CIC:	Hospital UPN:	Patient UIC	HSCT Date:
	HSCT - Min	imum Essential C	
		Centre Identification	
	Unit:		
		Patient Data	
□ No □ Yes	yyyy - mm - dd national / international study / s: Name of study / trial atient Number or Code (UPN) ations will not be accepted withour med in the same patient must be	U	nknown
	(first name(s)	_family name(s)) Sex:	Female
	Prir	mary Disease Diagnosis	
	nosis: yyyy - mm - dd DIAGNOSIS (CHECK THE DISEAS	 SE FOR WHICH THIS TRANSPLANT WAS PERFO	DRMED)
related Pre Precursor Therapy relat Secondary Ac Chronic Leuk	elogenous Leukaemia (AML) ecursor Neoplasms Lymphoid Neoplasms (old ALL) ted myeloid neoplasms (old cute Leukaemia) aemia yeloid Leukaemia (CML) mphocytic Leukaemia (CLL) kin Disease	 Myeloma/Plasma cell disorder Solid Tumour Myelodysplastic syndromes / Myeloproliferative neoplasm MDS MDS/MPN Myeloproliferative neoplasm Bone marrow failure including Aplastic anaemia Inherited disorders Primary immune deficiencies Metabolic disorders 	 ☐ Histiocytic disorders ☐ Autoimmune disease ☐ Juvenile Idiopathic Arthritis ☐ Multiple Sclerosis ☐ Systemic Lupus ☐ Systemic Sclerosis ☐ Haemoglobinopathy

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd
	LYMF	PHOMAS (main disease code 3		
		Hodgkin Lymphomas		
		Disease		
Date of Initial Diagnosis:	уууу - mm - dd	-		
Classification:				
Nodular lymphocyte				
☐ Classical predominan☐ Other , specify:				

CIC:	Hospital UPN:	Patient UIC	HS	CT Date:	уууу - mm - dd
		ALL LV/ADLIONAAO			yyyy - mm - dd
		ALL LYMPHOMAS			
		Treatment Pre-HSCT			
Treatment pre-HSCT	Enter first day	y of treatment and mark all drugs fron	n that date until co	onditionin	g
Yes Date of treatmer	nt yyyy - mm - da				
Drugs given					
Antibodies:	Alemtuzuma	b (MabCampath) (CD52)			
	Brentuximat	(Adcetris) (CD30)			
	Obinutuzum	ab (Gyzeva) (CD20)			
	Ofatumumal	b (Azerra) (CD20)			
	☐ Rituximab (N	/labthera) (CD20)			
	other antibo	dy, specify			
Radioimmunotherapy:	☐ Bexxar (CD2)	0) (radiolabelled MoAB)			
	☐ Zevalin (CD2	0) (radiolabelled MoAB)	Relapse	/progre	ssion under this drug
			Yes	No l	Jnknown
Specific inhibitors:	☐ ABT-199 (BC	L2-Inhibitor)			
	Crizotinib (A	LK-Inhibitor)			
	☐ CC-292 (B ce	ll receptor kinase inhibitor)			
	☐ Ibrutinib (B o	cell receptor kinase inhibitor)			
	☐ Idelalisib (B o	cell receptor kinase inhibitor)			
	other inhibit	or, specify			
Other:	☐ Bortezomib	(Velcade)			
	Lenalidomid	e (Revlimid)			

Other, specify _____

CIC: Ho:	spital UPN:	Patient U	ır	HSCT Da	to:
CIC. TIO	spitai OFN.			TISCT Da	te: yyyy - mm - dd
		ALL LYMPH	OMAS		
		Status at F	ISCT		
Date of this HSCT:	im - dd				
Number of prior lines of treatment	_ 1	<u> </u>	3 or more:	none	Unknown
(since diagnosis if 1st transplant, or since	e last reported transplant)				
Technique used for disease a	ssessment:				
CT scan c	done No	☐ Yes			
	PET Neg	gative	sitive 🗌 N	Not evaluated	
STATUS					
Never treated					
Complete remission (CR) Unconfirmed (CRU*)	Confirmed	d			
-	onse with persistent sca	an abnormalities of u	nknown significand	ce	
Partial response (PR) – (with o Stable disease	r without a prior CR)				
Untreated relapse (from a pre-	vious CR) / untreated pr	ogression (from a pr	evious PR)		
☐ Chemorefractory relapse or pr☐ Disease status unknown	ogression, including pri	mary refractory disea	ise		
Was this patient refractory to any l	line of chemotherapy be	efore this HSCT?	☐ No [Yes	
Number of Complete (CR, CRu) ach Count <u>all</u> CR including this one if applic		or to this HSCT:			
Number of Partial remissions (PR) a Count <u>all</u> PR including this one if apple		prior to this HSCT: _			

