#### TRAINING UPDATE

Lab Location: Department: SGAH and WAH Blood Bank Date Implemented: Due Date: 6.3.2014 6.8.2014

#### **DESCRIPTION OF PROCEDURE REVISION**

## Name of procedure:

**Blood Component Irradiation** 

## **Description of change(s):**

- 1. Nothing has changed for content with regards to irradiating regular blood products for inventory.
- 2. The component prep function for irradiating DIRECTED red cells will be "ID####" instead of "IE####."
- 3. Blood label check will be performed in Sunquest following irradiation.

Title	Blood Component Irradiation	
Prepared by	Stephanie Codina	Date: 2/15/2010
Owner	Stephanie Codina	Date: 2/15/2010

Print Name and Title	Signature	Date
Refer to the electronic signature page for approval and approval dates.		
Local Issue Date:	Local Effective Date:	

Review:		
Print Name	Signature	Date
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#### 1. PURPOSE

Blood components that contain viable lymphocytes (red blood cells, platelets, and granulocytes) may be irradiated to prevent proliferation of T-lymphocytes. Proliferation of T-lymphocytes causes transfusion-associated graft-versus-host disease (TA-GVHD), a disease in which donor lymphocytes engraft in the recipients and mount an attack on host tissue. Irradiated blood is prepared by exposing the component to a radiation source. The standard dose of gamma irradiation is 25 Gy targeted to the central portion of the blood product container with a minimum dose of 15 Gy delivered to any part of the component.

#### 2. SCOPE

Red blood cell products, platelet products, and granulocyte products must be irradiated in the following situations:

- When requested by the treating physician for a hospital-approved reason or with pathologist approval
- When a directed-donor unit from any blood relative will be transfused
- When the recipient is receiving HLA-matched and/or crossmatched platelets
- When red cells and platelets are provided for neonatal transfusion, including exchanges and intrauterine transfusion.

SGAH blood bank will irradiate blood products for the WAH blood bank.

#### 3. RESPONSIBILITY

All blood bank staff must demonstrate competency for component irradiation.

form revised 3/31/0

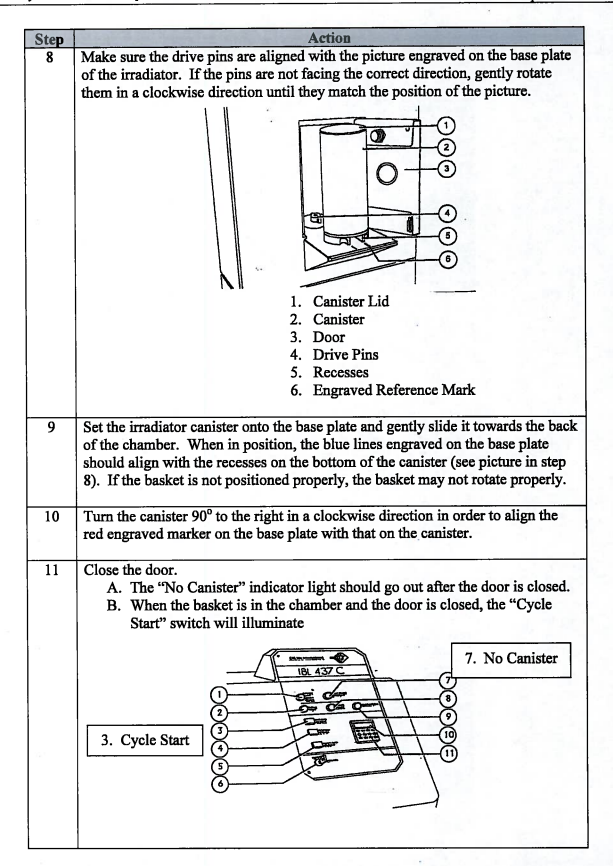
#### **DEFINITIONS** 4.

