

Hemovigilance Module Adverse Reaction

Post Transfusion Purpura *Required for saving

*Facility ID#: NHSN Ad	verse Reaction #:					
Patient Information						
*Patient ID:	*Gender: M F Oth	ner *Date of Birth://				
Social Security #:	Secondary ID:	Medicare #:				
Last Name:	First Name:	Middle Name:				
Ethnicity Hispanic or Latino	☐ Not Hispanic or Not Latino					
Race	a Native 🗌 Asian 🔲 Bl	lack or African American				
☐ Native Hawaiian/Other Pacific Islander ☐ White						
*Blood Group: ☐ A- ☐ A+ ☐ B-	□B+ □ AB- □ AB+ □ O	- ☐ O+ ☐ Blood type not done				
☐ Transitional ABO / R	h + Transitional ABO / Rh -	☐ Transitional ABO / Transitional Rh				
☐ Group A/Transitional Rh ☐ Group I	3/Transitional Rh 🔲 Group O/Transi	itional Rh Group AB/Transitional Rh				
Patient Medical History						
List the patient's admitting diagnos	s. (Use ICD-10 Diagnostic codes/de	escriptions)				
Code:	Description:					
Code:	Description:					
Code:	Description:					
List the patient's underlying indicati	on for transfusion. <i>(Use ICD-10 Dia</i>	gnostic codes/descriptions)				
Code:	Description:					
Code:	Description:					
Code:	Description:					
List the patient's comorbid condition reaction. (Use ICD-10 Diagnostic c	ns at the time of the transfusion related ones/descriptions)	ted to the adverse UNKNOWN NONE				
Code:	Description:					
Code:	Description:					
Code:	Description:					
of any individual or institution is collected wit stated, and will not otherwise be disclosed of Sections 304, 306 and 308(d) of the Public Fublic reporting burden of this collection of in reviewing instructions, searching existing data collection of information. An agency may no unless it displays a currently valid OMB continuous contents.	h a guarantee that it will be held in strict or released without the consent of the individent of the	nd 242m(d)).				



	dical procedure including past procedures and procedures to be hospital or outpatient stay. (Use ICD-10 Procedure NONE				
Code:	Description:				
Code:	Description:				
Code:	Description:				
Additional Information					
Transfusion History					
Has the patient received a pr	evious transfusion?				
Blood Product:					
Date of Transfusion:	/				
Was the patient's adverse i	reaction transfusion-related?				
If yes, provide information a	about the transfusion adverse reaction.				
Type of transfusion adverse					
	PTP TACO TAD TA-GVHD TRALI UNKNOWN				
OTHER Specify					
Reaction Details					
*Date reaction occurred:/_	/ *Time reaction occurred::				
*Facility location where patier					
Is this reaction associated with a	an incident? Yes No If Yes, Incident #:				
Investigation Results					
Investigation Results *☐ Post transfusion purpura	(PTP)				
	(PTP)				
* Post transfusion purpura *Case Definition Check all that occurred af	ter cessation of transfusion : atient directed against HPA or other platelet specific antigen detected at or after				
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*Case Definition Check all that occurred af Alloantibodies in the p development of thror Thrombocytopenia (i.e. Decrease in platelets Check all that apply: PTP is suspected, but patient has a drop in p	ter cessation of transfusion: latient directed against HPA or other platelet specific antigen detected at or after imbocytopenia. e., decrease in platelets to less than 20% of pre-transfusion count). to levels between 20% and 80% of pre-transfusion count. laboratory findings and/or information are not sufficient. NOTE: For example, the platelet count to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not to less than 80% of pre-transfusion count but HPA antibodies were not the less than 80% of pre-transfusion count but HPA antibodies were not the less	t			
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	Renal:	☐ Hematuria	☐ Hemoglobinuria ☐ Oli	guria
	Respiratory:	☐ Bilateral infiltrates on che	est x-ray	☐ Cough
		Hypoxemia	☐ Shortness of breath	
	Other: (specify)			
	*Severity			
	Did the patient receive or e	xperience any of the following	1?	
	☐ No treatment require	ed 🗌 Syr	nptomatic treatment only	
		ıding prolonged hospitalization	_	
	Disability and/or inca	· <u> </u>	ngenital anomaly or birth defect(s)	
	Other medically impo	ortant conditions	ath Unknown or no	t stated
	*Imputability			
		elationship between the transf		
		conditions to explain thrombo	• •	
	There are other poter likely cause.	ntial causes present that could	d explain thrombocytopenia, but tr	ansfusion is the most
	Alternate explanation	s for thrombocytopenia are m	ore likely, but transfusion cannot l	be ruled out.
	Evidence is clearly in	favor of a cause other than th	e transfusion, but transfusion can	not be excluded.
	☐ There is conclusive ev	vidence beyond reasonable de	oubt of a cause other than the trai	nsfusion.
	☐ The relationship between	een the adverse reaction and	the transfusion is unknown or not	stated.
	Did the transfusion occur at	your facility?	□NO	
	When did the reaction occu	r in relation to the transfusion	?	
	Occurred 5-12 days p	oost-transfusion		
	Occurred less than 5	or more than 12 days post-tra	nsfusion	
	dule-generated Design			
		efinition, severity, and imputal in the corresponding investig	oility will be automatically assigned ation results section above.	d in the NHSN
	*Do you agree with the ca	ase definition designation?	☐ YES	□NO
	^Please indicate your designation	gnation		
	*Do you agree with the se	everity designation?	☐ YES	□NO
	*Please indicate your designate	gnation		
	*Do you agree with the <u>in</u>		☐ YES	□NO
Do	^Please indicate your designment	gnation		_
	•	nent for the transfusion reaction	n? YES NO	UNKNOWN
	If yes, select treatment(s):	a type of modication)		
	☐ Medication (Select the	<u> </u>	oc//acoprocess Dranshadila	tor Divretice
	<u> </u>		es/Vasopressors	
	☐ Intravenous imm	nunoglobulin Intravenous	steroids Corticosteroids Other	Antibiotics
	☐ Antitriymocyte g	lobulin		