CIC: He	ospital UPN:	Patient UIC	HSCT Date:	уууу -	mm - d	'd
		HSCT				
	system used	sky □ 50 □ 60 □	70 🗆 80 🗆 90 🗀	□ 1 00)	
forror et al., Blood, 2005 Oct 1		orbidity Index /www.ncbi.nlm.nih.gov/t	omc/articles/PMC1895304/			
Vas there any <i>clinically signific</i> oreparative regimen? No Yes				to the		
Comorbidity		Definitions		No	Yes	N/E
Solid tumour, previously present	Treated at any time point melanoma skin cancer Indicate type		ory, excluding non-			
nfammatory bowel disease	Crohn's disease or ulcerat	tive colitis				
Rheumatologic	SLE, RA, polymyositis, mix	ked CTD, or polymyalgia r	rheumatica			
nfection	Requiring continuation of	f antimicrobial treatment	t after day 0			
Diabetes	Requiring treatment with diet alone	insulin or oral hypoglyca	nemics but not			
Renal: moderate/severe	Serum creatinine > 2 mg/ transplantation	dL or >177 μmol/L, on di	alysis, or prior renal			
Hepatic: mild moderate/ severe	ULN, or AST/ALT betweer	n ULN and 2.5 × ULN	AST/ALT greater than 2.5			
Arrhythmia	Atrial fibrillation or flutte arrhythmias	r, sick sinus syndrome, o	r ventricular			
Cardiac	Coronary artery disease, 6 50%, or shortening fraction	_	myocardial infarction, EF ≤			
Cerebrovascular disease	Transient ischemic attack	or cerebrovascular accid	lent			
Heart valve disease	Except mitral valve prola	pse				
Pulmonary: moderate	DLco and/or FEV1 66-80%	6 or dyspnoea on slight a	ctivity			
severe	DLco and/or FEV1 ≤ 65%	or dyspnoea at rest or re	quiring oxygen			
Obesity	Patients with a body mas	s index > 35 kg/m2				
Peptic ulcer	Requiring treatment					
Psychiatric disturbance	Depression or anxiety req	quiring psychiatric consul	tation or treatment			
	,			1		

Were there any other major clinical abnormalities prior to the preparative regimen? Specify.....

CIC:	Hospital UP	N: Pa	ient UIC	HSCT Date:	yyyy - mm - dd
		Type of HSC	T (Autologous	5)	
□ Au	itologous				
	Source of the Stem cells (check all that apply):	☐ Bone marrow☐ Cord blood		pheral blood er:	
	Graft manipulation ex-vivo other than for RBC removal o	r volume reduction			
	☐ No ☐ Yes: G	enetic manipulation of the	e graft: 🔲 No	☐ Yes:	
	☐ IF AUTOLOGOUS, O	CONTINUE TO "CHRONOLO	OGICAL NUMBER OF H	ISCT"	

CIC: Hospital UPN:	Patient UIC	HSCT Date: yyyy - mm - dd
ŀ	HSCT (Continued)	
Chronological number of HSCT for this patient? If >1, date of last HSCT before this one If >1, type of last HSCT before this one		
If >1, was last HSCT peformed at another institut If >1, please submit an Annual follow up for subsequent transplant as the date of last co (This is so we can capture relapse data and other part of a planned multiple (sequential) graft p	Name of the institution City TM before proceeding, giving the date of the ontact other events between transplants).	
☐ No ☐ Yes		
Pr	reparative Regimen	
Preparative (conditioning) regimen given? No (Usually Paed Inherited Disorders only) Go Yes Prugs No Yes No Yes	☐ Unknown	

