

*Required for saving

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Infection

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

*☐ Infection

*Case Definition

Was a test to detect a specific pathogen performed on the recipient post-transfusion?

☐ Yes

☐ No

If Yes, positive or reactive results?

☐ Yes

☐ No

Org1 _____

Org2 _____

Org3 _____

Was a test to detect a specific pathogen performed on the donor post-donation?

☐ Yes

☐ No

If Yes, positive or reactive results?

☐ Yes

☐ No

Org1 _____

Org2 _____

Org3 _____

Was a test to detect a specific pathogen performed on the unit post-transfusion? (i.e., culture, serology, NAT)

☐ Yes

☐ No

If Yes, positive or reactive results?

☐ Yes

☐ No

Org1 _____

Org2 _____

Org3 _____

Check all that apply:

☐ Temporally associated unexplained clinical illness consistent with infection

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 ☐ Cough ☐ 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 potential exposures to the pathogen could be identified in the recipient.
☐ Evidence is clearly in favor of a cause other than 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.

Check all that apply:

- ☐ Evidence of the pathogen in the transfused component.
☐ Evidence of the pathogen in the donor at the time of donation.
☐ Evidence of the pathogen in an additional component from the same donation.
☐ Evidence of the pathogen in an additional recipient of a component from the same donation.
☐ Evidence that the identified pathogen strains are related by molecular or extended phenotypic comparison testing with statistical confidence ($p < 0.05$).
☐ Evidence that the transfused component was negative for this pathogen at the time of transfusion
☐ Evidence that the donor was negative for this pathogen at the time of donation.
☐ Evidence that additional components from the same donation were negative for this pathogen.
☐ Evidence that the recipient was not infected with the pathogen prior to transfusion.
☐ Laboratory evidence that the recipient was infected with this pathogen prior to transfusion.

Did the transfusion occur at your facility? ☐ 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

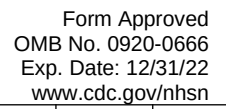
***Was a particular unit implicated in (i.e., responsible for) the adverse reaction?**

☐ 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

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