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	Title: Adverse Event (AE) Report Form		

Reporting Instructions

Name/Initials:

Please email the completed AE report form, and a copy of all relevant source documents, **within 24 hours** of becoming aware of an AE to **aereports@azurity.com**.

NOTE: Please redact all patient personal information (medical record number, social security number, address, etc.)

Return To:

Azurity Pharmaceuticals

Attn: Drug Safety

Phone: 1-800-461-7449

Email: aereports@azurity.com

Date of This Report (DDMMYY):

Patient Information:

Name/Initials:

Male Female

If Female, Pregnant?

Yes No Unknown

Date of Birth:

Age:

Age Category:

Neonate Infant Child Adolescent Adult Elderly

Report Details:

Report Type:

Patient Health Care Professional (HCP) Other (Specify): _____

Name:

HCP Profession (MD/DO/PA/NP/RN/PharmD):

Phone:

Fax:


Street Address:

City/State/Zip:

Email:


IMPORTANT: If reporter is a healthcare professional, is it their opinion that the AE is related to the product?

Yes No

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Adverse Event(s) (AE) Information:			
Product:		Indication for Use:	
Dose Form:		Strength:	
Dose Regimen:		Expiration:	
Lot Number (If Available):			
Dates of Products Use:			
Action Taken with the Product: Continued, discontinued, unknown, increase/decrease dose			
Severity of Event (Mild, Moderate, Severe):			
Start Date of Event		Stop Date of Event	
Outcome of the Event:	<input type="checkbox"/> Resolved <input type="checkbox"/> Recovered with Minor Sequelae <input type="checkbox"/> Recovered with Major Sequelae <input type="checkbox"/> Ongoing/Continuing Treatment <input type="checkbox"/> Condition Worsening <input type="checkbox"/> Death <input type="checkbox"/> Unknown		

Briefly describe a summary of the adverse event(s) experienced by the patient, and include, any hospitalization, treatment given, and current outcome of the event(s).
Did the patient recover from the event; if so, what were the start date and resolution dates?

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Concomitant/Other Medication:	<input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown			
Name, Brand Name	Dose	Route (Oral, IV, etc.)	Start Date	Stop Date

-Please Provide an additional Page(s) if needed-

Thank you for taking time in providing this information

Reported by Azurity Representative:			
Name:		Date:	
Email Address:			
Phone:			