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	
				yyyy - mm - dd

Specification and dose of the preparative regimen

TOTAL PRESCRIBED CUMULATIVE DOSE* as per protocol:					
DRUG (given before day 0)	DOSE		UNIT	S	
Ara-C (cytarabine)	2002	mg/m2	mg/kg		
ALG, ATG (ALS/ ATS)		mg/m2	mg/kg		
Animal origin: Horse					
Rabbit					
Other, specify					
Bleomycin		mg/m2	mg/kg		
Busulfan		mg/m2	☐ mg/kg	mg x hr/L	
Oral IV Both				micromol x min/L mg x min/mL	
BCNU		mg/m2	mg/kg		
Bexxar (radio labelled MoAB)		☐ mCi	☐ MBq		
CCNU		mg/m2	mg/kg		
Campath (AntiCD 52)		mg/m2	mg/kg		
☐ Carboplatin		mg/m2	☐ mg/kg	mg x hr/L micromol x min/L mg x min/mL	
☐ Cisplatin		mg/m2	mg/kg		
☐ Clofarabine		mg/m2	mg/kg		
Corticosteroids		mg/m2	mg/kg		
☐ Cyclophosphamide		mg/m2	mg/kg		
☐ Daunorubicin		mg/m2	mg/kg		
Doxorubicin (adriamycine)		mg/m2	☐ mg/kg		
Epirubicin		mg/m2	mg/kg		
Etoposide (VP16)		mg/m2	mg/kg		
☐ Fludarabine		mg/m2	mg/kg		
☐ Gemtuzumab		mg/m2	mg/kg		
☐ Idarubicin		mg/m2	mg/kg		
☐ Ifosfamide		mg/m2	☐ mg/kg		
☐ Imatinib mesylate		mg/m2	mg/kg		
☐ Melphalan		mg/m2	mg/kg		
Mitoxantrone		mg/m2	mg/kg		
☐ Paclitaxel		mg/m2	☐ mg/kg		
Rituximab (mabthera, antiCD20)		mg/m2	☐ mg/kg		
☐ Teniposide		☐ mg/m2	☐ mg/kg		
☐ Thiotepa		mg/m2	☐ mg/kg		
☐ Treosulphan		mg/m2	☐ mg/kg		
Zevalin (radiolabelled MoAB)		☐ mCi	☐ MBq		
Other radiolabelled MoAB		☐ mCi	☐ MBq		
Specify					
Other MoAB, specify		mg/m2	mg/kg		
Other, specify		mg/m2	mg/kg		

^{*}Report the total prescribed cumulative dose as per protocol. Multiply daily dose in mg/kg or mg/m² by the number of days; e.g. for Busulfan given 4mg/kg daily for 4days, total dose to report is 16mg/kg

^{**}AUC = Area under the curve

CIC:	Hospital UPI	l:	Patient UIC	HSCT D	ate: yyyy - mm - dd
Total Body Irradiation (TBI) <u>No</u>	☐ Yes		tion dose as per protocol	
		Nu	mber of fractions	over	radiation days
TLI, TNI, TAI (lymphoid, nodal, abdominal)	□ No	☐ Yes	: Total prescribed radi	iation dose as per protocol	Gу
			0 : 10: 1		
			Survival Stat	ius	
Survival Status on date	of HSCT lead				
		n of the prep	parative regimen and date o	of HSCT	
Main Cause of De	•	only one m	ain cause):		
Relapse or Prog		ent disease			
Unknown	4450				
Other			 heck as many as approp	riatal.	
GVHD	ory cause or i	Jeatii (C	песк аз тапу аз арргор	natej.	
	itial pneumonit	is			
☐ Pulmor	nary toxicity on:				
	acterial				
	ral				
	ingal arasitic				
	nknown				
	on/Poor graft f	unction			
	of severe Ven		isorder (VOD)		
☐ Haemo	rrhage				
Cardiad	toxicity				
Centra	l nervous syste	n (CNS) toxio	city		
Gastro	intestinal (GI) t	oxicity			
Skin to					
Renal f					
	le organ failure				
Utner,	эресту				