- 1. Irradiator = a CIS Bio International IBL 437C
- 2. Gy = Gray; Gray  $\times$  100 = rad. The real dose of energy per gram of tissue (blood product) delivered by irradiation.

#### **PROCEDURE** 5.

Step	Action
1	Determine if the recipient needs blood product irradiation.  A. When the physician requests irradiated blood products for the recipient for the first time,  a. Verify the reason for irradiation is an appropriate indication per hospital-defined criteria outlined in appendix B. Pathologist approval is required for irradiation outside of hospital-defined criteria.  b. Enter the irradiation marker into patient's blood bank administrative data file per procedure. Once the irradiation marker has been added, all subsequent transfusions must be irradiated until a physician sends a written order to blood bank indicating transfusion of irradiated blood products is no longer clinically necessary.  B. All directed-donor blood products from all blood relatives will be irradiated.  C. All HLA-matched and/or crossmatched platelets will be irradiated.  D. All granulocyte products will be irradiated.  E. All red blood cell and platelet products provided to neonates <4 months in age will be irradiated. This includes neonatal, exchange, and intrauterine transfusions.  Note: Irradiation applies to cellular products only (red blood cell, platelet, and granulocyte products). Non-cellular products such as plasma and cryoprecipitate are not routinely irradiated.
2	Select a blood product that meets all patient criteria including blood type, antigen status, and any other special attributes.
3	Complete the "Irradiation Log Sheet" by filling in the following information.  A. Tech performing irradiation B. Date of irradiation C. Unit number or donor identification number (DIN) D. Product code prior to irradiation E. Product code following irradiation F. Division G. Expiration date of original product H. Expiration date of irradiated product (28 days from date of expiration or original expiration date; whichever is first)

Step	Action
4	Place a RAD-SURE label on each component to be irradiated (Refer to appendix A).
19	A. Verify the lot number and expiration date of the Rad-Sure indicators.  Document the verification on the Irradiation Log Sheet.  B. Verify that the word "NOT" is showing on each Rad-Sure indicator to be
	used. C. Place the label on the front of the blood product below the product label. D. DO NOT cover any of the base label during application.
5	<ul> <li>Place each component that needs to be irradiated into the basket.</li> <li>A. Do not commingle products that are not stored at the same temperature.</li> <li>Do not place red blood cells and platelets into the carrier at the same time.</li> <li>B. Do not irradiate more than two blood products at one time.</li> <li>C. Do not overfill the irradiation canister. There should be at least 1 inch of space between the blood products and the canister lid.</li> <li>D. The canister will accommodate a 60 mL syringe if you remove the yellow doughnut from the canister.</li> </ul>
6	Replace the canister lid turning it counter-clockwise to lock it into place.
7	Open the door of the irradiation chamber.  Door



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64	Action
Step 12	Action  The IBL 437-C should always have the power turned (contact key) to the "On"
12	position. Ensure the following indicator lights are illuminated:
	• Timer display screen (#10)
	• Mains (#2)
	• Battery (#8)
	Notify a supervisor if the lights are not illuminated.
	15. 437 C
9	1. Continue Detation Links 7. No Continue
	1. Canister Rotation Light 7. No Canister 2. Mains (On/Off Light) 8. Battery
	2. Mains (On/Off Light) 8. Battery 3. Cycle Start 9. Irradiation
	4. Test Lamps 10. Timer Display Screen
	5. Cycle Break 11. Timer Keyboard
	6. Key Switch (On/Off)
13	Press the "Test Lamps" button. All indicators should light up except the "Test Lamps" indicator. Notify a supervisor if the lights do not illuminate.
14	The programmed irradiation time is calculated yearly at the time of preventative maintenance by the contracted blood irradiator service company.  A. Check timer display screen to make sure the correct time displays.  B. Document the initial irradiator setting on the Irradiation Log Sheet.  C. DO NOT TOUCH ANY OF THE BUTTONS ON THE TIMER DISPLAY SCREEN.
15	Press the "Cycle Start" switch to start the irradiation cycle. At this time, the drum will begin its rotation (this can be seen by looking in the window of the door), the "Canister Rotation" light will blink, the "Irradiation" light will illuminate, and the time on the "Timer Display Screen" will count down.  A. Observe that each of these occurs.  B. Document the canister rotation on the Irradiation Log Sheet.

