DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION

BIOLOGICAL PRODUCT DEVIATION REPORT

FDA USE ONLY
Date Received:
Date Reviewed:
BPD ID:
BPD No.

* Indicates required information	on
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* Indicates required information			
A. FACILITY INFORMATION		B. BIOLOGICAL PRODU	JCT DEVIATION (BPD) INFORMATION
1. Reporting Establishment Information 1. Establishment Tracking #		ng #	
* Reporting Establishment Name		2. Date BPD Occurred	
* Street Address Line 1		3. * Date BPD Discovere	ed
Street Address Line 2		4. * Date BPD Reported	
* City	* State	5. * Description of BPD ((use Page 2 for additional space)
Country	*Zip Code		
* Point of Contact			
* Telephone #		6. * Description of Contr (use Page 3 for additi	ibuting Factors or Root Cause ional space)
E-mail			
2. *Reporting Establishment Identification Number			
FDA Registration #			
CLIA#		7. * Follow-Up (use Pag	e 4 for additional space)
If the BPD occurred somewhere other than the above facility, please complete this Section and Section A4, otherwise continue onto Section B1.			
* Establishment Name			
Street Address Line 1		8. * Please Enter the 6 C	Character BPD Code
.Street Address Line 2			
* City	* State		
		C. UNIT / PRODUCT INF	FORMATION
* Country	* Zip Code		
4. Establishment Identification Number:		Please check the type	Blood (Continued on Page 5)
FDA Registration #		of product:	Non-Blood (Continued on Page 6)
CLIA#			

See OMB Statement on Page 8.

B5. DESCRIPTION OF BPD (continued)	

В6.	DESCRIPTION OF CONTRIBUTING FACTORS OR ROOT CAUSE (continued)

B7. FOLLOW-UP (continued)	

C1. BLOOD PRODUCTS / COMPONENTS

TOTAL NUMBER OF LOTS:		

Unit #	Collection Date (MM/DD/YYYY)	Expiration Date (MM/DD/YYYY)	Product Code	Disposition	Notification (Y,N,RN**)
1.)					
2.)					
3.)					
4.)					
5.)					
6.)					
7.)					
8.)					
10.)					
11.)					
12.)					
13.)					
14.)					
15.)					
16.)					
17.)					
18.)					

C2. NON-BLOOD PRODUCTS

TOTAL NUMBER OF LOTS:	

Lot #	Expiration Date (MM/DD/YYYY)	Product Type	Product Code	Disposition	Notification (Y, N)
1.)					
2.)					
3.)					
4.)					
5.)					
6.)					
7.)					
8.)					
10.)					
11.)					
12.)					
13.)					
14.)					
15.)					
16.)					
17.)					
18.)					

D.	ADDITIONAL COMMENTS

Public reporting burden for this collection of information is estimated to average 2 hours per response, including the time for reviewing instructions, searching existing data sources, adhering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to:

Department of Health and Human Services Food and Drug Administration Center for Biologics Evaluation and Research Office of Compliance and Biologics Quality 1401 Rockville Pike, Suite 200N, HFM-600 Rockville, MD 20852-1148

An agency may not initiate a collection activity without first obtaining OMB approval. The approved collection instrument should display a current and valid OMB control number, expiration date, public protection provision, and a burden statement on the approved collection instrument.