



# UKALL14

## Serious Adverse Event (SAE) Report



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

Please complete all sections 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	EudraCT number	2009-012717-22						
Patient details									
Patient initials	<input type="text"/> <input type="text"/> <input type="text"/>	Patient trial number	14 - <input type="text"/> - <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>						
		<i>Randomised patients only 14-1-XXX or 14-2-XXX Safety data not collected for 'Registration only' patients</i>							
Sex	<input type="checkbox"/> Male <input type="checkbox"/> Female	Date of birth	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>						
Hospital		Treating Clinician							
Type of report	<input type="checkbox"/> First <input type="checkbox"/> Update <input type="checkbox"/> Final	Height	<input type="text"/> <input type="text"/> <input type="text"/> cm	Weight	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> kg				
Trial arm	<b>B Randomisation:</b> <input type="checkbox"/> 0=N/A 1=B1 2=B2 <b>T Randomisation:</b> <input type="checkbox"/> 0=N/A 1=T1 2=T2 <b>P Randomisation:</b> <input type="checkbox"/> 0=N/A 1=P1 2=P2								
Trial treatment									
<input type="checkbox"/> Tick if no IMPs given to date									
Drug Name	Brand	Dose	Unit	Frequency	Is this full dose?	Route	Start date	Ongoing?	End date
					<input type="checkbox"/> Y <input type="checkbox"/> N		<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>
Pegylated Asparaginase	Oncaspar				<input type="checkbox"/> Y <input type="checkbox"/> N		<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>
Rituximab	Mabthera				<input type="checkbox"/> Y <input type="checkbox"/> N		<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>
Nelarabine	Atriance				<input type="checkbox"/> Y <input type="checkbox"/> N		<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>
Palifermin	Kepivance				<input type="checkbox"/> Y <input type="checkbox"/> N		<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>
<b>Most recent phase of protocol treatment</b> <input type="checkbox"/> (1=Phase 1 induction, 2= Phase 2 Induction, 3= Intensification, 4= Consolidation, 5= Maintenance, 6= Myeloablative transplant, 7= Non-Myeloablative transplant)					<b>Start Date of most recent phase of protocol treatment, prior to SAE:</b> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>		<b>What was the last IMP given prior to SAE?</b> (1=Pegylated asparaginase, 2= Rituximab, 3= Nelarabine, 4= Palifermin)		



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

**Serious Adverse Event (SAE)**

COMPLETE A SEPARATE PAGE FOR EACH EVENT THAT MEETS THE DEFINITION OF SERIOUS (photocopy this page as necessary for each event)

Name of event (use CTCAE version 4.0)	Grade	Date of onset	Ongoing?	Date resolved
	<input type="text"/>	<input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	<input type="text"/> Y <input type="text"/> N	<input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>

Why was the event serious? (choose most serious)	Outcome
<input type="checkbox"/> Resulted in death	<input type="checkbox"/> Fatal
<input type="checkbox"/> Life-threatening	<input type="checkbox"/> Not resolved
<input type="checkbox"/> Required new or prolonged hospitalisation	<input type="checkbox"/> Resolved
<input type="checkbox"/> Resulted in persistent or significant disability/incapacity	<input type="checkbox"/> Resolved with sequelae
<input type="checkbox"/> Resulted in congenital anomaly/birth defect	<input type="checkbox"/> Resolving
<input type="checkbox"/> Other (specify) _____	<input type="checkbox"/> Not assessable

**SAE Assessment**

Drug Name	Causal relationship to event <i>(Enter <u>one</u> code only)</i> 0 = None 1 = Unlikely 2 = Possibly 3 = Probably 4 = Definitely	Action taken <i>(Enter <u>one</u> code only)</i> 0 = Dose not changed 1 = Dose reduced 2 = Drug withdrawn 3 = Not applicable	OFFICE USE ONLY Event expected for the drugs  1 = Expected 2 = Not Expected
Pegylated Asparaginase	<input type="text"/>	<input type="text"/>	<input type="text"/>
Rituximab	<input type="text"/>	<input type="text"/>	<input type="text"/>
Nelarabine	<input type="text"/>	<input type="text"/>	<input type="text"/>
Palifermin	<input type="text"/>	<input type="text"/>	<input type="text"/>

Office use only			
Event No: <b>14</b> - <input type="text"/> - <input type="text"/> <input type="text"/> <input type="text"/> - <input type="text"/> <input type="text"/> - <input type="text"/> <input type="text"/>	Was the event a SUSAR? <input type="text"/> *Y <input type="text"/> N	Date SAE entered on database <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	
*Date reported to MHRA: <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	*Date reported to Main REC <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	*Reported to Principal Investigators <input type="text"/> Y	
Form checked by (signature)	Date <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	Date checked by clinical reviewer <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <small>d d m m m y y</small>	



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**Did the patient receive any treatment for this SAE?**  Y  N *(If yes, please specify below)* **Continued on a separate sheet?**  Y  N

Drug Name	Brand	Indication	Dose	Units	Frequency	Route	Start date		Ongoing?	End date	
							d	d		m	m
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>
							<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y <input type="checkbox"/> N	<input type="text"/>	<input type="text"/>

**Any relevant tests / laboratory data?**  Y  N *(If yes, please specify below)*

Date	Test	Results									
d	d	m	m	y	y						
<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y									
<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y									
<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y									
<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y									
<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y									
<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y									
<input type="text"/>	<input type="text"/>	<input type="checkbox"/> Y									

**Any relevant medical history / concurrent conditions?**  Y  N *(If yes, please specify below)*

Was the event expected in view of patient's medical history?  Y  N

<b>Signature:</b> PI or other participating clinicians only	<input type="text"/>	<b>Print name:</b>	<input type="text"/>	<b>Date of report:</b>	<input type="text"/>
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