



Serious Adverse Event - A

		1. SAE INITIAL DETAILS
1.1.	DATE of SAE onset	/// D D/M M/ Y Y Y Y
1.2.	DATE child seen or information received by research team	/// D D/M M/ Y Y Y Y
1.3.	Classification at presentation When the study team first became aware of the SAE. Tick the highest one applicable	□ Death □ Readmission to study hospital □ Readmission to non-study hospital □ Readmission is indicated but parent/carer declines admission □ Life-threatening event □ Persistent or significant disability/incapacity □ Event that prolongs hospitalisation whilst already in hospital (deterioration) □ Other serious medical event where medical intervention was required e.g. new diagnosis of TB, sickle cell disease
1.4.	Reported by (Tick one)	☐ Parent/caregiver ☐ Health Professional ☐ From medical records or discharge letter
1.5.	On study drugs at the onset of the SAE? (tick all that apply)	□ None □ Pancreatic enzymes/Placebo □ Urso/Placebo
1.6.	Any other medication in the last 7 days prior to onset of SAE? (tick all that apply) Give details with specific medications used in the text box below (section 1.07)	□ No medication □ Antibiotics □ Antimalarial □ IV fluids □ Blood Transfusion □ Anticonvulsants □ Anti-TB □ ART □ Traditional or Herbal □ Co-trimoxazole □ Yes, but unknown □ Other
1.7.		the SAE, where and when did it occur, was there any relation to timing of study drug flure, or relation to any other medication, who was involved? Describe any background factors or contributed to the SAE event

Patient Initials [][][]

1.8.	Describe the <u>new</u> clinical features of this SAE event (describe clinical findings from examination, medical records, or from caregiver)
1.9.	Describe any investigations or changes to lab results <u>RELEVANT</u> to this SAE event for its diagnosis or management* (i.e. Were there investigations done to investigate this SAE? The text needs to be sufficient to enable an independent reviewer to assess severity, type of illness and whether there is relationship to study drugs/medications)
1.10.	Describe the initial treatment given or other actions taken for this SAE*
1.11.	Describe the response to initial treatment *
1.12.	Describe any further clinical investigations and clinical progress*
1.13.	* write N/A if not applicable (e.g. death in the community) Suspected initial diagnosis for the cause of the SAE (e.g. pneumonia, sepsis etc)

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	2.0	Additional Docume	entation
2.1	Is there any additional documentation for this SAE		☐ YES ☐ NO
2.2	If Yes, select all that apply		☐ Clinical notes
	(Make copies of selected documents, certify and attach to the		☐ Lab results
	SAE CRF)		□ X ray
			☐ Death certificate
			☐ Discharge summary
			☐ Other, Specify
		3.0 PART A CRF C	OMPLETION
3.1	a) CRF Completed by (Initials) – to be signed when complete. Do not sign if any fields are empty		
	b) Date	/// 	- <u>Y</u>
	c) Time	: 24 h clock	
3.2.	a) CRF Reviewed by (Initials)		
	b) Date	/////////	-
	c) Time	:::24 h clock	



Serious Adverse Event - B

	4.0 SAE FINAL DETAILS
4.1	Circumstances : What was the SAE, where and when did it occur, was there any relation to timing of study drug administration or other procedure, or relation to any other medication, who was involved? Describe any background factors or co-morbidities that may have contributed to the SAE event
4.2	Describe the additional clinical features of this SAE event (describe clinical findings from examination, medical records, or from caregiver)
4.3	Describe any investigations or changes to results <u>RELEVANT</u> to this SAE event for its diagnosis or management* (i.e. Were there investigations done to investigate this SAE? The text needs to be sufficient to enable an independent reviewer to assess severity, type of illness and whether there is relationship to study drugs/medications)
4.4	Describe the treatment given or other actions taken for this SAE*
4.5	Describe the response to treatment, any further clinical investigations and clinical progress* * write N/A if not applicable (e.g. death in the community)

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	5.0 SAE CLASSIFICATION		
5.1	End date (dd/mm/yyyy)	/ or if no end date (died, or recovered) D D/M M/ Y Y Y On-going & receiving care Unknown	
5.2	Final Classification Tick the highest <u>one</u> applicable	□ Death □ Readmission to study hospital □ Readmission to non-study hospital □ Readmission is indicated but parent/carer declines admission □ Life-threatening event □ Persistent or significant disability/incapacity □ Event that prolongs hospitalisation whilst already in hospital □ Other serious medical event where medical intervention was required e.g. new diagnosis of TB, sickle cell disease	
	Was this event a suspected unexpected serious adverse reaction (SUSAR)?		

