

Hemovigilance Module Monthly Incident Summary

*Required for saving

*Facility ID#: ____

*Month:

*Year: _____

All reporting is facility-wide. Include numbers of individual incident reports in the totals.

*Process Code	*Incident Code	*Total Incidents	*Total Adverse Reactions
	PC 00 Detail not specified	incidents	Reactions
shipment and receipt of products into the	PC 00 Detail not specified PC 01 Data entry incomplete/incorrect/not performed		
	PC 02 Shipment incomplete/incorrect		
	PC 03 Products and paperwork do not match		
	PC 04 Shipped/transported under inappropriate conditions		
	PC 05 Inappropriate return to inventory		
transfusion service from the	PC 06 Product confirmation incorrect/not performed		
supplier, another hospital site, satellite storage, or	PC 07 Administrative check not incorrect/not performed (record review/audit)		
clinical area.	PC 08 Product label incorrect/missing		
	US 00 Detail not specified		
US: Product Storage	US 01 Incorrect storage conditions		
(Transfusion Service) Events that occur during	US 03 Inappropriate monitoring of storage device		
product storage by the	US 04 Unit stored on incorrect shelf (e.g., ABO/autologous/directed)		
transfusion service.	US 05 Incorrect storage location		
	IM 00 Detail not specified		
IM: Inventory	IM 01 Inventory audit incorrect/not performed		
Management	IM 02 Product status incorrectly/not updated online (e.g., available/discarded)		
(Transfusion Service)	IM 03 Supplier recall/traceback not appropriately addressed/not performed		
Events that involve quality	IM 04 Product order incorrectly/not submitted to supplier		
management of the blood	IM 05 Outdated product in available inventory		
product inventory.	IM 06 Recalled/quarantined product in available inventory		
	PR 00 Detail not specified		
	PR 01 Order for wrong patient		
PR: Product/Test	PR 02 Order incompletely/incorrectly ordered (online order entry)		
Request	PR 03 Special processing needs not indicated (e.g., CMV negative, autologous)		
(Clinical Service) Events that occur when the	PR 04 Order not done		
clinical service orders	PR 05 Inappropriate/unnecessary (intended) test ordered		
patient tests or blood	PR 06 Inappropriate/unnecessary (intended) blood product ordered		
products for transfusion.	PR 07 Incorrect (unintended) test ordered		
	PR 08 Incorrect (unintended) blood product ordered		
	OE 00 Detail not specified		
OE: Product/Test	OE 01 Order entered for wrong patient		
Order Entry	OE 02 Order incompletely/incorrectly entered online		
(Transfusion Service) Events that occur when the transfusion service receives a patient order. This process may be excluded if clinical service uses online ordering.	OE 03 Special processing needs not entered (e.g., CMV-, autologous)		
	OE 04 Order entry not done		
	OE 05 Inappropriate/unnecessary (intended) test order entered		
	OE 06 Inappropriate/unnecessary (intended) blood product order entered		
	OE 07 Incorrect (unintended) test ordered		
	OE 08 Incorrect (unintended) blood product ordered		

Assurance of Confidentiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or institution is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)).

Public reporting burden of this collection of information is estimated to average 30 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering, and maintaining the data needed, and completing and reviewing the collection of information. An agency may not CDC 57.302 Rev. 3, v8.1



conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).

