

Hemovigilance Module Adverse Reaction Transfusion Associated Circulatory Overload

***Required for saving**

*Facility ID#: _____		NHSN Adverse Reaction #: _____	
Patient Information			
*Patient ID: _____		*Date of Birth: ____/____/____	
*Sex: ☐ M ☐ F			
Social Security #: _____		Secondary ID: _____ Medicare #: _____	
Last Name: _____		First Name: _____ Middle Name: _____	
Ethnicity (Specify): ☐ Hispanic or Latino ☐ Not Hispanic or Latino ☐ Unknown ☐ Declined to respond			
Race (Select all that apply): ☐ American Indian or Alaska Native ☐ Asian ☐ Black or African American ☐ Middle Eastern or North African ☐ Native Hawaiian or Pacific Islander ☐ White ☐ Unknown ☐ Declined to respond			
Preferred Language (Specify from the list provided): _____		Interpreter Needed: ☐ Yes ☐ No ☐ Unknown ☐ Declined to Respond	
*Blood Group: ☐ A- ☐ A+ ☐ B- ☐ B+ ☐ AB- ☐ AB+ ☐ O- ☐ O+ ☐ Blood type not done ☐ Transitional ABO / Rh + ☐ Transitional ABO / Rh - ☐ Transitional ABO / Transitional Rh ☐ Group A/Transitional Rh ☐ Group B/Transitional Rh ☐ Group O/Transitional Rh ☐ Group AB/Transitional Rh			
Patient Medical History			
List the patient's admitting diagnosis. (Use ICD-10 Diagnostic codes/descriptions)			
Code: _____		Description: _____	
Code: _____		Description: _____	
Code: _____		Description: _____	
List the patient's underlying indication for transfusion. (Use ICD-10 Diagnostic codes/descriptions)			
Code: _____		Description: _____	
Code: _____		Description: _____	
Code: _____		Description: _____	
List the patient's comorbid conditions at the time of the transfusion related to the adverse reaction. (Use ICD-10 Diagnostic codes/descriptions)			☐ UNKNOWN ☐ NONE
Code: _____		Description: _____	
Code: _____		Description: _____	
Code: _____		Description: _____	

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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) ☐ UNKNOWN ☐ NONE

Code: _____ Description: _____

Code: _____ Description: _____

Code: _____ Description: _____

Additional Information _____

Transfusion History

Has the patient received a previous transfusion? ☐ YES ☐ NO ☐ UNKNOWN

Blood Product: ☐ WB ☐ RBC ☐ Platelet ☐ Plasma ☐ Cryoprecipitate ☐ Granulocyte

Date of Transfusion: ____/____/____ ☐ UNKNOWN

Was the patient's adverse reaction transfusion-related? ☐ YES ☐ NO

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? ☐ Yes ☐ No If Yes, Incident #: _____

Investigation Results

*☐ **Transfusion associated circulatory overload (TACO)**

*Case Definition

Check all that occurred within 12 hours of cessation of transfusion (new onset or exacerbation):

☐ Acute respiratory distress (dyspnea, orthopnea, cough)

☐ Elevated brain natriuretic peptide (BNP)

☐ Elevated central venous pressure (CVP)

☐ Evidence of left heart failure

☐ Evidence of positive fluid balance

☐ Radiographic evidence of pulmonary edema

Other signs and symptoms: (check 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	☐ Infusion site pain
Renal:	☐ Hematuria	☐ Hemoglobinuria
	☐ Oliguria	
Respiratory:	☐ Bilateral infiltrates on chest x-ray	☐ Bronchospasm
	☐ Hypoxemia	☐ Shortness of breath
☐ Other: (specify) _____		

*Severity

Did the patient receive or experience any of the following?

- | | |
|---|---|
| ☐ No treatment required | ☐ Symptomatic treatment only |
| ☐ Hospitalization, including prolonged hospitalization | ☐ Life-threatening reaction |
| ☐ Disability and/or incapacitation | ☐ Congenital anomaly or birth defect(s) of the fetus |
| ☐ Other medically important conditions | ☐ Death |
| | ☐ Unknown or not stated |

*Imputability

Which best describes the relationship between the transfusion and the reaction?

- ☐ No other explanations for circulatory overload are possible.
- ☐ Transfusion is a likely contributor to circulatory overload
- ☐ The patient has a history of a pre-existing condition(s) that most likely explains circulatory overload.
- ☐ Evidence is clearly in favor of a cause other than the transfusion, but transfusion cannot be excluded.
- ☐ There is conclusive evidence beyond reasonable doubt of a cause other than the transfusion.
- ☐ The relationship between the adverse reaction and the transfusion is unknown or not stated.

Did the transfusion occur at your facility? ☐ YES ☐ NO

Does the patient have a history of cardiac insufficiency?

- ☐ Yes, the patient has a history of cardiac insufficiency that could explain the circulatory overload, but transfusion is just as likely to have caused the circulatory overload.
- ☐ Yes, the patient has a history of pre-existing cardiac insufficiency that most likely explains circulatory overload.
- ☐ No, the patient does not have a history of cardiac insufficiency.

Did the patient received other fluids in addition to the transfusion? ☐ YES ☐ NO

Module-generated Designations

NOTE: Designations for case definition, severity, and imputability will be automatically assigned in the NHSN application based on responses in the corresponding investigation results section above.

*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 _____

Patient Treatment

Did the patient receive treatment for the transfusion reaction? ☐ YES ☐ NO ☐ UNKNOWN

If yes, select treatment(s):

☐ Medication (*Select the type of medication*)

☐ Antipyretics ☐ Antihistamines ☐ Inotropes/Vasopressors ☐ Bronchodilator ☐ Diuretics

☐ Intravenous

Immunoglobulin

☐ Intravenous steroids

☐ Corticosteroids

☐ Antibiotics

☐ Antithymocyte globulin

☐ Cyclosporin

☐ Other

☐ Volume resuscitation (Intravenous colloids or crystalloids)

☐ Respiratory support (*Select the type of support*)

☐ Mechanical ventilation

☐ Noninvasive ventilation

☐ Oxygen

☐ 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

Date of Death: ____/____/____

^If recipient died, relationship of transfusion to death:

☐ Definite

☐ Probable

☐ Possible

☐ Doubtful

☐ Ruled Out

☐ Not determined

Cause of death: _____

Was an autopsy performed? ☐ Yes ☐ No

Component Details

***Was a particular unit implicated in (i.e., responsible for) the adverse reaction?**

☐ Yes

☐ No

☐ N/A

Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	^Unit number (Required for Infection and TRALI)	*Unit expiration Date/Time	*Blood group of unit	Implicated Unit?
____/____/____ ____:____:____	☐ ISBT-128 ☐ Codabar	☐ Entire unit ☐ Partial unit _____ mL	_____ _____ _____	____/____/____ ____:____:____	☐ A- ☐ A+ ☐ B- ☐ B+ ☐ AB- ☐ AB+ ☐ O- ☐ O+ ☐ N/A	Y
____/____/____ ____:____:____	☐ ISBT-128 ☐ Codabar	☐ Entire unit ☐ Partial unit _____ mL	_____ _____ _____	____/____/____ ____:____:____	☐ A- ☐ A+ ☐ B- ☐ B+ ☐ AB- ☐ AB+ ☐ O- ☐ O+ ☐ N/A	N

Custom Fields

Label	Label
_____ / _____ / _____ _____	_____ / _____ / _____ _____
Comments	