigation Results (cor	itinued)									
Indicate the case definition (check all that apply):										
PTP is suspected, but laboratory findings and/or information are not sufficient. NOTE: For example, the patient has a drop in platelet count to less than 80% of pre-transfusion count but HPA antibodies were not tested or were negative.										
None of the above										
Other signs and symptoms: (c	heck all that apply)									
Generalized:	Chills/rigors Fever Nausea/vomiting									
Cardiovascular:	Blood pressure decrease Shock									
Cutaneous:	Edema Flushing Jaundice									
	Other rash Pruritus (itching) Urticaria (hives)									
Hemolysis/Hemorrhage:	Disseminated intravascular coagulation Hemoglobinemia									
	Positive antibody screen									
Pain:	Abdominal pain Back pain Flank pain pain									
Renal:	Hematuria Hemoglobinuria Oliguria									
Respiratory:	Bilateral infiltrates on chest x-ray Bronchospasm Cough									
Tespiratory.	Hypoxemia Shortness of breath									
Other: (specify)										
<u>*Severity</u>										
Did the patient receive or e	xperience any of the following? (Response definitions listed in protocol)									
Symptomatic treatm	ent only Hospitalization, inlcuding prolonged hospitalization									
Life-threatening rea	ction Disability and/or incapacitation									
Congenital anomaly	or birth defect(s) of the fetus									
Other medically imp	ortant conditions Unknown or not stated									
<u>*Imputability</u>										
	elationship between the transfusion and the reaction?									
Patient has no other	conditions to explain thrombocytopenia.									
There are other potential causes present that could explain thrombocytopenia, but transfusion is the most likely cause.										
Alternate explanation	ns for thrombocytopenia are more likely, but transfusion cannot be ruled out.									
Evidence is clearly in	favor of a cause other than the transfusion, but transfusion cannot be excluded.									
There is conclusive e	vidence beyond reasonable doubt of a cause other than the transfusion.									
The relationship betw	een the adverse reaction and the transfusion is unknown or not stated.									
Did the transfusion occur at	your facility? YES NO									
When did the reaction occur in relation to the transfusion?										
Occurred 5-12 days post-transfusion										
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Occurred less than 5 or more than 12 days post-transfusion

Continued >>



Investigation Results (continued)		
Designations for case definition, severity, and imputability will be automatically a based on responses in the corresponding investigation results section above.	ssigned in the NF	ISN application
Do you agree with the case definition designation?	YES	NO
Please indicate your designation		
Do you agree with the severity designation?	YES	NO
Please indicate your designation		
Do you agree with the imputability designation?	YES	NO
Please indicate your designation		
Additional Information		
Patient Treatment		
*Did the patient receive treatment for the transfusion reaction?	NO [UNKNOWN
If yes, select treatment(s):		
Medication (Select the type of medication)		
Antipyretics Antihistamines Inotropes/Vasopressors	Bronchodilator	Diuretics
Intravenous		
	rticosteroids	Antibiotics
Antithymocyte globulin Cyclosporin H1 receptor	blockers	Other
Volume resuscitation (Intravenous colloids or crystalloids)		
Respiratory support (Select the type of support)		
	ygen	
Renal replacement therapy (Select the type of therapy)	I I a a a Changa	
Hemodialysis Peritoneal Continuous Veno-Venous	Hemotilitration	
Phlebotomy Other Specify:		
Other Specify.		
Outcome		
*Outcome: Death Major or long-term sequelae sequelae		Not determined
*Outcome: Death Major or long-term sequelae sequelae Date of Death: / /		Not determined
^*If recipient died, relationship of transfusion to death:		
	d Out Not	determined
Cause of death:		
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Was an autopsy performed? Yes No

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Component Details (Use worksheet on page 4 for additional units.)									
*Was a partic	ular unit implicated ir		ble for) the	advers	se reaction?	Yes	s 🗌	No	N/A
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	Unit numbe	r	*Unit expiration Date/Time	on *Blood group of unit			Implic ated Unit?
^IMPLICATED UNIT									
	ISBT-128 Codabar	Entire unit Partial unitmL		_		A- B+	A+ AB- O+	B- AB+ N/A	Y
!! !! :	ISBT-128 Codabar	Entire unit Partial unitmL				A- B+	A+	B- AB+ N/A	N
	ISBT-128 Codabar	Entire unit Partial unitmL				A- B+	A+	B- AB+ N/A	N
Custom Field	ds								
Label			Labe	el					
		<u>'</u>	- - -			_			
Comments									



Hemovigilance Module Additional Worksheet

(part 1) List the pa	ient's admitting diagnosis. (Use ICD-10 Diagnostic codes/descriptions)
Code:	Description:
Code:	Description:
Code:	Description:
Code:	
Code:	Description:
Code:	
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Code:	Description:
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adverse reaction. (Code: Code:	Description:
adverse reaction. (Code: Code:	Description:
adverse reaction. (Code: Code:	Description:



Hemovigilance Module Additional Worksheet

Transfusion History
Has the patient received a previous transfusion?
**If yes, provide information about the transfusion event. If not, skip to Reaction Details section.
Blood Product: WB RBC Platelet Plasma Cryoprecipitate Granulocyte
Date of Transfusion:// UNKNOWN
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If yes, provide information about the transfusion adverse reaction.
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OTHER Specify



Hemovigilance Module Additional Worksheet

Component Details							
*Was a particul	ar unit implicated in ((i.e., responsi	ible for) the adverse read	ction? Y	es No N	I/A	
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	Unit number	*Unit expiration Date/Time	*Blood group of unit	Implic ated Unit?	
	ISBT-128 Codabar	Entire unit Partial unit mL		<i>l 1</i>	A- A+ B- B AB- AB+ O- O+ N/A	N	
	ISBT-128 Codabar	Entire unit Partial unit mL			A- A+ B- B- AB- AB+ O- O+ N/A	N	
	ISBT-128 Codabar	Entire unit Partial unit mL			A- A+ B- B- AB- AB+ O- O+ N/A	N	
	ISBT-128 Codabar	Entire unit Partial unitmL			A- A+ B- B- AB- AB+ O- O+ N/A	N	
	ISBT-128 Codabar	Entire unit Partial unit mL	 	<i>I. I.</i>	A- A+ B- B- AB- AB+ O- O+ N/A	N	
	ISBT-128 Codabar	Entire unit Partial unitmL		<u> </u>	A- A+ B- B AB- AB+ O- O+ N/A	N	
	SBT-128	Entire			A+ B-	IN	

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	unit Partial unit mL		A- + -		AB+	