

EBMT Centre Identification Code (CIC):	Treatment Type	☐ HCT
Hospital Unique Patient Number (UPN):		
Patient Number in EBMT Registry:	Treatment Date _	II (YYYY/MM/DD)

HAEMATOPOIETIC CELL TRANSPLANTATION (HCT) --- Day 100 Follow-Up ---

SURVIVAL STATUS					
Date of follow-up:/_/_(YYYY/MM/DD) (if died: date of death, if lost to follow up: date last seen)					
Survival status: Alive Dead Lost to follow-up Main cause of death: (check only one main cause)					
Relapse or progression/persistent disease					
Secondary malignancy					
☐ CT-related	Select treatment related cause: (select all that apply) Graft versus Host Disease Non-infectious complication Infectious complication:				
☐ HCT-related	(select all that apply) Bacterial infection				
☐ GT-related	☐ Viral infection☐ Fungal infection☐				
☐ IST-related	Parasitic infection Infection with unknown pathogen				
☐ Unknown					
Other; specify:					
Autopsy performed: No Yes Unknown					
BEST RESPONSE Not applicable for Inborn Errors					
Best clinical/biological response after HCT* (observed before any subsequent treatment):					

Unknown

Date best response first observed: _ _ _ / _ _ (YYYY/MM/DD)

^{*} Indicate the best clinical/biological response after HCT corresponding to indication diagnosis by selecting from the list provided in Appendix 1



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RECOVERY

Absolute neutrophil count (ANC) recovery (neutrophils ≥ 0.5x10 ⁹ /L):
☐ No (Primary graft failure): Date of the last assessment: / / (YYYY/MM/DD) ☐ Unknown
 Yes: Date of ANC recovery: / / (YYYY/MM/DD) ☐ Unknown (first of 3 consecutive values after 7 days without transfusion containing neutrophils) ☐ Never below ☐ Unknown
Platelet reconstitution (platelets ≥ 20×10 ⁹ /L:):
☐ No: Date of the last assessment: / _ / _ (YYYY/MM/DD) ☐ Unknown
Yes: Date of platelet reconstitution: / / (YYYY/MM/DD) Unknown (first of 3 consecutive values after 7 days without platelet transfusion)
☐ Never below
☐ Unknown
Date of the last platelet transfusion: / (YYYY/MM/DD)

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Poor graft function (defined as: frequent dependence on blood and/or platelet transfusions and/or growth factor support in the absense of other explanations, such as disease relapse, drugs, or infection): No Yes; Date of poor graft function://(YYYY/MM/DD) Unknown Unknown Complete for every chimaerism test performed: (complete only if patient received an allogeneic HCT)
Chimaerism test date: / / (YYYY/MM/DD)
Source of cells tested: Peripheral blood
☐ Bone marrow
Select cell type and complete relevant test results: Global: % donor
Myeloid cells (i.e. CD33, CD15 or CD14):% donor ☐ Unknown☐ T-cells (CD3):% donor ☐ Unknown
B-cells (CD19 or CD20):% donor Unknown
CD34+ cells:% donor Unknown
Other cell type; specify cells; donor Unknown
copy and fill-in this table as many times as necessary.
PREVENTIVE THERAPIES (Complete only if the patient received an alloHCT)
Immunosuppression: No Yes; Immunosuppresion stopped: No Yes; End date:/(YYYY/MM/DD) Unknown Unknown
☐ Unknown
Letermovir used as CMV prophylaxis: No Yes; Start date: / _ / _ (YYYY/MM/DD) Unknown
Letermovir treatment stop?
☐ Unknown

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Unknown

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COMPLICATIONS POST HCT TREATMENT