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Step	Action  The state of the second state of the s
16	The irradiation cycle finishes when the programmed time has elapsed. The counter displays the value zero and the alarm will sound.  A. Timer accuracy must be verified each day of use.  B. Canister rotation must be documented for each cycle.  C. Refer to the irradiation quality control procedures.
'	C. Rolof to the fraction quarty contact procedures.
17	Open the door of the irradiation chamber to cease the alarm.  A. The timer will reset to the previously programmed value.  B. Turn the canister in a clockwise director in order to align the recesses at the bottom of the basket with the blue marks engraved in the base plate.  C. Slide the basket towards the base plate until it releases.  D. Remove the canister lid and blood products from the canister.  E. Store the empty canister inside the chamber. Do not leave the canister on the platform of the loading chamber.
eg	(a) (b) (c) (c) (c) (c) (c) (c) (c) (c) (c) (c
	1. Canister Lid 4. Drive Pins
	2. Canister 5. Recesses
	3. Door 6. Engraved Reference Mark
18	<ul> <li>Examine the "Rad-Sure" label for a black color.</li> <li>A. If the word "NOT" is visible, the unit did not receive the proper amount of irradiation and cannot be issued when the patient/situation requires irradiated blood products.</li> <li>B. Units that did not receive the proper irradiation dose cannot be reirradiated.</li> </ul>
10	
19	Complete the "Irradiation Log Sheet" by filling in the following information.  A. Ending timer setting  B. Indicate whether the timer check was performed (see Quality Control)  C. Indicate whether the word "NOT" is obscured on the rad-sure indicator label
	D. Time in and out of the storage container (refrigerator or platelet rotator)

#### **B.** LIS Documentation

Step	Action
1	Access Sunquest function "Blood Component Preparation."
2	<ul> <li>At the "Value" prompt, type the irradiation function that corresponds to the unit you are irradiating then press the "tab" key.</li> <li>A. For homologous units, the BCP code is I plus the E code of the original product. For example, if you are irradiating a E0226 product, type "IE0226."</li> <li>B. For directed products, the BCP code is ID plus the numerical portion of the E code. For example, if you are irradiating a directed "E0167" Sunquest will convert the product code to "ED0167." The irradiation code will be "ID0167."</li> <li>C. Refer to appendix D for additional information.</li> </ul>
3	Press the "Tab" key to default to the current date and time as the irradiation time. Enter the date and time on which the product was irradiated if irradiation took place at an earlier time (such as during computer downtime).
4	Click the "continue" button.
5	<ul> <li>A second "Blood Component Prep" screen will appear.</li> <li>A. At the "Unit #" prompt, scan the unit number DIN of the unit to be irradiated.</li> <li>B. At the "Component" prompt, scan the product code from the product to be irradiated. This will autofill both the product code and division fields.</li> </ul>

Step	Action
6	For red cell units only, enter the expiration date and time of the new, irradiated unit.  A. The new expiration date will be 28 days from the date of expiration or the original expiration date, whichever is sooner.  a. Sunquest will calculate the 28-day expiration if you type "T+28" in the expiration date field.  b. Compare the original expiration to the 28-day expiration and choose the date that is sooner.  B. The expiration time will be 2359 or the expiration time of the original unit, if sooner.
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7	Click the "save" button.
8	A "Preview Output / New Units" screen will appear. Review the information to ensure accuracy, then click on the "finish" button to generate new product/expiration date labels for the irradiated products.
9	Perform a blood label check of each irradiated unit in Sunquest per procedure.
10	Allocate and/or crossmatch each unit per procedure.
11	Return the unit to the appropriate storage container. Document the time the unit was returned to storage on the Irradiation Log Sheet.

