N	ledicinal Sid	e Effects Reporting Form (by all)	Ea	ast India Ph	arma P-Vig ICSR Form 2550 F1 (version 1.0)								
	(base	ed on Suspected ADR Reporting Form, Pharmacovigilance Pro	me of India, Ministry	of Health & Family Welfare, Govt of India)									
RE	PORT DATE	- 20 <b>Co. REF No</b>			For internal use only								
٨	PATIENT: 1	. Initials 2. Sex 3. Body	kg	4. Residing at									
A.	PAHENI. 1	. ilittais 2. Sex 3. Bouy	ייי [		4. Nesiung at								
5. I	Date of Birth	or Age	at t	ime of event	yrs / mths / days								
B. SUSPECTED MEDICINE: (May tick >1 box wherever applicable. Use additional sheets if required.)													
1.	Brand		2.	Strength									
3.	Dosage	Oral □ Susp □ Syr □ Gel □ Tab □ Cap	4.	Composition									
	Form	Drops □ Oral □ Eye □ Ear □ Nasal		(Chemical /									
		Skin		Compound / Molecular /									
		<b>Injection</b> □ SC □ IM □ IV □ Infusion		Generic)									
5.	Dose	□ 1 □ 2 □ 3 □ 4 Tab / Cap / TSF (5 mL)	6.	Dose	□ 1x □ 2x □ 3x □ 4x □ 5x □ 6x daily								
	Amount / Quantity	or drops / puffs / mL / mg		Frequency & Duration	☐ Stat ☐ SOS ☐ Wkly FOR:days								
7.	START DATE		8.	STOP DATE ~									
	(& Time)			(last taken on)									
9.	Reason for		10.	Reason for	☐ Course / treatment complete								
	starting (Indication /			stopping	☐ Symptoms relieved / cured : by%								
	Disease /				Oversight / negligence by patient / provider								
1 1	Symptoms)		43	D	Appearance of side effects								
11.	Suggested * by (details)		12.	Prescriber / Doctor Name									
13.	Batch No.		14.	Expiry Date									
· Wı	ite CONT if con	tinuing * Doctor, Pharmacist, Self (from past expe	rience	e / no such). Frie	nd / Relative, Advertisement (media / online), etc.								
		) / ADVERSE EXPERIENCE : (May tick >1 box whe											
		(way tiens 1 box wife		START DATE	additional streets in required.)								
			۷.	(& Time)									
			3.	STOP DATE ~ (last noted on)									
1.	Detailed description			(.act notes only	☐ Barely noticeable ☐ Tolerable ☐ Intolerable								
	ueser iption		Δ	Severity /	Daily activity ☐ Disrupted ☐ Somewhat ☐ Not								
			••	Intensity	Based on these, state if :								
					☐ Mild 1-3 ☐ Moderate 4-7 ☐ Severe 8-10								
		☐ Continued drug without much discomfort			☐ Death (dd-mm-yyyy)								
		☐ Continued drug despite discomfort			☐ Life threatening ☐ Hospitalised / Prolonged								
5	Action	☐ Reduced drug amount / dose quantity			☐ Temporary impairment								
	taken	☐ Reduced dose frequency	6.	Seriousness	☐ Permanent diasability								
		☐ Had to stop medicine altogether ☐ Required treatment / antidote (specify):			☐ Birth defect / cancer								
		= nequired treatment, untidate (speeny).			□ Needed specific treatment								
_	0	□ Resovered □ Resovering □ Ongoing	0	D++'									
7.	Outcome of action	☐ Recovered ☐ Recovering ☐ Ongoing ☐ Fatal ☐ Unknown ☐ Others	8.	Past reaction to same drug	□ None □ Same □ Other times yrs ago □ NO prior use								
9.	When drug	☐ Reaction also reduced ☐ Continued unabated	10.	If / when re-	☐ Reaction reappeared ☐ Did not reappear								
	reduced	☐ Unknown ☐ Not applicable		introduced	☐ Unknown ☐ Not applicable								
11	Lab tests		12.	History of pre-existing	☐ Pregnancy ☐ Allergy ☐ Smoking ☐ Alcohol								
11.	with date			/ ongoing	□ Diabetes □ High BP Diseases of : □ Liver								
	(if any)			conditions /	□ Lungs □ Heart □ Kidney □ Nerve □ Eyes								
				diseases	☐ Bones ☐ Skin ☐ Others ☐ NIL								

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D.	ANY C	NY OTHER SIMULTANEOUS MEDICATION(s): (including herba										rbal	& self-medications, except those used (iii)									to treat side-effects) (iV)						
1.	Indica	ition																										
2.	Brand Form,																											
3.	Comp	ositio	n																									
4.	Dose &																											
5.	STAR																											
6. STOP DATE ~																						_						
7. Expiry Date																												
E. REPORTER: 1. Relation to patient 2. Date 20																												
3. Initials 4. Age yrs 5. Sex 6. Edu Qualification																												
7. <b>C</b> e	ell : +													8. <b>v</b>	wApp :	+												
9.	FULL I														10. <b>e</b> -	mai	il(s)											
• <u>lf</u> :	Profes ADDR th Desi Dept / U Bldg, Ir etc ± R Seal, S filled U	gnation Unit / estitution legn No ignature	i- n, e) <u>beh</u>	alf of reporter) <b>by company i</b>							prese	enta	tive ,	/ an	12. Residential ADDRESS (with Locality, Village / Town, PO, PS, Block, District, State, PIN code, Landmark, etc.)						14.	Λαο			— ¬,	urs 1	15. <b>S</b>	av 🗔
		IVAIVIL	. (DIO	ok oap	ntais	" <u>—</u>											$\overline{}$				<u> </u>	<b>Т</b>	·	<u> </u>		13 .	. J. <b>J</b>	
18.	Profes ADDR th Desi Dept / I Headqu Unit, Se	gnation Branch a uarters a eal, etc.	i- , , , )												Lo To St La	eside DDR cality wn, late, f	RESS ( y, Villa PO, Di PIN co lark, et	with ge / istrict de, c.)										
W W W W	REPORT ADRs (adverse drug reactions / side effects of medicines) – Just a Few Min could Save Many:  WHY: ADRs are the 4 <sup>th</sup> -6 <sup>th</sup> major cause of deaths worldwide. Filling out this form could help us ensure safer medicines, as the data generated will help in continual risk-benefit analysis of medicinal drugs.  WHO: Anybody can report: • Laymen / General Public: Patients (consumer), their friends / relatives, etc.  Specially • Health care professionals: Doctors, Dentists, Nurses, Paramedics, Pharmacists, etc. whose participation is vital.  WHEN: Even if you • don't have all details • are not certain that reactions have been caused by that product only  WHAT: Any adverse experience: • Adverse Drug Reactions • Adverse Events • Product Problems  WHERE: Please download, fill-up & submit / send, hard / soft / scanned copy of this form, to any of:  • Our company representatives • email to reportsideeffects@eastindiapharma.org  • Quality Control Dept, East India Pharmaceutical Works Ltd., 119 Biren Roy Road (West), Kolkata 700061, India.  • Online version of this form available at dedicated Medicinal Safety webpage of our website www. eastindiapharma.org  Disclaimer: Patient's identity will be kept Strictly Confidential & protected to the fullest extent, at all times & circumstances.  Reporter's identity too is not for public disclosure. Reporting is purely voluntary & aimed at patient safety only. Merely its compilation / submission DOES NOT amount to an admission of causality / contribution by the physician, manufacturer or																											