

EBMT Centre Identification Code (CIC): Hospital Unique Patient Number (UPN):	Treatment Type CT
Patient Number in EBMT Registry:	Treatment Date / / (YYYY/MM/DD)
CELLULAR THER	APY

CELLULAR THER	APY
Day 0	

PRE-INFUSION				
Cell collection procedure - Apheresis  Date of collection: / _ / _ (If more than one collection enter to the collection of collections:	_ (YYYY/MM/DD) the date of the <u>first</u> collection.)	☐ Date unknown (e.g. allogeneic product from unknown donor)		
	ATION FOR PLANNED CELL	ULAR THERAPY		
☐ Treatment of a primary disease:  Indication diagnosis for this of the indication diagnosis for the indication diagnosis for cellular therapy: (	osis has been registered first, usi	ing the relevant diagnosis form)		
☐ Induction therapy ☐ Prevention of disease relaps ☐ Rescue from disease relaps ☐ Minimal residual disease red ☐ Refractory disease ☐ Other; specify:	se or progression e or progression Juction			
☐ Treatment or prevention of complice (derived from a previous treatment or Date of the last treatment:	expected from a subsequent trea	atment)		
		has been registered and that relevant follow-up form unsplants and/or cellular therapies can be captured.		
Reason for cellular therapy: ☐ GvHD	☐ Treatment of GvHD☐ Preventive treatment for Gv	HD		
☐ Graft function	☐ Graft failure treatment ☐ Prevention of rejection/Pron ☐ Graft enhancement	notion of cell engraftment		
☐ Immune reconstitution				
Other indication; specify:				

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# BASIC INFORMATION ON THE PLANNED CELLULAR THERAPY

Clinical setting: (check only one)		
As per marketing approval / Standard of care / Institutional guidelines		
☐ Hospital exemption		
Compassionate use / Accelerated access		
Investigational drug product (IDP)/ Clinical trial	Phase:	
Cell origin:		
Autologous (Proceed to 'Planned cellular therapy infusion product(s)' section on page 3)		
☐ Allogeneic:		
This product is manufactured from:		
☐ A known donor never used to treat this patient (e.g. from a donor registry or related)		
(Proceed to 'Donor information' section on page 3.)		
A donor that is already registered as part of a previous treatment		
(Proceed to 'Planned cellular therapy infusion product(s)' section on page 3.)		
☐ An unknown donor with no data available (e.g. from a commercial product)  (Proceed to 'Planned cellular therapy infusion product(s)' section on page 3.)		

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DONOR INFORMATION  Complete only if cell source was allogeneic				
Did the donor consent to having their data in the EBM  ☐ No (complete only fields marked with '*' in this section ☐ Yes				
Date of birth: / (YYYY/MM/DD)	OR:	*Age at time of donation: years		
		If the donor was younger than 2 years:		
*Sex (at birth):		*Age in months:		
☐ Male				
☐ Female				
Donor Identification:				
Donor ID given by the treating centre (mandatory):	,			
Global registration identifier for donors (GRID):				
ION code of the Donor Registry or Cord Blood Ban	ık (mandatory	<i>)</i> :		
EuroCord code for the Cord Blood Bank (if applical	ble):			
Name of Donor Registry or Cord Blood Bank:				
<u>Donor</u> ID given by the Donor Registry or Cord Bloo	od Bank:			
Patient ID given by the Donor Registry or Cord Blo	od Bank:			
PLANNED CELLUI	LAR INFUSIO	ON PRODUCT(S)		
Will the planned cellular infusion product consist of n  □ No	nore than on	e infusion unit?		
Yes: Number of infusion units:				
Unknown				
Tissue source (check all that apply):				
Bone marrow				
Peripheral blood				
Cord blood				
☐ Tumour ☐ Other: specify:				
L L CILIEL, SUECIIV.				