*Process Code	*Incident Code	*Total Incidents	*Total Adverse Reactions
	SC 00 Detail not specified		
	SC 01 Sample labeled with incorrect patient ID (intended patient drawn)		
	SC 02 Sample not labeled		
SC: Sample	SC 03 Wrong patient collected (sample labeled for intended patient)		
Collection	SC 04 Sample collected in wrong tube type		
(Service collecting the	SC 05 Sample quantity not sufficient (QNS)		
samples) Events that occur during patient sample collection.	SC 06 Sample hemolyzed		
	SC 07 Sample label incomplete/illegible for patient identifiers		
	SC 08 Sample collected in error (e.g., unnecessary/duplicate)		
	SC 09 Patient sample not collected (in error)		
	SC 10 Patient wristband incorrect/not available		
	SC 11 Sample contaminated		
	SH 00 Detail not specified		
	SH 00 Detail not specified SH 01 Sample sent without requisition		
	SH 02 Requisition and sample label don't match		
SHI Somplo	SH 02 Requisition and sample label don't match	-	
SH: Sample			
Handling (Service collecting the	SH 04 No Patient ID on requisition		
samples)	SH 05 No phlebotomist/witness identification	-	
Events that occur when a	SH 06 Sample sent with incorrect requisition type		
patient sample is sent for	SH 07 Patient information (other than ID) missing/incorrect on requisition		
testing.	SH 08 Requisition sent without sample		
	SH 09 Data entry incorrect/incomplete/not performed		
	SH 10 Sample transport issue (e.g., sample broken/inappropriate conditions)		
	SH 11 Duplicate sample sent in error		
SR: Sample Receipt	SR 00 Detail not specified		
(Transfusion Service)	SR 01 Sample accepted in error		
Events that occur when a	SR 02 Historical review incorrect/not performed		
sample is received by the transfusion service.	SR 03 Demographic review/ data entry incorrect/not performed		
transfusion service.	SR 04 Sample incorrectly accessioned		
	ST 00 Detail not specified		
	ST 01 Data entry incomplete/incorrect/not performed		
	ST 02 Appropriate sample checks incomplete/incorrect/not performed		
	ST 03 Computer warning overridden in error or outside SOP		
	ST 05 Sample test tube incorrectly accessioned		
	ST 07 Sample test tubes mixed up		
	ST 09 Sample test tube mislabeled (wrong patient identifiers)		
ST: Sample Testing	ST 10 Equipment problem/failure/not properly QC'd		
(Transfusion Service)	ST 12 Sample testing not performed		
Events that occur during patient sample testing by the transfusion service.	ST 13 Incorrect sample testing method chosen		
	ST 14 Sample testing performed incorrectly		
	ST 15 Sample test result misinterpreted		
	ST 16 Reagents used were incorrect/inappropriate/expired/not properly QC'd		
	ST 17 ABO/Rh error caught on final check		
	ST 18 Current/historical ABO/Rh mismatch		
	ST 19 Additional testing not performed		
	ST 20 Confirmatory check incorrect/not performed (at time work performed)		
	ST 21 Administrative check incorrect/not performed (record review/audit)		