-- GvHD --

Allogeneic HCT only

Did gr	aft versus host dis	ease (GvHD) occur?					
□N	o (proceed to 'Comp	lications since the last report - Non-infectious complications')					
☐ Y	 Yes: Did the patient receive a systemic/immunosuppressive treatment for GvHD? No Yes: Date treatment started:// (YYYY/MM/DD) ☐ Unknown 						
	Treatment stopped: No Yes; Stop date of treatment:// (YYYY/MM/DD) Unknown Unknown						
	☐ Unknown						
	Inknown (proceed to	'Complications since the last report - Non-infectious complications')					
Did a	cute GvHD occur d	uring this follow-up period?					
□и	0						
☐ Ye	es: Date of onse	t: / / (<i>YYYY/MM/DD</i>)					
	Maximum ohse	rved organ severity score:					
[Skin:	☐ 0 (none) ☐ 1 ☐ 2 ☐ 3 ☐ 4 ☐ Not evaluated ☐ Unknown					
	Liver:	☐ 0 (none) ☐ 1 ☐ 2 ☐ 3 ☐ 4 ☐ Not evaluated ☐ Unknown					
	Lower GI tract:	☐ 0 (none) ☐ 1 ☐ 2 ☐ 3 ☐ 4 ☐ Not evaluated ☐ Unknown					
	Upper GI tract:	☐ 0 (none) ☐ 1 ☐ Not evaluated ☐ Unknown					
	Other site affected:	☐ No ☐ Yes; specify:					
	Overall maximum (grade observed: 1 2 3 4 Unknown Not evaluated					
Steroid-refractory acute GvHD: No Yes: Date of onset:/(YYYY/MM/DD) Unknown							
	aGvHD resolved:	☐ Unknown☐ No☐ Yes; Date of aGvHD resolution://(YYYY/MM/DD)☐ Unknown☐ Unknown					

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-- GvHD --

	Allogeneic HCT only							
Did chro	Did chronic GvHD occur during this follow-up period?							
☐ No								
☐ Yes:	Date of onset:	_//(YY	YY/MM/D	DD) Unkno	own			
	Maximum NIH score: Mild Moderate Severe Unknown Not evaluated							
	Date of maximum N Maximum observed			(YYYY/MM/	<i>′DD)</i> ∏ Unkr	nown		
	Skin:	0 (none)		П 2	□ 3	☐ Not evaluared	☐ Unknown	
	Oral:	0 (none)		☐ 2	3	☐ Not evaluated	 ☐ Unknown	
	Gastrointestinal:	□ 0 (none)	□ 1	□ 2	□ 3	☐ Not evaluated	 ☐ Unknown	
	Eyes:	□ 0 (none)	<u> </u>	□ 2	□ 3	☐ Not evaluated	☐ Unknown	
	Liver:	☐ 0 (none)	□ 1	□ 2	□ 3	☐ Not evaluated	☐ Unknown	
	Joints and fascia:	□ 0 (none)	<u> </u>	□ 2	□ 3	☐ Not evaluated	☐ Unknown	
	Lungs:	☐ 0 (none)	□ 1	□ 2	□ 3	☐ Not evaluated	☐ Unknown	
	Genitalia:	☐ 0 (none)	□ 1	□ 2	□ 3	☐ Not evaluated	☐ Unknown	
	Other site affected:	☐ No	☐ Yes;	specify:				
Steroid-refractory chronic GvHD: No Yes: Date of onset://(YYYY/MM/DD) Unknown Unknown								
C	GvHD resolved:	No						
	☐ Yes; Date of cGvHD resolution: / (YYYY/MM/DD) ☐ Unknown							
	□ Unknown							

☐ No ☐ Yes ☐ Unknown

Was overlap syndrome observed: (features of both chronic and acute GvHD)

☐ Unknown

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COMPLICATIONS SINCE THE LAST REPORT

-- Non-infectious complications --

Did non-infectious complications occur during the follow-up period? (Please only report toxic events here that are above Grade 2 and not linked to GvHD and/or infections)
No (proceed to 'Complications since the last report - Infectious complications')
Yes (report in the table below)
Secondary graft failure
Complication observed? No
☐ Yes
☐ Unknown
Maximum grade observed during this period: Non-fatal Fatal
Onset date (YYYY/MM/DD):/ _ Unknown
Resolved: No
Yes; Stop date (YYYY/MM/DD):/ _ Unknown
☐ Unknown
Cardiac event
Complication observed? No*
☐ Yes:
Unknown
Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown
Onset date (YYYY/MM/DD):/ _ Unknown Resolved: No
☐ Yes; Stop date (<i>YYYY/MM/DD</i>): / ☐ Unknown
☐ Unknown
Central nervous system (CNS) toxicity
Complication observed?
Ţes:
Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown
Onset date (YYYY/MM/DD): / Unknown
Resolved: No
☐ Yes; Stop date (<i>YYYY/MM/DD</i>): / ☐ Unknown
☐ Unknown
Gastrointestinal (GI) Toxicity (non-GvHD and non-infectious related)
Complication observed? No*
☐ Yes:
☐ Unknown
Maximum CTCAE grade observed: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ _ Unknown
Resolved: No
Yes; Stop date (YYYY/MM/DD):/ _ Unknown
☐ Unknown