☐ Volu	me resuscitation (Intr	avenous colloid	s or c	rystalloids)					
Res	piratory support <i>(Sele</i>] Mechanical ventilati	<u> </u>) e ventilation	☐ Oxygen	l			
 □ Renal replacement therapy (Select the type of therapy) □ Hemodialysis □ Peritoneal □ Continuous Veno-Venous Hemofiltration 									
☐ Phlebotomy									
Other Specify:									
Outcome *Outcome: ☐ Death ☐ Major or long-term sequelae ☐ Minor or no sequelae ☐ Not determined									
*Outcome: Date of		ajor or long-tern	n sequ	ieiae _	Minor or no se	queiae		ot detern	ninea
	ecipient died, relation	/ nship of transfus	ion to	death:					
	Definite Probable	·		Doubtful	☐ Ruled Ou	t [] Not de	etermine	ed
Cause	of death:								
Was an	autopsy performed?	☐ Yes	☐ No)					
Component									
*Was a partion?	cular unit implicate	d in (i.e., respo			adverse	☐ Ye	s 🗌	No [□ N/A
Transfusion Start and End	*Component code	Amount transfused at	(Requ	t number uired for ion and	*Unit expiration	*Bloc	od grou	р	Implicate d
Date/Time	(check system used)	reaction onset	TRAL	.l)	Date/Time	of un	it		Unit?
Date/Time ^IMPLICATED	(check system used)	reaction onset		l)		of un	it		Unit?
	(check system used)	reaction onset				of un	it		Unit?
	(check system used) UNIT	☐ Entire unit		. — — —		of un	A+	□ B-	
	(check system used) UNIT ISBT-128			. — — —				□ B-	Y
	(check system used) UNIT ISBT-128	☐ Entire unit		. — — — . — — — —		□ A-	□ A+		
	(check system used) UNIT ISBT-128	☐ Entire unit		. — — — — . . — — — — .		□ A-	□ A+	☐ AB+	
	(check system used) UNIT ISBT-128 Codabar	☐ Entire unit ☐ Partial unit ☐ mL ☐ Entire unit		 		□ A-	□ A+	☐ AB+	Y
	(check system used) UNIT ISBT-128 Codabar ISBT-128	☐ Entire unit ☐ Partial unit ☐ mL				□ A- □B+ □ O-	□ A+ □ AB- □ O+	□ AB+	
	(check system used) UNIT ISBT-128 Codabar ISBT-128	☐ Entire unit ☐ Partial unitmL ☐ Entire unit ☐ Partial unit				□ A- □B+ □ O-	□ A+ □ AB- □ O+	□ AB+ □ N/A □ B-	Y
	UNIT ISBT-128 Codabar ISBT-128 Codabar Codabar	☐ Entire unit ☐ Partial unitmL ☐ Entire unit ☐ Partial unit				□ A- □B+ □ O- □ A- □B+	□ A+ □ AB- □ O+ □ A+ □ AB-	□ AB+ □ N/A □ B- □ AB+	Y
*IMPLICATED	UNIT ISBT-128 Codabar ISBT-128 Codabar Codabar	☐ Entire unit ☐ Partial unitmL ☐ Entire unit ☐ Partial unit		Label		□ A- □B+ □ O- □ A- □B+	□ A+ □ AB- □ O+ □ A+ □ AB-	□ AB+ □ N/A □ B- □ AB+	Y
^IMPLICATED //://://:	UNIT ISBT-128 Codabar ISBT-128 Codabar Codabar	☐ Entire unit ☐ Partial unitmL ☐ Entire unit ☐ Partial unit				□ A- □B+ □ O- □ A- □B+	□ A+ □ AB- □ O+ □ A+ □ AB-	□ AB+ □ N/A □ B- □ AB+	Y
^IMPLICATED //://://:	UNIT ISBT-128 Codabar ISBT-128 Codabar Codabar	☐ Entire unit ☐ Partial unitmL ☐ Entire unit ☐ Partial unit				□ A- □B+ □ O- □ A- □B+	□ A+ □ AB- □ O+ □ A+ □ AB-	□ AB+ □ N/A □ B- □ AB+	Y
^IMPLICATED	UNIT ISBT-128 Codabar ISBT-128 Codabar Codabar	☐ Entire unit ☐ Partial unitmL ☐ Entire unit ☐ Partial unit				□ A- □B+ □ O- □ A- □B+	□ A+ □ AB- □ O+ □ A+ □ AB-	□ AB+ □ N/A □ B- □ AB+	Y
^IMPLICATED	UNIT ISBT-128 Codabar ISBT-128 Codabar Codabar	☐ Entire unit ☐ Partial unitmL ☐ Entire unit ☐ Partial unit				□ A- □B+ □ O- □ A- □B+	□ A+ □ AB- □ O+ □ A+ □ AB-	□ AB+ □ N/A □ B- □ AB+	Y

