

**Instructions:** For all adverse events, complete sections 1, 2 and 3.

In addition, for:

- suspected transfusion transmitted infectious disease events (other than bacterial), complete section 4.
- suspected TRALI reactions, complete section 5.
- suspected bacterial contamination events, complete section 6.

*You may be required to report this adverse event to your state department of health. Follow your local procedures for state reporting.*

Please sign the last page and submit the completed form to the **facility that shipped implicated blood unit(s) to you.** Contact information for each facility is included below.

|  |   |
|--|---|
| <p><b><u>Community Blood Centers- Kansas City</u></b></p> <ul style="list-style-type: none"><li>• TRALI- Fax to IRL at 816-277-0757 or email to <a href="mailto:Immuno@cbckc.org">Immuno@cbckc.org</a></li></ul> <p><b>Contact IRL immediately if TRALI is involved in a patient fatality (816-968-4053)</b></p> <ul style="list-style-type: none"><li>• Bacterial Contamination – Fax to QM at 816-277-0798 or email to <a href="mailto:QAGroupALL@cbckc.org">QAGroupALL@cbckc.org</a></li><li>• Post Transfusion Disease- Fax to Donor Notification at 816-277-0785 or email to <a href="mailto:TherapeuticCollectionServices@cbckc.org">TherapeuticCollectionServices@cbckc.org</a></li></ul> | <p><b><u>New York Blood Center</u></b></p> <p>Special Donor Services Department</p> <ul style="list-style-type: none"><li>• Phone: 800-688-0900</li><li>• Fax: 212-288-8464</li></ul> |
| <p><b><u>Blood Bank of Delmarva</u></b></p> <p>Submit reports through Blood Hub. If not available, send report to:</p> <p>Reference Laboratory</p> <ul style="list-style-type: none"><li>• Fax: 302-709-6155</li><li>• Then call 302-737-8405 ext. 716</li></ul>   | <p><b><u>Rhode Island Blood Center</u></b></p> <p>Laboratory Supervisor</p> <ul style="list-style-type: none"><li>• Phone: 401-453-8374</li><li>• Fax: 401-248-5750</li></ul>         |
| <p><b><u>Innovative Blood Resources</u></b></p> <p>Memorial Blood Centers<br/>Nebraska Community Blood Bank</p> <p>Physician Services Donor Advocates</p> <ul style="list-style-type: none"><li>• Phone 651-332-7287, Fax 651-332-7001</li></ul>   |   |

|   |  |   |
|---|--|---|
| <b>1</b>  | <b>FACILITY INFORMATION AND DESCRIPTION OF EVENT</b> |   |
| <b>Reporting Facility Information</b>               |  |   |
| Date of Report:                                     | Name of person reporting:                            | Title of person reporting:                            |
| Telephone number:                                   | Email address:                                       |   |
| Reporting Facility Name:                            | Reporting Facility Address:                          |   |
| Transfusion Medicine Physician Name:                | Transfusion Medicine Physician Phone Number:         |   |
| <b>Select Suspected Category for Adverse Event:</b> |  |   |
| Check all that apply<br>▶                           | <input type="checkbox"/>                             | Anaplasma   |
|   | <input type="checkbox"/>                             | Babesiosis  |
|   | <input type="checkbox"/>                             | HBV   |
|   | <input type="checkbox"/>                             | HCV   |
|   | <input type="checkbox"/>                             | HIV 1-2   |
|   | <input type="checkbox"/>                             | HTLV I-II   |
|   | <input type="checkbox"/>                             | Septic Transfusion Reaction (Bacterial Contamination) |
|   | <input type="checkbox"/>                             | Transfusion Related Acute Lung Injury (TRALI)         |
|   | <input type="checkbox"/>                             | Other ▼ (if selected, describe below)                 |
| Additional Information<br>▶                         |  |   |

|   |  |  |
|---|--|--|
| <b>2</b>  | <b>PATIENT INFORMATION</b>   |  |
| <b>Patient Recipient General Information</b>        |  |  |
| Medical Record Number:                              | Patient Date of Birth:   | Patient Sex:<br><input type="checkbox"/> Female<br><input type="checkbox"/> Male |
| <b>Medical Information</b>                          |  |  |
| Attending Physician Name:                           |  | Attending Physician Phone Number:  |
| Admitting or Primary Diagnosis:                     |  | Indication for Transfusion:  |
| Relevant Severe Co-Morbidities (if applicable):     | Current Status of Patient:   |  |
|   | <input type="checkbox"/> Expired (Transfusion Related fatality) ** Report to FDA within 24 hours |  |
|   | <input type="checkbox"/> Reaction continues  |  |
|   | <input type="checkbox"/> Returned to pre-transfusion status                                      |  |
|   | <input type="checkbox"/> Unknown   |  |
| <input type="checkbox"/> Other ▼ describe if other: |  |  |
| <b>Treatment and Clinical Course</b>                |  |  |
| Treatment   | Check all Treatments Administered  | Indicate YES if patient Responded to administered treatment                      |
| Acetaminophen                                       | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Antihistamines                                      | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Bronchodilators                                     | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Diuretics   | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Epinephrine   | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Intubation Ventilatory Support                      | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Oxygen Supplementation                              | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Steroids  | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Other (specify) ►                                   | <input type="checkbox"/> YES   | <input type="checkbox"/> YES   |
| Describe if <b>Other</b> :                          |  |  |
| Additional Comments:                                |  |  |