### C. Troubleshooting

Step	Action			
1	In situations where the cycle must be interrupted,			
	A. Press the "Cycle Break" button.			
	B. The corresponding red indicator light will illuminate.			
	C. The drum will return to the loading/unloading position (ie facing the			
	door).			
	D. The timer counter will display the remaining time.			
	E. Open the chamber door and remove the canister.			
	When ready to resume the cycle,  A. Replace the canister.			
	B. Press the "Cycle Break" button a second time.			
	C. The red indicator light will go out.			
	D. The "Start Cycle" green light will illuminate.			
	E. Press the "Start Cycle" switch and the timer will resume the countdown			
	starting from the value reached at the time of the interruption.			
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	2. Mains (On/Off Light)  7. No Cantister  8. Battery			
	3. Cycle Start 9. Irradiation			
	4. Test Lamps 10. Timer Display Screen			
	5. Cycle Break 11. Timer Keyboard			
	6. Key Switch (On/Off)			
e e				

Step	Action					
2 2	When extreme emergency situations such as smoke, fire, or insulation failure occur,  A. Press the "Emergency Stop Punch."  B. The equipment will be electronically disconnected, including the battery.  C. The drum will remain in the position that it was in at the time of the disconnect.  D. The counter will stop the countdown and will restore the time programmed at the beginning of the cycle.  E. Notify a supervisor immediately.  NOTE: This action can cause over-exposure of the blood products. The blood product cannot be issued if the "Emergency Stop Punch" was pressed during its irradiation cycle.					

### D. Power Failure

Step	Action						
1	In the event of a power failure, the battery backup power supply ensures the						
	return of the drum to the loading/unloading position.						
	A. The "Mains" indicator light will flash.						
	B. The "Cycle Break" indicator light will illuminate, indicating						
	interruption of the cycle.						
	C. The timer will store the remaining irradiation time at the time of the						
	power failure.						
	D. If the power is interrupted for less than 3 seconds, the irradiation cycle						
	will not be interrupted.						
2	All blood products must be immediately removed from the canister and						
	returned to proper storage conditions in a power failure. The blood products						
	should be returned to the canister and cycle continued with the remaining time						
	when power is restored. To resume the cycle,						
	A. Press the "Cycle Break" switch to acknowledge the interruption (the						
	light will be blinking).						
	B. The indicator light will go out.						
	C. The timer countdown will resume from the value saved at the time of						
	the power failure until the end of the cycle.						

#### E. Basket Rotation Fault

Step	ep Action				
1	The "Canister Rotation" light will stop flashing in the event of a fault or stoppage in the canister's rotation.				
2	Press the "Cycle Break" button.				
3	Look through the eye-piece of the door to see if the drum returned to the loading/unloading position. The drum should automatically return to the loading/unloading position.				
4	If the drum does NOT return to the loading/unloading position:  A. Press the "Emergency Stop Button" to disconnect the power supply.  This is very important in preventing electrocution.				
	Emergency Stop Button  Front Panel				
	B. Remove the front panel of the machine and remove the safety control lever which is found in its holder.				
	<ul> <li>C. Insert the lever into the holes of the cam-holder disc and turn the drum in a clockwise direction to the loading/unloading position.</li> <li>D. Look through the eye-piece of the door to visualize.</li> <li>E. Remove the blood products from the canister and return them to their</li> </ul>				
	proper storage conditions.				

F. Consult a supervisor for further instructions.

#### F. Emergency Procedures

Step	Action				
The irradiator must be taken out of service <b>IMMEDIATELY</b> if the malfunctions during loading/unloading or during the irradiation cycle Evidence of malfunction shall include binding of moving parts or the of metal shavings or chips.					
	Press the emergency stop button if necessary to stop the irradiator.				
2	Secure the room housing the irradiator.				
3	Notify the Radiation Safety Officer and the contracted blood irradiator service company.				
4	Log and describe the abnormal occurrence(s) in the Blood Bank Communication Log.				
5	DO NOT attempt to repair or modify the irradiator.				
6	DO NOT attempt to operate the irradiator until clearance is obtained from either the Radiation Safety Officer or the contracted blood irradiator service company.				
7	Refer all questions regarding the safety of the irradiator to the Radiation Safety Officer.				