ST 22 Sample storage incorrect/inappropriate

		*Tetel	*Total
*Process Code	*Incident Code	*Total Incidents	Adverse Reactions
	UM 00 Detail not specified		
	UM 01 Data entry incomplete/incorrect/not performed		
	UM 02 Record review incomplete/incorrect/not performed		
UM: Product	UM 03 Incorrect product (type) selected		
Manipulation/	UM 04 Incorrect product (patient) selected		
Processing/Testing	UM 05 Product labeled incorrectly (new/updated)		
(Transfusion Service)	UM 06 Computer warning overridden in error or outside SOP		
Events that occur while	UM 07 Special processing needs not checked		
testing, manipulating (e.g.,	UM 08 Special processing needs misunderstood or misinterpreted		
pooling, washing, aliquoting, irradiating),	UM 09 Special processing needs performed incorrectly		
processing, or labeling	UM 10 Special processing needs not performed		
blood products.	UM 11 Equipment problem/failure/not properly QC'd		
	UM 12 Reagents used were incorrect/inappropriate/expired/not properly QC'd		
	UM 13 Confirmatory check incorrect/not performed (at time work performed)		
	UM 14 Administrative check incorrect/not performed (record review/audit)		
	NB 01 Inventory less than usual par level due to supplier unable to meet usual		
	steady demand		
	NB 02 Demand for blood product exceeding usual par inventory level		
No Blood	NB 03 Incompatible/inappropriate units issued due to inventory constraints		
	when demand for blood product exceeds usual par inventory levels.		
	NB 04 Suboptimal dose (less than optimal quantity) transfusion or no		
	transfusion due to inventory constraints when demand for blood product		
	exceeds usual par inventory levels.		
RP: Request for	RP 00 Detail not specified		
Pick-Up	RP 01 Request for pick-up on wrong patient		
(Clinical Service)	RP 02 Incorrect product requested for pick-up		
Events that occur when the	RP 03 Product requested prior to obtaining consent		
clinical service requests	RP 04 Product requested for pick-up, but patient not available		
pick-up of a blood product from the transfusion	RP 05 Product requested for pick-up, but IV not ready		
service.	RP 06 Request for pick-up incomplete (e.g., patient ID/product type missing)		
	RP 07 Pick-up slip did not match patient information on product		
	UI 00 Detail not specified		
	UI 01 Data entry incomplete/incorrect/not performed		
	UI 02 Record review incomplete/incorrect/not performed		
	UI 03 Product issued for wrong patient		
	UI 04 Product issued out of order		
	UI 05 Product issue delayed		
UI: Product Issue	UI 06 LIS warning overridden in error or outside SOP		
(Transfusion Service)	UI 07 Computer issue not completed		
Events that occur when the	UI 08 Issued visibly defective product (e.g., clots/aggregates/particulate matter)		
transfusion service issues blood product to the clinical service.	UI 09 Not/incorrect checking of unit and/or patient information		
	UI 10 Product transport issues (e.g., delayed) by transfusion service		
	UI 11 Unit delivered to incorrect location by transfusion service		
	UI 12 Product transport issue (from transfusion service to clinical area)		
	UI 18 Wrong product issued for intended patient (e.g., incompatible)		
	UI 19 Inappropriate product issued for patient (e.g., not irradiated, CMV+)		
	UI 20 Confirmatory check incorrect/not performed (at time work performed)		
	UI 21 Administrative check incorrect/not performed (record review/audit)		
	UI 22 Issue approval not obtained/documented		



UI 23 Receipt verification not performed (pneumatic tube issue)

*Process Code	*Incident Code	*Total	*Total Adverse
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CS: Satellite Storage (Clinical Service)	CS 00 Detail not specified		
	CS 01 Incorrect storage conditions of product in clinical area		
	CS 02 Incorrect storage location in the clinical area		
Events that occur while product is stored and	CS 03 Labeling issue (by clinical staff) CS 04 Floor/clinic did not check for existing products in their area		
handled by the clinical	CS 04 Product transport issues (to or between clinical areas)		
service.			
	CS 06 Monitoring of satellite storage incorrect/incomplete/not performed		
	CS 07 Storage tracking/documentation incorrect/incomplete/not performed		
	UT 00 Detail not specified	_	
	UT 01 Administered intended product to wrong patient	_	
	UT 02 Administered wrong product to intended patient		
	UT 03 Transfusion not performed in error		
	UT 05 Bedside check (patient ID confirmation) incomplete/not performed		
	UT 06 Transfused product with incompatible IV fluid		
	UT 07 Transfusion delayed beyond pre-approved timeframe		
UT: Product	UT 09 Transfused unsuitable product (e.g., outdated/inappropriately stored)		
Administration	UT 10 Administered components in wrong order		
(Clinical Service)	UT 11 Appropriate monitoring of patient not performed		
Events that occur during the	UT 14 Transfusion volume too low (per order or SOP)		
administration of blood products.	UT 15 Transfusion volume too high (per order or SOP)		
products.	UT 16 Transfusion rate too slow (per order or SOP)		
	UT 17 Transfusion rate too fast (per order or SOP)		
	UT 18 Inappropriate preparation of product		
	UT 19 Transfusion protocol not followed (not otherwise specified)		
	UT 22 Order/consent check incorrect/not performed		
	UT 23 Transfusion documentation incorrect/incomplete/not performed		
	UT 24 Transfusion documentation not returned to transfusion service		
	UT 26 Transfusion reaction protocol not followed		
MS: Other	MS 99 Other		
	Total		