|                        |  |                                       |
|------------------------|--|---------------------------------------|
| 2- Date of transfusion | 2-Start Time of Unit Transfusion (hh:mm) | 2-End Time of Unit Transfusion(hh:mm) |
| 3-DIN:                 | 3-Component Type:                        |                                       |
| 3- Date of transfusion | 3-Start Time of Unit Transfusion (hh:mm) | 3-End Time of Unit Transfusion(hh:mm) |
| 4-DIN:                 | 4-Component Type:                        |                                       |
| 4- Date of transfusion | 4-Start Time of Unit Transfusion (hh:mm) | 4-End Time of Unit Transfusion(hh:mm) |
| 5-DIN:                 | 5-Component Type:                        |                                       |
| 5- Date of transfusion | 5-Start Time of Unit Transfusion (hh:mm) | 5-End Time of Unit Transfusion(hh:mm) |
| 6-DIN:                 | 6-Component Type:                        |                                       |
| 6- Date of transfusion | 6-Start Time of Unit Transfusion (hh:mm) | 6-End Time of Unit Transfusion(hh:mm) |
| 7-DIN:                 | 7-Component Type:                        |                                       |
| 7 Date of transfusion  | 7-Start Time of Unit Transfusion (hh:mm) | 7-End Time of Unit Transfusion(hh:mm) |
| 8-DIN:                 | 8-Component Type:                        |                                       |
| 8- Date of transfusion | 8-Start Time of Unit Transfusion (hh:mm) | 8-End Time of Unit Transfusion(hh:mm) |
| 9-DIN:                 | 9-Component Type:                        |                                       |
| 9- Date of transfusion | 9-Start Time of Unit Transfusion (hh:mm) | 9-End Time of Unit Transfusion(hh:mm) |

|  |   |  |
|--|---|--|
| 10-DIN:                                  | 10-Component Type:                        |  |
| 10- Date of transfusion                  | 10-Start Time of Unit Transfusion (hh:mm) | 10-End Time of Unit Transfusion(hh:mm) |
| Specify any modifications made to units: |   |  |

|   |   |
|---|---|
| <b>4</b>  | <b>INFECTIOUS DISEASE AND TESTING</b>                                     |
| <b>Infectious Diseases</b>  |   |
| Has the patient been assessed for risks from exposure (e.g. IV drug use, tattoos, acupuncture-ear piercing-venereal disease-sexual contact with infected partner)?                        | <input type="checkbox"/> YES explain below<br><input type="checkbox"/> NO |
| Could the event be related to causes other than the transfusion (dialysis-receipt of clotting factors in the past-occupational exposure to blood or body fluids-needle stick-spill-bite)? | <input type="checkbox"/> YES explain below<br><input type="checkbox"/> NO |
| Explain (if YES):   |   |
| <b>Testing</b>  |   |
| Was the recipient tested for this infectious disease prior to transfusion?  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO               |
| List application Pre and Post Txn test results below:   |   |
|   |   |
| <b>Hepatitis Testing</b>  |   |
| PRE-TXN   | POST-TXN  |
| Pre-Txn test Date:  | Post-Txn test Date:   |
| Pre-Txn HBsAg Result:   | Post-Txn HBsAg Result:  |
| Pre-Txn Anti-HBs Result:  | Post-Txn Anti-HBs Result:   |
| Pre-Txn Anti-HBc Result:  | Post-Txn Anti-HBc Result:   |