#### 6. RELATED DOCUMENTS

SOP: Entering Special Transfusion Attributes into the LIS

Form: Irradiation Log Sheet (AG.F109)

SOP: Quality Control of the Blood Component Irradiator

SOP: Blood Label Check

SOP: Crossmatch

SOP: Platelets for Transfusion

#### 7. REFERENCES

- 1. IBL 437C Irradiator Type H Operator's Manual. CIS Bio International.
- 2. Standards for Blood Banks and Transfusion Services, 29<sup>th</sup> ed., 2014. AABB Publishing, Bethesda, Maryland.
- 3. CIS-US, Inc Technical Bulletin TB-001, Quarterly Safety Checks, 9/28/06.
- 4. Guidance for Industry, Gamma Irradiation of Blood and Blood Components: A Pilot Program for Licensing, US Department of Health and Human Services, FDA-CBER, Feb 2000.

 Circular of information for the use of human blood and blood components. Prepared by AABB, the American Red Cross, America's Blood Centers, and the Armed Services Blood Program. Bethesda, MD: AABB, 2009.

#### 8. REVISION HISTORY

Version	Date	Reason for Revision	Revised By	Approved By
		Supersedes SOP SHB.012.001		
000	2/15/10	Updated owner Section 5: changed to table format and diagrams added, revised item B for LIS upgrade Section 9: added appendix A, renumbered subsequent appendices	S.Codina	N. Cacciabeve
001	4.25.11	Section 5: LIS documentation—Update BCP values; Section 6: Updated Irradiation Log Sheet; Section 9: forms moved to Related Documents; Deleted Component Modification Form	SCodina	NCacciabeve
002	10.9.12	Section 5: Updated criteria for irradiation, removed QC and calibration (moved to separate procedure) Section 6: Removed Rad-Sure SOP and QC form Section 9: Added Appendix A (Rad-Sure Indicators), removed Irradiation Time Calculation and Programming, renumbered subsequent appendices	SCodina	NCacciabeve
003	5.1.13	Section 5: Updated step 2 of LIS documentation (moved information to appendix D). Section 9: Added appendix D.	SCodina	NCacciabeve
004			SCodina	NCacciabeve
005	5.27.14	Section 5: Updated wording for clarity. Removed references to codabar labeling. Updated LIS information to reflect the v6.4 upgrade. Removed references to Pharmalucence (no longer exists). Section 9: Revised appendix D. Footer: version # leading zero's dropped due to new EDCS in use as of 10/7/13.	SCodina	NCacciabeve

#### 9. ADDENDA AND APPENDICES

- A. Rad-Sure Indicator Labels
- B. Indications for Blood Product Irradiation
- C. COMAR Regulations
- D. Irradiation Functions

# Appendix A Rad-Sure Indicator Labels

#### **PURPOSE**

When attached to blood products, Rad-Sure Type 15 indicators are a qualitative mechanism to indicate that a product has been exposed to radiation. They do not measure the quantitative dose from an irradiator or whether the IBL-437C is operating properly.