	6.0 RELATIONSHIP OF EVENT TO STUDY DRUGS	
6.1	No temporal relationship to drug; and alternate aetiology (clinical state, environmental or other interventions); and does not follow known pattern of response to study product	☐ No Relationship
	Unlikely temporal relationship to drug; and alternate aetiology likely (clinical state, environmental or other interventions); and does not follow known typical or plausible pattern of response to drug. Reasonable temporal relationship to drug; or event not readily produced by clinical state, environmental or other interventions; or similar pattern of response to that seen with other drugs Reasonable temporal relationship to drug; and event not readily produced by clinical state, environment, or other interventions or known pattern of response seen with other drugs	
	Reasonable temporal relationship to drug; and event not readily produced by clinical state, environment, or other interventions; and known pattern of response seen with study drugs	Definite

Patient Initials [][][]

	7.0 DIAGNOSES OF THE CAUSES OF SAE			
	Do not include unchanged conditions that existed prior to the SAE. Tick up to THREE diagnoses.			REE diagnoses.
	agnosis should be based on examination an			
7.1	General	☐ Anaemia	☐ Sickle Cell Disease	☐ Thalassaemia
		☐ Renal impairment	☐ Ileus	☐ Nephritis
		☐ Liver dysfunction	☐ Nephrotic syndrome	
		☐ Congenital cardiac disc	ease confirmed by echo	
7.0				
7.2	Respiratory	☐ LRTI/pneumonia	☐ Bronchiolitis	☐ URTI
		☐ Pulmonary TB		
	<u> </u>	☐ Otitis media	·	piration e.g. of feed
7.3	Infection	☐ Gastroenteritis	☐ Sepsis	☐ Confirmed
		Malaria —		_
		☐ Extra pulmonary TB	☐ Soft tissue infection	
		☐ HIV related illness	☐ Measles	☐ Varicella
		☐ Osteomyelitis	☐ Febrile illness unsp	ecified
		•	·	
		☐ Confirmed enteric feve	er 🛘 Typhoid/paratypho	oid with perforation
7.4	CNS	☐ Febrile convulsions	☐ Epilepsy	
		☐ Other encephalopathy	/ ☐ Hydrocephalus	☐ Cerebral palsy
		☐ Probable meningitis ☐	Clinically suspected mer	ningitis
		☐ LP confirmed meningit	is	
		☐ Confirmed diagnosis c	ongenital syndrome	
7.5	Other confirmed diagnosis	☐ Failed appetite test on	nly/severe malnutrition on	lly (readmission).
		☐ Suspected drug toxicit	: y (if due to study drug, comp	lete toxicity CRF)
		☐ Other known diagnosis	s:	

		8.0 Additional Docum	entation
	8.1	Is there any additional documentation for this SAE	☐ YES ☐ NO
	8.2	If Yes, select all that apply (Make copies of selected documents, certify and attach to the	☐ Clinical notes ☐ Lab results
		SAE CRF)	☐ X ray ☐ Death certificate
			☐ Discharge summary
			☐ Other, Specify
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9.0 Part B CRF Completion 9.1 a) CRF Completed by (Initials) – to be signed when complete. Do not sign if any fields are empty b) Date	
be signed when complete. Do not sign if any fields are empty	
Do not sign if any fields are empty	
$\frac{1}{D}\frac{1}{D}\frac{1}{M}\frac{1}{M}\frac{1}{Y}\frac{1}{Y}\frac{1}{Y}\frac{1}{Y}$	
D D/M M/Y Y Y Y	
c) Time	
:	
9.2 a) CRF Reviewed by (Initials)	
b) Date	
D D / M M / Y Y Y	
c) Time	
:	
Z-T II GIOGN	
Additional notes (Not for entry into database) all entries should be initialled and dated	
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END of SAE CRF