|   |                                   |                           |
|---|-----------------------------------|---------------------------|
| Pre-Txn Anti-HCV Result:                      | Post-Txn Anti-HCV Result:         |                           |
| Pre-Txn HBV PCR Result:                       | Post-Txn HBV PCR Result:          |                           |
| Pre-Txn HCV PCR Result:                       | Post-Txn HCV PCR Result:          |                           |
| <b>HIV Testing</b>                            |                                   |                           |
| PRE-TXN                                       | POST-TXN                          |                           |
| HIV Pre-Txn Test Date                         | HIV Post-Txn Test Date            |                           |
| Pre-Txn Anti-HIV Result                       | Post-Txn Anti-HIV Result          |                           |
| Pre-Txn HIV PCR Result                        | Post-Txn HIV PCR Result           |                           |
| Other HIV Tests (Specify and provide result): |                                   |                           |
| <b>Babesiosis Testing</b>                     |                                   |                           |
| PRE-TXN                                       | POST-TXN                          |                           |
| Babesiosis Pre-Txn Testing Date:              | Babesiosis Post-Txn Testing Date: |                           |
| Pre-Txn Antibody Result:                      | Post-Txn Antibody Result:         |                           |
| Pre-Txn PCR Result:                           | Post-Txn PCR Result:              |                           |
| <b>Additional Testing</b>                     |                                   |                           |
| Other Testing:                                | Other Test Pre-Txn Date:          | Other Test Post-Txn Date: |
| Other Test Pre-Txn Result:                    | Other Test Post-Txn Result:       |                           |

|  |  |  |                        |
|--|--|--|------------------------|
| <b>5</b>   | <b>TRALI REACTION INFORMATION</b>  |  |                        |
| <b>Risk Factors for Acute Lung Injury</b> check all that apply ▼   |  |  |                        |
| <input type="checkbox"/> Acute Pancreatitis<br><input type="checkbox"/> Acute Respiratory Distress Syndrome(ARDS)<br><input type="checkbox"/> Amiodarone<br><input type="checkbox"/> Aspiration<br><input type="checkbox"/> Burn<br><input type="checkbox"/> Cardiopulmonary Bypass<br><input type="checkbox"/> Chemotherapy | <input type="checkbox"/> Diffuse Alveolar Damage<br><input type="checkbox"/> Disseminated Intravascular Coagulation<br><input type="checkbox"/> Drug Overdose<br><input type="checkbox"/> Lung Contusion<br><input type="checkbox"/> Massive Blood Transfusion<br><input type="checkbox"/> Multiple Trauma<br><input type="checkbox"/> Near Drowning | <input type="checkbox"/> Pneumonia<br><input type="checkbox"/> Severe Sepsis<br><input type="checkbox"/> Shock<br><input type="checkbox"/> Renal Failure<br><input type="checkbox"/> Radiation to Thorax<br><input type="checkbox"/> Upper Airway Obstruction<br><input type="checkbox"/> Toxic Inhalation |                        |
| Additional Comments (Other risk factors):  |  |  |                        |
|  |  |  |                        |
| <b>Pre-Transfusion Diagnostics</b>   |  |  |                        |
|  | Diagnostic Test  | Test performed?  | Pre-Transfusion Values |
| 1  | O2 sat ≤ 90% on room air   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed  | Pre-Txn Value:         |
| 2  | PaO2FIO2 ≤ 300mm Hg  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed  | Pre-Txn Value:         |
| 3  | Chest X-ray: Bilateral infiltrates   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed  |                        |
| 4  | Chest X-Ray: Widened Cardiac Silhouette (Cardiomegaly)   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed  |                        |
| 5  | Elevated BNP (Provide value in pg per mL)  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed  | Pre-Txn Value:         |
| 6  | Elevated Central Venous Pressure greater than 12mm Hg (Provide values.)  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed  | Pre-Txn Value:         |
| 7  | Elevated Pulmonary Artery Pressure greater than 18 mm Hg (Provide values.)   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed  | Pre-Txn Value:         |



|   |  |   |   |
|---|--|---|---|
| 8   | Positive Fluid Value (in mL)   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| 9   | Transient decrease White Blood Cell Count                                  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| <b>Post-Transfusion Diagnostics</b>                         |  |   |   |
| Diagnostic Test   |  | Test performed?   | Pre-Transfusion Values                                |
| 1   | O2 sat ≤ 90% on room air   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| 2   | PaO2FIO2 ≤ 300mm Hg  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| 3   | Chest X-ray: Bilateral infiltrates   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed |   |
| 4   | Chest X-Ray: Widened Cardiac Silhouette (Cardiomegaly)                     | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed |   |
| 5   | Elevated BNP (Provide value in pg per mL)                                  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| 6   | Elevated Central Venous Pressure greater than 12mm Hg (Provide values.)    | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| 7   | Elevated Pulmonary Artery Pressure greater than 18 mm Hg (Provide values.) | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| 8   | Positive Fluid Value (in mL)   | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| 9   | Transient decrease White Blood Cell Count                                  | <input type="checkbox"/> YES<br><input type="checkbox"/> NO<br><input type="checkbox"/> Not Performed | Pre-Txn Value:  |
| <b>If TRALI is diagnosed, please provide the following:</b> |  |   |   |
| Recipient HLA Type:   |  | Recipient HNA Type:   | Recipient HLA-HNA antibody status and identification: |

|  |                               |
|--|-------------------------------|
| Donor HLA-HNA antibody status and identification (if performed on unit): | Donor HLA type (if available) |
|--|-------------------------------|

|          |                                |
|----------|--------------------------------|
| <b>6</b> | <b>BACTERIAL CONTAMINATION</b> |
|----------|--------------------------------|