#### **PROCEDURE**

Step	Action
1	Each box of Rad-Sure indicators contains a temperature history indicator card. Verify the color of the temperature film.  a. A blue black color indicates the Rad-Sure indicators have been maintained at an appropriate temperature range and the indicators are safe to use.  b. A red/orange color indicates the Rad-Sure indicators have been exposed to excessive heat during transit. DO NOT use these indicators. Contact the manufacturer as indicated on the card. The manufacturer will send a replacement box of Rad-Sure indicators.
2	Remove a sheet of Rad-Sure indicators from the box and look for the word "NOT" in the window of each indicator. Do not use the indicators unless the word "NOT" is clearly visible.
	15 QY INDICATOR IR RADIATED Let No. \$174A1915 Le
3	Write the date of irradiation and your tech initials or tech code on the designated spaces on the indicator.
4	Peel the indicator from the backing and apply firmly to the appropriate clean, dry place on the blood product.
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Step	Action
6	Following irradiation, check the indicator to verify that the window is black and obscures the word "NOT." The indicator should now read "IRRADIATED." Immediately notify a supervisor if the word "NOT" is still visible.  [INTERIOR OF PRINTING   10   10   10   10   10   10   10   1
7	Rad-Sure indicators are stored refrigerated at temperatures between 0-6°C.  a. Prolonged exposure to heat and/or light can damage the indicators.  b. The indicators must be stored away from all radiation sources, including gamma rays, electron beam devices, and microwaves.
8	Rad-Sure indicators cannot be used to calibrate an irradiator or measure radiation dosage. They are a qualitative indicator, not quantitative.

#### REFERENCE:

Rad-Sure Type 15 Gy and Type 25 Gy Blood Irradiator Indicator Package Insert, International Specialty Products, Rev 01-05.

# Appendix B Indications for Blood Product Irradiation

	For All Patients		For Pediatric Patients		For Patients with Congenital Immune Deficiency
•	Malignant lymphoma, from diagnosis to death Currently or previously on purie analogue treatment (fludarabine, cladribine, 2-CDA, pentastatin) Products from first- or second-degree relatives HLA-matched components Granulocyte components Chronic graft-vs-host disease (GVHD) on purine analogue drugs Allogeneic marrow transplant from start of conditioning to end of GVHD prophylaxis 7 days before harvesting of autologous stem cell transplant to 3 months after transplant or 6 months if total body irradiation is used Aplastic anemia Undiagnosed pancytopenia ALL or AML for stem cell transplant		All red cell and platelet transfusions given to infants <4 months of age Intrauterine transfusions (IUT) of red cells or platelets "Top-up" transfusions if previous IUT Exchange transfusions (ET) or platelet transfusions following IUT Any ET if delay for irradiation does not compromise care Small blue cell tumors in childhood Acute lymphoblastic leukemia (ALL) Acute myeloblastic leukemia (AML) Burkitt's lymphoma/leukemia Solid tumors, eg  Ewing's sarcoma  Hepatoblastoma  Neuroblastoma  Retinoblastoma  Retinoblastoma  Rhabdomyosarcoma  Langerhan's cell histiocytosis	• • • • • •	Deficiency  Di George's syndrome Congenital heart deficiency or open-heart surgery in patient <6 months old Congenital cell-mediated immune deficiency Severe combined immune deficiency Wiskott-Aldrich syndrome Purine nucleoside phosphorylase deficiency Reticular dysgenesis Adenosine deaminase deficiency Major histocompatibility complex (HLA) I or II deficiency Leukocyte adhesion molecular deficiency Cell-mediated deficiency, not otherwise specified
		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	Open-heart surgery in patient <6 months old		

Berte LM. Transfusion Service Manual of Standard Operating Procedures, Training Guides, and Competence Assessment Tools, 2<sup>nd</sup> ed. 2007. AABB Press: Bethesda, Maryland.

# Appendix C COMAR Regulations

Due to the presence of an IBL 437C Irradiator in the Blood Bank, the following guidelines are to be followed as additional controls in accordance with Maryland Code Ann., Env. Art. 8-501, COMAR 26.12.01.01 C.30(b), and COMAR 26.12.01.01C.50.