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PLANNED CELLULAR INFUSION PRODUCT(S)
s the planned cell infusion product a commercial product?
□ No
☐ Yes
dentification:
Name of manufacturer:
☐ Autolus
Celgene/ Bristol-Myers Squibb
☐ Celyad
☐ GlaxoSmithKline (GSK)
☐ Janssen (Johnson & Johnson)
☐ Kite Gilead
☐ Miltenyi
☐ Novartis
☐ Local hospital or university
Other; specify:
Name of product:
Abecma
☐ Breyanzi
☐ Carvykti
☐ Kymriah
☐ Tecartus
☐ Yescarta
☐ No product name available
☐ Other: specify:

#### **END OF PRE-INFUSION SECTION**

PLEASE PROCEED WITH THE CELLULAR THERAPY SECTION TO COMPLETE

THE CELLULAR THERAPY DAY 0 REPORT

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# **CELLULAR THERAPY**

Date of (planned) cell infusion: / / (YYYY/MM/DD)				
Centre where infusion took place (CIC): (if the product was not infused, report the centre where the infusion was planned to take place)				
Patient UPN for this treatment:				
Team or unit where treatment took place (select all that apply):				
☐ Adults ☐ Pediatrics ☐ Haematology☐ Oncology ☐ Allograft ☐ Autograft ☐ Other; specify:				
Unit number:				
Was the cellular therapy product infused during this treatment/procedure?				
☐ No: Reason why the treatment did not take place: ☐ Production failure				
Select all reasons that apply  Out of specification product rejected by physician				
☐ Disease progression or patient condition worsening				
☐ Patient became ineligible for treatment				
☐ Patient died				
Other reason; specify:				
Yes: B-cell aplasia at time of cellular therapy?				
☐ Absent				
Present: Percentage of B-cells:%				
☐ Not evaluated				

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THERAPY & CELL INFUSION(S)					
Chronological number of cellular therapy treatment for this patient:  (Please do not include any transplants the patient has had in the past)					
Complete this section only if this is the second or a subsequent cellular therapy for this patient and the previous cellular treatments cannot be registered.					
If > 1:					
Same product as for the previous cellular therapy?  No Yes					
Date of the last cellular therapy before this one:I(YYYY/MM/DD)					
Type of the last cellular therapy before this one:  ☐ Autologous					
☐ Allogeneic: Was the same donor used both for prior and current cellular therapy? ☐ No					
☐ Yes  Was the last cellular therapy performed at another institution?					
No					
☐ Yes: CIC (if known):					
Name of institution:					
City:					
If > 1 submit an annual follow-up form before proceeding using the latest assessment date before this cellular therapy; this is so relapse data and other events between transplants/cellular therapies can be captured.					
Did the patient receive a previous HCT?					
□ No					
Yes: Date of the last HCT before this CT:/(YYYY/MM/DD)					
Type of the last HCT before this CT:  Autologous					
□ Allogeneic					

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☐ Yes

For same indication as the cellular therapy?  $\hfill \square$  No



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	_		_					_

	(before transplant/cellular the	rapy)
Was the par	tient treated before this cellular therapy procedure?	
☐ No (prod	ceed to 'Cellular therapy infusion unit(s)' on page 8)	
		1
☐ Yes	complete the "Treatment — non-HCT/CT/GT/IST" form	
_		
☐ Unknow	n (proceed to 'Cellular therapy infusion unit(s)' on page 8)	

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	Treatment Type	□ ст			
-	Treatment Date	1	1	(YYYY/MM/DD)	

CELLULAR THERAPY INFUSION UNIT(S)
Was there more than one cell infusion unit administered during this treatment?  No Yes: Number of different cell infusion units that were part of this treatment:
CELLULAR THERAPY INFUSION UNIT(S) DESCRIPTION
If the CT product was not infused proceed to 'Survival status' section on page 14.
If more than one cell infusion unit please copy and fill-in this section for each one of them.
Unique ID of the product: (If applicable)
Batch number: (If applicable)
Identification of the cell infusion unit given by the centre:
(If there is only one cell infusion unit enter "1")
Was the infused cellular product consistent with the specifications?
☐ No: specify the difference from specifications:
☐ Yes
Unknown
Was the cellular therapy product cryopreserved prior to infusion?
□ No □ Ves: Date of cryonreservation: / / (VVVV/MM/DD) □ Unknown
Tes. Date of dryopheservation.
☐ Unknown

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# CELLULAR THERAPY INFUSION UNIT(S) MANIPULATION

Complete <u>only for non-commercial products</u>. If more than one cell infusion unit please copy and fill-in this section for each one of them.

