CIC:	Hospital UPN:	Patient UIC	HSCT Date: yyyy - mm - dd					
	HSCT - Min	imum Essential REGISTRATION - DAY 0						
Centre Identification								
	Unit:	_						
Patient Data								
No Yes: Hospital Unique Pati Compulsory, registratic All transplants perform the patient and not to	•	ut this item. e registered with the same patient identific	Jnknown					
Date of birth:	yyyy - mm - dd	Sex: Male	Female					
	Prir	mary Disease Diagnosis						
Acute Leukaem Acute Myelo related Precursor Ly Therapy related Secondary Acute Chronic Leukaem Chronic Myelo	nia ogenous Leukaemia (AML) ursor Neoplasms omphoid Neoplasms (old ALL) d myeloid neoplasms (old te Leukaemia) emia eloid Leukaemia (CML) phocytic Leukaemia (CLL)	☐ Myeloma/Plasma cell disorder ☐ Solid Tumour ☐ Myelodysplastic syndromes /	Histiocytic disorders Autoimmune disease Juvenile Idiopathic Arthritis Multiple Sclerosis Systemic Lupus Systemic Sclerosis Haemoglobinopathy					
Other diagnosis	, specify:							

CIC:	Hospital UPN:	Patient UIC	HSCT	Date:
	AC	UTE LEUKAEMI Other acute leuk	AS (main disease code 1)	
		Disease		
Date of initial diagno	sis: yyyy - mm - dd			
Classification: Acute Leukaemias of	ambiguous lineage			
☐ Mixed pheno ☐ Natural killer (NK)-		a/lymphoma		
		Secondary O	rigin?	
			No Yes Unknown TE LEUKAEMIA, ANSWER THE FO	LLOWING QUESTION
		Status at H	SCT	
Date of this HSCT:	yyyy - mm - dd			
STATUS		NUMBER	TYPE OF REMISSION	
Primary induction	failure			
Complete haemato	ological remission (CR)	☐ 1st ☐ 2nd ☐ 3rd or higher	CYTOGENETIC REMISSION No Yes Not evaluated Not Applicable* Unknown	MOLECULAR REMISSION No Yes Not evaluated Not Applicable* Unknown
Relapse		☐ 1st ☐ 2nd ☐ 3rd or higher		

^{*} No abnormalities detected prior to this time point

CIC: Hosp	oital UPN: Patient UIC H	SCT Date:	уууу -	mm - d	d
	HSCT				
Performance score Score		□ 90 □	100		
Weight (kg):	neight (chi):				
	Comorbidity Index				
forror et al., Blood, 2005 Oct 15;	106(8): 2912-2919: http://www.ncbi.nlm.nih.gov/pmc/articles/PM	IC1895304/			
Vas there any <i>clinically significar</i> oreparative regimen? No Yes	t co-existing disease or organ impairment at time of patient assessr	ment just prior	to the		
Comorbidity	Definitions		No	Yes	N/E
Solid tumour, previously present	Treated at any time point in the patient's past history, excluding no melanoma skin cancer	on-			
	Indicate type				
nfammatory bowel disease	Crohn's disease or ulcerative colitis				
Rheumatologic	SLE, RA, polymyositis, mixed CTD, or polymyalgia rheumatica				
nfection	Requiring continuation of antimicrobial treatment after day 0				
Diabetes	Requiring treatment with insulin or oral hypoglycaemics but not diet alone				
Renal: moderate/severe	Serum creatinine > 2 mg/dL or >177 μ mol/L, on dialysis, or prior retransplantation	enal			
Hepatic: mild	Chronic hepatitis, bilirubin between Upper Limit Normal (ULN) and ULN, or AST/ALT between ULN and 2.5 × ULN				
moderate/ severe	Liver cirrhosis, bilirubin greater than 1.5 × ULN, or AST/ALT greate × ULN	r tnan 2.5			
Arrhythmia	Atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias				
Cardiac	Coronary artery disease, congestive heart failure, myocardial infar 50%, or shortening fraction in children (<28%)	rction, EF ≤			
Cerebrovascular disease	Transient ischemic attack or cerebrovascular accident				
Heart valve disease	Except mitral valve prolapse				
Pulmonary: moderate	DLco and/or FEV1 66-80% or dyspnoea on slight activity				
severe	DLco and/or FEV1 ≤ 65% or dyspnoea at rest or requiring oxygen				
Dbesity	Patients with a body mass index > 35 kg/m2				
Peptic ulcer	Requiring treatment				
Psychiatric disturbance	Depression or anxiety requiring psychiatric consultation or treatm	ent			