- IC 1. The Irradiator is secured to prevent unauthorized access into the Blood Bank. The irradiator remains in a locked cage at all times to prevent unauthorized access. A second level of protection is offered by limiting access to the laboratory via locked doors.
  - a. Only the Blood Bank staff members who have been deemed trustworthy and reliable are permitted unescorted access to use the irradiator. A Blood Bank staff member is present in the room at all times.
  - b. All Blood Bank staff members are verified for employment by Quest Diagnostics Human Resources staff with background checks of previous employment, education, and references.
  - c. Routine maintenance is only conducted by an authorized NRC service provider. A member of the Blood Bank staff is present during routine maintenance.
  - d. A current list of Blood Bank staff members is posted.
  - e. The irradiator is contained within a locked cage within the Blood Bank. Only staff that has had a completed background check by the Nuclear Regulatory Commission is permitted to have unescorted access to the irradiator.
- IC 2. Any unauthorized access to the radioactive material is documented and any action taken is according to the Shady Grove Adventist Hospital Radiation Contamination Accident Plan (Policy 38016).
- IC 3. Any shipment of the licensed material is provided by the IBL 437C vendor CIS-US. The Blood Bank does not ship licensed radioactive material. All shipments are handled by the contracted irradiator service company and will be required to meet all NRC guidelines. The Blood Bank will establish the time of delivery, confirm receipt of radioactive material, and notify vendor of any delay in receipt.
- IC 4. There are no portable or mobile devices containing radioactive material in the Blood Bank.
- IC 5. All documentation pertaining to these guidelines will be retained for 3 years after they are no longer effective.
- IC 6. All documentation pertaining to these guidelines is to be maintained by the Blood Bank Supervisor and stored in the laboratory.

### Appendix D Irradiation Functions

**Irradiated Red Cells** 

Original Product	Component Prep Function	Final (Irradiated) Product
E0167	IE0167	E0178
E0181	IE0181	E0179
E0212	IE0212	E0223
E0226	IE0226	E0224
E0276	IE0276	E0274
E0316	IE0316	E0331
E0336	IE0336	E0332
E0366	IE0366	E0378
E0382	IE0382	E0379
E0404	IE0404	E0419
E0424	IE0424	E0420
E0605	IE0605	E4537
E0678	IE0678	E0661
E0685	IE0685	E0668
E0686	IE0686	E0669
E0693	IE0693	E0676
E0694	IE0694	E0677
E4519	IE4519	E4521
E4520	IE4520	E4522
E4531	IE4531	E4526
E4532	IE4532	E4527
E4533	IE4533	E4528
E4536	IE4536	E0660
E4543	IE4543	E4538
E4544	IE4544	E4539
E4545	IE4545	E4540
E4546	IE4546	E4539
E4547	IE4547	E0677

#### **Irradiated Platelets**

Original Product	Component Prep Function	Final (Irradiated) Product
E3077	IE3077	E3046
E3087	IE3087	E3056
E3088	IE3088	E3057
E3089	IE3089	E3058
E4643	IE4643	E4647
E4644	IE4644	E4648

Form revised 3/31/00

#### **Irradiated Directed Donor Red Cells**

Original Product	Component Prep Function	Final (Irradiated) Product
ED0167	ID0167	ED0178
ED0181	ID0181	ED0179
ED0212	ID0212	ED0223
ED0226	IE0226	ED0224
ED0276	ID0276	ED0274
ED0316	ID0316	ED0331
ED0336	ID0336	ED0332
ED0366	ID0366	ED0378
ED0382	ID0382	ED0379
ED0404	ID0404	ED0419
ED0424	ID0424	ED0420
ED0605	ID0605	ED4537
ED0678	ID0678	ED0661
ED0685	ID0685	ED0668
ED0686	ID0686	ED0669
ED0693	ID0693	ED0676
ED0694	ID0694	ED0677
ED4519	ID4519	ED4521
ED4520	ID4520	ED4522
ED4531	ID4531	ED4526
ED4532	ID4532	ED4527
ED4533	ID4533	ED4528
ED4536	ID4536	ED0660
ED4543	ID4543	ED4538
ED4544	ID4544	ED4539
ED4545	ID4545	ED4540
ED4546	ID4546	ED4539
ED4547	ID4547	ED0677