Identification of the cell in	fusion uni	t (given by	y the centre):	
Manipulation:				
Processing/Manufacturing ☐ Onsite, by local cell proc	_	lity		
☐ Offsite, by a non-comme	ercial facility	/		
Gene manipulation:  ☐ No ☐ Yes: <u>Type</u>				
Gene transfer:	<del>_</del>	Vector: [	Retroviral vector Lentiviral vector Other vector; specify:	
	Tra	nsgene: [	CAR; specify all targets:	See appendix 1 for a list of target antigens
			TCR; specify all targets:	
			specify HLA element:	
			Suicide gene; specify:	_
		С	Other: specify:	
Other:	□ No			

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Yes: specify:



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### **CELLULAR THERAPY INFUSION UNIT(S) MANIPULATION** continued

Complete only for non-commercial products. If more than one cell infusion unit please copy and fill-in this section for each one of them.

Manipulation aims:			
Recognition of a speci	fic target/antigen:		
Yes: <u>Type (check all</u>	that apply):		
│		☐ Funç	gal:
│	novirus	Г	↑ Candida
 ☐ BK V	'irus		- ↑ Aspergillus
☐ Covid	d-19 (SARS-CoV-2)		
☐ Cyto	megalovirus (CMV)	L	
☐ Epste	ein-Barr virus		
☐ Hum	an herpes virus 6		
☐ Hum	an immunodeficiency virus	HIV)	
☐ RSV-			
☐ Othe	r virus; specify:	<del></del>	
☐ Tumour/cancer	antigen(s); specify all:		
_	ecify:		
		<del></del>	
Cell types administere	<b>d</b> (check all that apply):		
CD3+ lymp			
CD4+ lymp			
CD8+ lymp	hocytes		
☐ CD34+			
☐ Dendritic ce			
☐ Gamma-De			
☐ Mesenchyr	nal cells		
☐ NK cells			
☐ Regulatory	T-cells		
☐ Other; spec	cify:	_	
Expansion:	Activation:	Induced differen	itiation:
☐ No	☐ No	☐ No	
☐ Yes	☐ Yes	☐ Yes	
Unknown	Unknown	☐ Unknown	
<u> </u>			



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# PREPARATIVE REGIMEN

Do not include lines of therapy given for disease treatment, bridging therapy or maintenance, these should be reported in other form.
Preparative conditioning / lymphodepletion regimen given?
□ No
Yes: <b>Drugs given?</b> (any active agent, including chemotherapy, monoclonal antibody, polyclonal antibody, serotherapy, etc.)
□ No
Yes (provide details in the table on pages 12-13)
Unknown

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#### **PREPARATIVE REGIMEN continued**

# Specification and dose of the preparative regimen:

(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.

Report dosages and units only for individual drugs.)

Chemotherapy		Dose	Units
☐ Alemtuzumab			☐ mg/m² ☐ mg/kg
☐ Anti-Thymocyte Globulin Anti-Lympho	cyte Globulin		☐ mg/m² ☐ mg/kg
Product name:			
Origin:			
☐ Bendamustine			☐ mg/m² ☐ mg/kg
Bleomycin			☐ mg/m² ☐ mg/kg
☐ Busulfan			
	ral / oth		☐ mg/m² ☐ mg/kg
Drug monitoring performed:   N  Ye	o es; total AUC:		
	☐ mg x hr/L ☐ micromol x min/L ☐ mg x min/mL		
☐ Carboplatin			
Drug monitoring performed: 🔲 N	lo		☐ mg/m² ☐ mg/kg
□ Y	'es; total AUC: ☐ mg x hr/L ☐ micromol x min/L ☐ mg x min/mL		
Carmustine			☐ mg/m² ☐ mg/kg
☐ Cisplatin			☐ mg/m² ☐ mg/kg
☐ Clofarabine			☐ mg/m² ☐ mg/kg
Corticosteroids:			
Beclometasone			☐ mg/m² ☐ mg/kg
Budesonide			☐ mg/m² ☐ mg/kg
Dexamethasone			☐ mg/m² ☐ mg/kg
Methylprednisolone			☐ mg/m² ☐ mg/kg
☐ Prednisolone			☐ mg/m² ☐ mg/kg
☐ Cyclophosphamide			☐ mg/m² ☐ mg/kg



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#### PREPARATIVE REGIMEN continued

# Specification and dose of the preparative regimen:

(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.)