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

CIC:	Hospital UPN	I: Patient UIC		HSCT Date:	nm - dd
		Type of HSCT (Aut	ologous)		
	Autologous				
	Source of the Stem cells (check all that apply):	☐ Bone marrow☐ Cord blood	☐ Peripheral bl		
	Graft manipulation ex-vivo other than for RBC removal or	volume reduction			
	☐ No ☐ Yes: Ge	enetic manipulation of the graft:	□ No □ Yes	s:	
	☐ IF AUTOLOGOUS, C	ONTINUE TO "CHRONOLOGICAL NU	JMBER OF HSCT"		

CIC:	Hospital UPN:	Patient UIC	HSCT Date:
			HSC1 Date:
		HSCT (Continued)
Chronol	agical number of USCT for this nationt?	1 1	
Chronoi	ogical number of HSCT for this patient?	1 1	
	If >1, date of last HSCT before this one	yyyy - mm - dd	-
	If >1, type of last HSCT before this one	☐ Allo ☐ Auto	
	, -, -, -,		
	If >1, was last HSCT peformed at another	institution? No	Vac. CIC if known
	11 > 1, was last riser perorimed at another		Yes: CIC if known
		Name of the ins	titution
		City	
	If >1, please submit an Annual follow	up form before proceeding, givin	g the date of the
,	subsequent transplant as the date of	last contact	
	(This is so we can capture relapse date	a and other events between trans	plants).
⊔sct ,	art of a planned multiple (sequential)	graft protocol (program)?	
		grant protocor (program):	
	No Yes		
		Preparative Regime	en
Prepar	ative (conditioning) regimen given?		
	No (Usually Paed Inherited Disorders o	nly) Go to GvHD Prophylaxis	
	Yes		
Davida			
Drugs	☐ No ☐ Yes		
(include	any active agent be it chemo, monoclonal o	antibody, polyclonal antibody, serothe	rapy, etc.)

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)		mg/m2	mg/kg			
ALG, ATG (ALS/ ATS)		mg/m2	mg/kg			
Animal origin: Horse						
Rabbit						
Other, specify						
			□ ma/lea			
Bleomycin Busulfan		☐ mg/m2	☐ mg/kg			
		mg/m2	mg/kg	mg x hr/L micromol x min/L		
☐ Oral ☐ IV ☐ Both				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		
Carbopiatiii		IIIg/IIIZ	□ IIIg/kg	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			
			1			

^{*}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:	Hospi	ital UPN:			Patient UIC		HSCT Date:	yyyy - mm - dd
Total Body Irradiation (TBI)		No		Yes	: Total prescribed radia			
					imber of fractions			
TLI, TNI, TAI (lymphoid, nodal, abdominal)		No		Yes	: Total prescribed radi			
					Survival Stat	TUS		
Survival Status on date o	f LICC	`T			- Carvivar Ctat			
Alive De		. I						
		istration	of th	e prep	parative regimen and date o	of HSCT		
Main Cause of Dea	•		-		ain cause):			
Relapse or Progro		/Persister	it dis	ease				
Unknown								
Other								
Contributor	y Cau	ise of De	eath	(0	check as many as approp	riate):		
☐ GVHD ☐ Interstiti	al nne	umonitis						
Pulmona								
Infection	:							
	terial							
□ vira								
☐ fun	gai asitic							
	nown							
Rejection			ctio	า				
					isorder (VOD)			
☐ Haemori	hage							
Cardiac t	oxicity	/						
☐ Central r	ervou	s system	(CNS) toxi	city			
Gastroin	testina	al (GI) tox	icity					
Skin toxi								
Renal fai								
☐ Multiple								
☐ Other, sp	ecity.		• • • • • • • • • • • • • • • • • • • •					