

## Adverse Drug Reaction (ADR) Reporting Form

A. Patient Details	
Patient initials:	Date of Birth: Day/Month/Year
Sex: <input type="checkbox"/> Male <input type="checkbox"/> Female [ <input type="checkbox"/> Pregnant <input type="checkbox"/> Not Pregnant ]	Weight:                      Height:

B. Suspected Drug/s						
Drug/s Name (Include generic name/s)	Manufacturer & Batch No.	Dose route	Dose frequency	Start date	End date	Indication/purpose of use

C. Concomitant Drug/s (Exclude those used to treat reaction)						
Drug/s Name (Include generic name/s)	Manufacturer & Batch No.	Dose route	Dose frequency	Start date	End date	Indication/purpose of use

D. Adverse Drug Reaction Description	
Adverse event including relevant tests/lab data and dates	Other relevant history, including preexisting medical conditions; (Diagnosis, allergies, pregnancy, hepatic, renal etc)
Date when event started:	Date when event disappeared (if applicable):

E. Action Taken					
<input type="checkbox"/> Drug discontinued	<input type="checkbox"/> Dose reduced	<input type="checkbox"/> Dose increased	<input type="checkbox"/> Dose not changed	<input type="checkbox"/> Unknown	<input type="checkbox"/> Not applicable

F. Outcome of ADR				
The patient: <input type="checkbox"/> Recovered; date:	<input type="checkbox"/> Recovering	<input type="checkbox"/> No improvement	<input type="checkbox"/> Died	<input type="checkbox"/> Unknown
Event subsided after stopping the suspected drug (Dechallenge)	<input type="checkbox"/> No	<input type="checkbox"/> Yes	<input type="checkbox"/> Unknown	<input type="checkbox"/> Unknown
Event reappeared after reintroducing to the suspected drug (Rechallenge)	<input type="checkbox"/> No	<input type="checkbox"/> Yes	<input type="checkbox"/> Not applicable	<input type="checkbox"/> Not applicable
Specific antagonist used	<input type="checkbox"/> No	<input type="checkbox"/> Yes; specify:		

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G. Seriousness of ADR		
<input type="checkbox"/> Patient died; date:	<input type="checkbox"/> Life threatening	<input type="checkbox"/> Hospitalization
<input type="checkbox"/> Permanent disability	<input type="checkbox"/> Congenital anomaly	<input type="checkbox"/> Prolonged hospitalization more than 24 hr.
<input type="checkbox"/> Required Emergency Room (ER) visit	<input type="checkbox"/> Required intervention to prevent permanent impairment/damage	
<input type="checkbox"/> None of the above (Not serious)		
Comments if any:		

H. Reporter Details		
Reporter Name:	Profession/Specialty:	
Center:	Address:	
Phone/Mobile:	E-mail:	
Fax:	Date:	Signature:

**Adverse Drug Reaction (ADR)** is a response to a medicinal product which is noxious and unintended. Response in this context means that a causal relationship between a medicinal product and an adverse event is at least a reasonable possibility

**Serious adverse reaction;** is an adverse reaction which:

- results in death,
- is life-threatening,
- requires in-patient hospitalization or prolongation of existing hospitalization,
- results in persistent or significant disability or incapacity or,
- is a congenital anomaly/birth defect.

**This form can be used by:**

- Physicians
- Pharmacists
- Dentists
- Nurses
- Other healthcare providers

**How to report:**

- Fill out the reporting form.
- Attach additional information, if needed.
- Use a separate form for each ADR.

**Please submit completed forms to:**

- Tabuk Pharmaceuticals
- Riyadh 11437 P.O.Box 28170
- Fax: +966 11 4782686
- Phone: +966 11 4777128 ext. 227, 233
- Email : pv.info@tabukpharmaceuticals.com

Thank You