* Grade 0-2



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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COMPLICATIONS SINCE THE LAST REPORT Non-infectious complications continued				
Liver disorder Complication observed? No* Yes: Unknown				
Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown				
Onset date (YYYY/MM/DD):/				
☐ Yes; Stop date (YYYY/MM/DD):/ _ ☐ Unknown ☐ Unknown				
Renal failure (chronic kidney disease, acute kidney injury)				
Complication observed? No* Yes: Unknown				
Maximum CTCAE grade observed: 3 5 (fatal) Unknown				
Onset date (YYYY/MM/DD):/				
☐ Yes; Stop date (YYYY/MM/DD): / ☐ Unknown ☐ Unknown				
Respiratory disorders				
Complication observed? No* Yes: Unknown				
Maximum CTCAE grade observed: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown				
Onset date (YYYY/MM/DD):/				
Yes; Stop date (YYYY/MM/DD):/ Unknown				
☐ Unknown				
Skin Toxicity (non-GvHD and non-infectious related)				
Complication observed? No*				
☐ Yes:				
Unknown				
Maximum CTCAE grade observed: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown				
Onset date (YYYY/MM/DD):/				
☐ Yes; Stop date (YYYY/MM/DD):/ ☐ Unknown ☐ Unknown				

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^{*} Grade 0-2



EBMT Centre Identification Code (CIC):	Treatment Type	☐ HCT	
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Vascular event
Complication observed? No*
☐ Yes:
☐ Unknown
Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown
Onset date (YYYY/MM/DD):/ _ Unknown
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / ☐ Unknown
☐ Unknown
Avascular necrosis (AVN)
Complication observed? No*
☐ Yes:
☐ Unknown
Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown
Onset date (YYYY/MM/DD):/ _ Unknown
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / ☐ Unknown
☐ Unknown
Cerebral haemorrhage
Cerebral haemorrhage Complication observed? No*
•
Complication observed? No*
Complication observed? No*
Complication observed? No* Yes: Unknown
Complication observed? No* Yes: Unknown Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown
Complication observed? No* Yes: Unknown Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown Onset date (YYYY/MM/DD):/ Unknown
Complication observed? No* Yes: Unknown Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown Onset date (YYYY/MM/DD):/ Unknown Resolved: No
Complication observed?
Complication observed?
Complication observed?
Complication observed? No* Yes: Unknown Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown Onset date (YYYY/MM/DD):/ Unknown Resolved: No Yes; Stop date (YYYY/MM/DD):/ Unknown Unknown Haemorrhage (other than cerebral haemorrhage) Complication observed? No*
Complication observed? No* Yes: Unknown Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown Onset date (YYYY/MM/DD):// Unknown Resolved: No Yes; Stop date (YYYY/MM/DD):// Unknown Unknown Haemorrhage (other than cerebral haemorrhage) Complication observed? No* Yes:
Complication observed?
Complication observed? No* Yes: Unknown Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown Onset date (YYYY/MM/DD):/ Unknown Resolved: No Yes; Stop date (YYYY/MM/DD):/ Unknown Unknown Haemorrhage (other than cerebral haemorrhage) Complication observed? No* Yes: Unknown Unknown Maximum CTCAE grade observed: 3 4 5 (fatal) Unknown
Complication observed?



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Cerebral thrombosis
Complication observed? No*
☐ Yes:
☐ Unknown
Maximum CTCAE grade observed: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ _ Unknown
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD):/ ☐ Unknown
☐ Unknown
Cytokine release syndrome (CRS)
Complication observed? No*
☐ Yes:
☐ Unknown
Maximum CTCAE grade observed: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ _ Unknown
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / ☐ Unknown
☐ Unknown
Haemophagocytic lymphohistiocytosis (HLH)
Complication observed? No*
☐ Yes:
☐ Unknown
Maximum CTCAE grade observed: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ Unknown
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / ☐ Unknown
☐ Unknown
Pure red cell aplasia (PRCA)
Complication observed? No
Yes:
☐ Unknown
 Maximum grade observed:
Onset date (YYYY/MM/DD): /
Resolved: No
☐ Unknown
_

^{*} Grade 0-2



Resolved: No

☐ Unknown

EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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COMPLICATIONS SINCE THE LAST REPORT

Non-infectious complications
continued
Posterior reversible encephalopathy syndrome (PRES)
Complication observed? No
☐ Yes:
☐ Unknown
Maximum grade observed: ☐ Non-severe ☐ Severe ☐ Fatal ☐ Unknown
Onset date (<i>YYYY/MM/DD</i>): /
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD):/ _ ☐ Unknown
☐ Unknown
Transplant-associated microangiopathy (TMA)
Complication observed? No*
☐ Yes:
☐ Unknown
Maximum grade observed: Non-severe Severe Unknown
Onset date (<i>YYYY/MM/DD</i>): /

