Patient Initials [][][]

PB-SAM Toxicity CRF v1.0



PRE-DEFINED SUSPECTED GRADE 3 or 4 DRUG TOXICITY

The medicines being used in PB-SAM are licensed drugs with a known profile of adverse reactions. Assessment of safety will therefore focus on <u>severe and causally related events</u>. Clinical or laboratory toxicity will be reported if Grade 3 or 4 according to the Division of AIDS table for grading severity of adverse events: http://rsc.techres.com/Document/safetyandpharmacovigilance/DAIDS AE Grading Table v2 NOV2014.pdf

If abnormalities are known to be already present at enrolment, or are due to other identifiable factors, this need not be reported as a causally-related toxicity. Only record new events whilst the child is receiving study drugs.

Always act in the best interests of the child. When possible, discuss any decisions regarding stopping study drugs because of suspected toxicity with the PB-SAM PI, clinical lead, designee or local safety monitor. They can also prove information on other aspect of management of adverse reactions.

A grade 3 or 4 toxicity event may also constitute an SAE, and should also be reported on an SAE CRF.

1. STUDY MEDICATION DETAILS				
1.1. Which study drugs is the child receiving? a) Pancreatic Enzymes/Placebo	☐ Pancreatic Enzymes/Placebo	: 24h Clock		
i. Date and time of start		m m		
ii. How many doses have been given?	doses			
b) Bile Acids/Placebo	☐ Bile Acids/Placebo			
i. Date and time of start		: 24h Clock m m		
ii. How many doses have been given?	doses			
tick all that apply	2. SUSPECTED GRADE 3 or 4 TOXICITY □ Not applicable	Definition		
2.1. Allergic & Cutaneous	☐ Generalized urticaria ☐ Angioedema with intervention indicated ☐ Symptoms of mild bronchospasm ☐ Acute anaphylaxis ☐ Life-threatening bronchospasm ☐ Laryngeal oedema	Clinical diagnosis Clinical diagnosis Clinical diagnosis Clinical diagnosis Clinical diagnosis Clinical diagnosis		

■ Not applicable

2.2. Diarrhoea

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Patient Initials [][][]

PB-SAM Number [2][0] [][][]

	□ Increase of ≥ 7 stools per 24-hour period			
	☐ IV fluid replacement indicated			
	☐ Life-threatening consequences (e.g., hypotensive shock)			
	□ Not applicable			
	= 1100 applicable			
	☐ Abnormal live	r function	ALT 5.0 to < 10.0 x ULN	
2.3. Hepatic				
•	☐ Abnormal live		Total bilirubin 2.6 to <5.0 x ULN	
	☐ Abnormal live		ALT > 10.0 x ULN	
	☐ Abnormal live		Total bilirubin >5.0 x ULN	
Write details in the description box on the next page,	ULN, upper limit of	normal based on local laborator	y reference values	
		If Y, Date:	Time:	
2.4.34				
3.1. Was the study drug stopped?	☐ Y* ☐ N	/ /		
		DD/MM/YYYY	:	
* if Y, also complete the drug discontinuation section	in the Study Conclus			
, , , also compress and aray alsoon an addition occurs.				
3.2. Description including concurrent med	ication manage	ment undertaken and out	come	
3.2. Description including concurrent incu	ication, manage	ment andertaken and oat	come	
4. Toxicity CRF completed by	Da	ate	Time	
The second secon				
initials		/ /		
initials		D/MM/YYYY	:	
F. Taviaita CDF Davisonad las	†		Time	
5. Toxicity CRF Reviewed by	Da	ate	Time	
initials	_	/	:	
	D	D/MM/YYYY		