UKALL14



Serious Adverse Event (SAE) Report

NOT REQUIRED FOR 'REGISTRATION ONLY' SUB-STUDY PATIENTS (14-'3'-XXX OR 14-'4'-XXX)

Please <u>complete all sections</u> with details of any SAE occurring during the reporting windows outlined in protocol section 12.2.2 (and outside these timeframes if the event is felt to be a long term side effect). For guidance on which events to report please see trial protocol.

Please fax this form to the UKALL14 Co-ordinator at the CR UK & UCL Cancer Trials Centre on 020 7679 9861 within 24 hours of becoming aware of the event.

Trial details														
Trial title	A randomised trial for adults with newly diagnosed acute lymphoblastic leukaemia													
Trial acronym	UKALL14	4				EudraCT nur	mber		2009-012717-22					
Patient details														
Patient initials						Patient trial Randomised p Safety data no	patients only		14 - 🗌 -	14				
Sex	Male Female					Date of birth	h							
Hospital						Treating Clir	nician							
Type of report	First Update Final					Height		cm		Weight kg				
Trial arm	B Randomisation: 0=N/A 1=B1 2=B2 T Randomisation: 0=N/A 1=T1 2=T2 P Randomisation: 0=N/A 1=P1 2=P2													
Trial treatment														
Trial treatment											Tick if no IMPs given to date			
Trial treatment Drug Name		Brand	Dose	Unit	Frequency	Is this full dose?	Route	d d	Start date	Ongoing?	Tick if no IMPs given to date End date			
	ase C	Brand Oncaspar	Dose	Unit	Frequency		Route			Ongoing?	End date			
Drug Name			Dose	Unit	Frequency	dose?	Route				End date			
Drug Name Pegylated Asparagin	N	Dncaspar	Dose	Unit	Frequency		Route				End date			
Drug Name Pegylated Asparagin Rituximab	A	Dncaspar Mabthera	Dose	Unit	Frequency		Route				End date			

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	NOT REQUIRED FOR 'REGISTRATION ONLY' SUB-STUDY PATIENTS (14-'3'-XXX O	R 14-'4'-XXX) Patient trial number: 44- 📖 - 📖 🦳
	a concise medical description of the event including all relevant symptoms and complete age for each event that meets the definition of serious)	Continued on a separate sheet:
Date site became aware of SAE:	If aware more than 24 hours before submission, reason for late reporting:	
No. of events included in this report:	If hospitalisation, please provide: Admission date	Discharge date

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NOT REQUIRED FOR 'REGISTRATION ONLY' SUB-STUDY PATIENTS (14-'3'-XXX OR 14-'4'-XXX) Patient trial number:

Serious Adverse Event (SAE)											
COMPLETE A SEPAR	ATE PAGE FOR EAC	H EVENT THAT N	IEETS THE DEFINITI	ON OF SERIOUS (photo	ocopy this page as neces	sary for each event)					
Name of event (use CTCAE version 4.0)	Grade	Date	of onset	Ongoing?	Date resolved						
		d d m		<u> у </u> N							
Why was the event serious? (choose most se			Outcome								
Resulted in death				Fatal							
Life-threatening				Not resol	ved						
Required new or prolonged hospita	lisation			Resolved							
Resulted in persistent or significant	disability/incapacity	y		Resolved	with sequelae						
Resulted in congenital anomaly/bir	th defect			Resolving	5						
Other (specify)				Not asses	ssable						
			SAE Assessn	nent							
Drug Name	ationship to eve one code only) e ely bly ably ably itely	ent	Action take (Enter <u>one</u> code of 0 = Dose not char 1 = Dose reduce 2 = Drug withdra 3 = Not applical	only) nged ed awn	OFFICE USE ONLY Event expected for the drugs 1 = Expected 2 = Not Expected						
Pegylated Asparaginase											
Rituximab											
Nelarabine											
Palifermin											
Office use only											
Event No: 14	Was the	event a SUSAR?	*Y	Date SAE entered of	ered on database						
*Date reported to MHRA:	*Date re	ported to Main RE		m y y *	*Reported to Principal Investigators						
Form checked by (signature)	Dat			Date checked by clinic							

UKALL14



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		NOT REQUIRE	D FOR 'RE	GISTRAT	ION ONLY' SU	B-STUDY I	PATIENTS (14-'3'-XXX OR 14-'4'->	(XX) Patient trial	number: 14 - 🗌 - 🗌 🗌					
Concomitant Me	dications	 List non-IMP drug 	s given with catments or	nin the <u>30</u> treatmen	<u>days</u> prior to SA ht for SAE, as the	AE onset, in	cluding non-IMP treatment for ALI orded elsewhere on the form.	Con	tinued on a rate sheet:					
Drug Name	Brand	Indication	Dose	Units	Frequency	Route	Start date	Ongoing?	End date					
							d d m m m y y		d d m m m y y					
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Serious Adverse Event (SAE) Report											лк									
NOT REQUIRED FOR 'REGISTRATION ONLY' SUB-STUDY PATIENTS (14-'3'-XXX OR 14-'4'-XXX) Patient trial number: 14-																				
Did the patient receive any treatment for this SAE?						es, please sp								a separate sheet		ү [N			
Drug Name Brand Indication Dose				Units	Frequency	Route		Start date C					Ongoing?	Ongoing? End date						
								b t								d d	m n		y] [<u>y</u>
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Any relevant test	s / laboratory	data?	N	(If yes, ple	ease specify be	low)														
Date		Test		Results																
d d m m m				Results pending:																
				Results pending:																
				Results pending:																
				Results pending:																
				Results pending:																
												Resu	ts per	ding	: [Y				
																Resu	ts per	ding	: [Y
Any relevant med	dical history / d	concurrent conditions?		Y (If yes, please specify below)																
																	Г	_	Г	
				•		Was the event expected in view of patient's medical history?										N				
Signature: PI or other participating clinicians only					nt name:	Date of report:														