**Suspected Bacterial Contamination Questions**

|   |  |                                  |
|---|--|----------------------------------|
| Were the suspected units returned to the blood bank?<br><br><input type="checkbox"/> YES<br><input type="checkbox"/> NO | On reinspection does the component present any abnormalities (e.g. clumps, discoloration, hemolysis)?<br><br><input type="checkbox"/> YES<br><input type="checkbox"/> NO | Describe abnormalities (if any): |
|---|--|----------------------------------|

|   |  |
|---|--|
| Suspect Component- Source Used:<br><br><input type="checkbox"/> Bag<br><input type="checkbox"/> Segment<br><input type="checkbox"/> Not performed | Does the patient have history of fever or of other infection-related to his / her underlying medical condition?<br><br><input type="checkbox"/> YES<br><input type="checkbox"/> NO |
|---|--|

|   |                                      |
|---|--------------------------------------|
| Was the patient on antibiotics at the time of transfusion?<br><br><input type="checkbox"/> YES ►<br><input type="checkbox"/> NO | Specify antibiotic (if <b>YES</b> ): |
|---|--------------------------------------|

|   |                                      |
|---|--------------------------------------|
| Is the patient currently being treated with antibiotics?<br><br><input type="checkbox"/> YES ►<br><input type="checkbox"/> NO | Specify antibiotic (if <b>YES</b> ): |
|---|--------------------------------------|

|   |
|---|
| Did the patient have an absolute neutropenia count (neutrophil less than 500 per µl) prior to transfusion?<br><br><input type="checkbox"/> YES<br><input type="checkbox"/> NO |
|---|

|                      |
|----------------------|
| Additional Comments: |
|----------------------|

**Suspected Bacterial Contamination Additional Testing**

|   |                    |
|---|--------------------|
| Gram Stain Results for unit:<br><br><input type="checkbox"/> Negative<br><input type="checkbox"/> Positive<br><input type="checkbox"/> Not Done | Result (Organism): |
|---|--------------------|

|   |                    |
|---|--------------------|
| Culture Performed on unit:<br><br><input type="checkbox"/> Negative<br><input type="checkbox"/> Positive<br><input type="checkbox"/> Pending<br><input type="checkbox"/> Not Done | Result (Organism): |
|---|--------------------|

|   |                                  |   |  |
|---|----------------------------------|---|--|
| Was a secondary test performed by the hospital for this component (PGD or equivalent)?<br><input type="checkbox"/> YES ►<br><input type="checkbox"/> NO                                     |                                  | Specify test performed if <b>YES</b> :        |  |
| Patient Pre-Transfusion Blood Culture<br><input type="checkbox"/> Negative<br><input type="checkbox"/> Positive<br><input type="checkbox"/> Pending<br><input type="checkbox"/> Not Done    | Date of Pre-Transfusion Culture: | Result of Pre-Transfusion Culture (Organism): |  |
| Patients Post-Transfusion Blood Culture:<br><input type="checkbox"/> Negative<br><input type="checkbox"/> Positive<br><input type="checkbox"/> Pending<br><input type="checkbox"/> Not Done | Date of Post-Transfusion Culture | Result of Post-Transfusion Culture (Organism) |  |

|                                      |            |       |
|--------------------------------------|------------|-------|
| <b>Signature of person reporting</b> | Signature: | Date: |
|--------------------------------------|------------|-------|

**Submit the completed form to the facility that shipped implicated blood unit(s).**

## Signature Manifest

**Document Number:** EW-FRM-0003

**Revision:** 02

**Title:** Suspected Transfusion Related Adverse Event Form

**Effective Date:** 03 May 2021

All dates and times are in Eastern Time.

### EW-FRM- 0003 rev 02 Suspected Transfusion Related Adverse Event Form

#### NEEC Approval

| Name/Signature        | Title | Date                     | Meaning/Reason |
|-----------------------|-------|--------------------------|----------------|
| Betsy Jett (NY-BJETT) |       | 23 Apr 2021, 03:42:05 PM | Approved       |

#### Set Effective Date

| Name/Signature                 | Title | Date                     | Meaning/Reason |
|--------------------------------|-------|--------------------------|----------------|
| MaryBeth Parache (NY-MPARACHE) |       | 26 Apr 2021, 04:47:04 PM | Approved       |

#### Quick Approval

#### Approve Now

| Name/Signature               | Title | Date                     | Meaning/Reason |
|------------------------------|-------|--------------------------|----------------|
| Ivette Augusto (NY-IAUGUSTO) |       | 03 May 2021, 05:25:42 AM | Approved       |