Chemotherapy	Dose	Unit
☐ Cytarabine		☐ mg/m² ☐ mg/kg
☐ Daunorubicin		☐ mg/m² ☐ mg/kg
☐ Doxorubicin		☐ mg/m² ☐ mg/kg
☐ Epirubicin		☐ mg/m² ☐ mg/kg
☐ Etoposide		☐ mg/m² ☐ mg/kg
Fludarabine		☐ mg/m² ☐ mg/kg
☐ Gemtuzumab ozogamicin		☐ mg/m² ☐ mg/kg
☐ Ibritumomab tiuxetan		☐ mCi ☐ MBq
☐ Idarubicin		☐ mg/m² ☐ mg/kg
☐ Ifosfamide		☐ mg/m² ☐ mg/kg
☐ Imatinib		☐ mg/m² ☐ mg/kg
Lomustine		☐ mg/m² ☐ mg/kg
☐ Melphalan		☐ mg/m² ☐ mg/kg
Mitoxantrone		☐ mg/m² ☐ mg/kg
☐ Paclitaxel		☐ mg/m² ☐ mg/kg
☐ Anti-CD20 antibodies		☐ mg/m² ☐ mg/kg
☐ Teniposide		☐ mg/m² ☐ mg/kg
☐ Thiotepa		☐ mg/m² ☐ mg/kg
☐ Tositumomab		☐ mCi ☐ MBq
☐ Treosulfan		☐ mg/m² ☐ mg/kg
Other; specify*:		☐ mg/m² ☐ mg/kg
		☐ mCi ☐ MBq
*Please consult the LIST OF CHEMOTHERAPY DRUGS/AGENTS AND names	REGIMENS on the EBM	E website for drugs/regimen
Total body irradiation (TBI):		
□ No		
Yes; total prescribed radiation dose as per protocol:	Gy	
number of fractions:		
number of radiation days:		

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Cell viability: \_\_\_\_\_\_ %

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CELL INFUSION EPISODE(S)			
Was there more than one cell infusion episodes during this treatment or procedure?  □ No			
Yes: Number of cell infusion episodes during this treatment/procedure:			
CELL INFUSION EPISODE(S) DESCRIPTION			
If more than one cell infusion unit please copy and fill-in this section for each one of them.			
Date of cell infusion episode:/_/_(YYYY/MM/DD)			
Route of infusion: (check all that apply)			
Intraveneous Intraveneous			
Intrathecal			
Intratumour injection			
Other route; specify:			
Did the patient receive concomitant therapy?			
□ No			
Yes; specify:			
Treatment given: Simultaneously to the cellular therapy			
☐ After the cellular therapy episode was finished			
If more than one unit was used, indicate the identification of the cell infusion given by the centre as described in the 'Cell Infusion Unit' section (This item is mandatory if more than one cell infusion unit was used.):			
Is the exact number of cells infused available?			
□ No			
☐ Yes: Number of cells: Unit (check only one): ☐ 10 <sup>6</sup> /kg ☐ 10 <sup>6</sup> ☐ 10 <sup>8</sup> /kg ☐ 10 <sup>8</sup> (not adjusted for cell viability)			

Treatment Type 

CT

END OF THE CELLULAR THERAPY DAY 0 REPORT proceed to form DISEASE STATUS AT HCT/CT/GT/IST

If more than one cell infusion unit was administered please copy and fill-in this section for each one of them.

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# Appendix 1 -- List of transgene CAR targets --

AFP (alpha fetoprotein) BAFF-R

 $\mathsf{BCMA}$ 

B7H3

CD11 CD16

CD19

CD20 CD22 CD30

CD33 CD38

CD56

CD123 CD138

CD171 CD229

CD229 CLL1 CS-1 (SLAMF7) EGFR GD2 GPRC5D HER2 HPV-16E6

Integrinβ7

Lewis-Y

MAGE-A4

MAGE-A10 Mesothelin (MSLN)

MUC16 NKG2D

NY-ESO-1

PRAME

PSCA SSX

Survivin TACI

WT-1

Other (specify)