SECTION I				
Location (unit):				
Date Event Reported Da	Date Event First Occurred			
Time Event Reported AM/PM Time	me Event First Occurred AM/PM			
Patient Name:				
SECTION II				
Type of Event (<i>Check one</i>): Potential Actual (Medicatio	n event reached patient)			
	[*] for each medication (For Actual Events ONLY):			
*Variances: Total number of doses that reached the patient or were omitted **If more than six (6) medications are involved, must submit additional form(s)				
SECTION III				
Event Categories (Check all items within each event category that apply):				
	C. Transcribing:			
Drug Count Discrepancy	 Order transcribed for incorrect patient 			
A. Prescribing: Order written for incorrect patient 	Illegible transcription to Kardex			
Order is illegible	Incorrect dose transcribed Incorrect route transcribed			
Incorrect dose ordered	Incorrect dosing time transcribed			
Incorrect route ordered	Incorrect medication transcribed			
Incorrect dosing regimen Incorrect medication ordered	Allergy not transcribed Telephone order transcribed incorrectly			
Order written for medication to which patient has documented allergy	Order not transcribed			
Order written for medication to which patient has potential allergy	Medication not transcribed to new MAR			
 Order contains prohibited abbreviation Unclear/ambiguous order 	□ Other:			
Was order: Written Verbal				
□ Incomplete paperwork (<i>Example: Omitted off Formulary Request</i> Form/Antibiotic Order Form)	D. Administering:			
Other:	Medication administered to incorrect patient Incorrect medication administered			
	Incorrect dose of medication administered			
B. Dispensing:	 Medication administered by incorrect route 			
Medication dispensed to incorrect patient	Medication administered at incorrect time Medication administered at incorrect rate			
Incorrect medication dispensed to drawer	Medication administered at incorrect rate Medication to which patient has documented allergy administered			
 Incorrect dose of medication dispensed to drawer Pharmacy dispensed medication formulated for incorrect route 	Medication to which patient has potential allergy administered			
 Pharmacy dispensed medication formulated for incorrect route Medication to which patient has known allergy dispensed 	Extra dose of medication administered Dose of medication omitted			
Medication to which patient has potential allergy dispensed	 Dose of medication omitted Medication incorrectly prepared 			
	Equipment failure or malfunction			
Formulary medication not dispensed by Pharmacy because medication not available	Look alike/Sound alike medication			
Look alike/Sound alike medication	PYXIS not validated with MAR Other:			
Labeling error Ambiguous strength designation in labeling				
Pharmacy Order Entry	E. Monitoring			
Other:	Physician did not order appropriate follow-up (<i>vital signs, lab work, studies</i>			
	 			
	Nursing did not perform appropriate follow-up (vital signs, process lab work/study orders)			
	Nursing did not perform appropriate evaluation of patient (Accucheck, BP)			
	Other:			

SECTION IV				
Report of Event:				
Action Taken:				
Do you feel lack of training contributed to this event? Do you feel unit staffing contributed to this event? Did redeployment of staff directly contribute to this event? Did per diem staffing directly contribute to this event? Did overtime staffing directly contribute to this event? Do you feel working conditions contributed to this event?	 Yes Yes Yes Yes Yes Yes Yes 	 No No No No No No No 	 Unknown Unknown Unknown Unknown Unknown Unknown Unknown 	
SECTION V				
Best Estimate of Severity (To be completed by Physician for ACTUAL EVENTS ONLY): No patient harm. (Level 1; ORYX 1) Event resulted in the need for increased patient monitoring but no change in vital signs and no patient harm. (Level 2; ORYX 1) Event resulted in the need for increased patient monitoring with a change in vital signs but no ultimate patient harm, or any event that resulted in the need for increased laboratory monitoring. (Level 3; ORYX 2) Event resulted in the need for treatment with another drug or and increased length of stay, or acute care hospital admission, or that affected patient participation in an investigational drug study. (Level 4; ORYX 2) Event resulted in permanent patient harm or was life threatening. (Level 5; ORYX 3) Event resulted in patient death. (Level 6; ORYX 3)				
NOTE: Severity rating of Level 4 or above should be forwarded immediately to Registered Nurse Supervisor for critical incident reporting.				
Physician's Comment (If applicable):				

Physician Signature

Print Name

Date