Yes; Stop date (YYYY/MM/DD): ____/ _ Unknown

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Resolved: No

☐ Unknown

EBMT	EBMT Centre Identification Code (CIC Hospital Unique Patient Number (UPN Patient Number in EBMT Registry:):	reatment Type	YY/MM/DD)		
COMPLICATIONS SINCE THE LAST REPORT Non-infectious complications						
Veno-occlusive disease (VOD)						
Complication	n observed? No* Yes	Unknown				
Maximum CT	CAE grade observed Mild	☐ Moderate ☐ Severe	☐ Very severe ☐ Fatal	Unknown		
Onset date ()	/YYY/MM/DD):] Unknown				

 \square Yes; Stop date (YYYY/MM/DD): ____/ \square Unknown

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EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
Hospital Unique Patient Number (UPN):			
Patient Number in EBMT Registry:	Treatment Date _	//(YYYY/MM/DD)

COMPLICATIONS SINCE THE LAST REPORT Non-infectious complications		
Other complication observed? No* Yes Unknown		
Specify: Consult appendix 4 for a list of complications that should not be reported		
(Indicate CTCAE term)		
Maximum CTCAE grade observed 3 4 5 (fatal) Unknown		
Onset date (YYYY/MM/DD): / Unknown		
Resolved: No		
Yes; Stop date (YYYY/MM/DD):/ Unknown		

If more other complications occurred, copy and fill-in this table as many times as necessary.

* Grade 0-2

☐ Unknown

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EBMT Centre Identification Code (CIC):	Treatment Type	□ HCT	
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Infectious complications
Do not report infections that were already reported as resolved on the previous assessment and did not reoccur. Did infectious complications occur during the follow-up period? No Consult appendix 4 for a list of complications that should not be reported Yes (report all infection-related complications below)
Bacterial infection: No Yes
1) Start date: / / (YYYY/MM/DD)
Gram-positive Gram-negative Other Pathogen*:
Infection with clinical implications: No Yes: (select all that apply during this period)
Symptoms/signs of disease
Administration of pathogen-directed therapy Unknown Indicate at least 1 location involved during this period:
Localisation 1 (CTCAE term)**:
Localisation 2 (CTCAE term)**:
Localisation 3 (CTCAE term)**:
Yes; specify***:
☐ Unknown
Resolved: No Yes Unknown
(if patient died) Contributory cause of death: No Yes Unknown
2) Start date://(YYYY/MM/DD) Gram-positive Gram-negative Other Pathogen*:
Infection with clinical implications: No Yes: (select all that apply during this period) Symptoms/signs of disease
☐ Administration of pathogen-directed therapy
Unknown Indicate at least 1 location involved during this period: Localisation 1 (CTCAE term)**:
Localisation 2 (CTCAE term)**:
Localisation 3 (CTCAE term)**:
Intravascular catheter-related infection No Yes; specify***:
Unknown Resolved: No Yes Unknown
(if patient died) Contributory cause of death: No Yes Unknown
If more than 2 bacterial infections, copy and fill-in this table as many times as necessary.

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^{*} Indicate the pathogen and sub-type (if applicable) by choosing from the list of pathogens provided in Appendix 2
** Indicate CTCAE term by choosing from the list provided in Appendix 3
*** If intravascular catheter-related infection, specify it by choosing from the list provided in Appendix 5



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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Viral infection: No Yes	
1) Start date: / / (YYYY/M	M/DD)
If the pathogen was CMV/EBV: Was th	is infection a reactivation? No
Infection with clinical implications:	☐ No ☐ Yes: (select all that apply during this period) ☐ Symptoms/signs of disease
	☐ Administration of pathogen-directed therapy ☐ Unknown
Indicate at least 1 location involved during Localisation 1 (CTCAE term)**:	
Localisation 2 (CTCAE term)**:	
Localisation 3 (CTCAE term)**:	
Resolved: No Yes	☐ Unknown
(if patient died) Contributory cause of death: □ N	Jo ☐ Yes ☐ Unknown
2) Start date : / / (YYYY/M	M/DD)
Pathogen*:	
If the pathogen was CMV/EBV: Was ti	nis infection a reactivation?
Infection with clinical implications:	☐ No ☐ Yes: (select all that apply during this period) ☐ Symptoms/signs of disease
	Administration of pathogen-directed therapy
Indicate at least 1 location involved during Localisation 1 (CTCAE term)**:	Unknown g this period:
Localisation 2 (CTCAE term)**:	
Localisation 3 (CTCAE term)**:	
Resolved: No Yes	Unknown
(if patient died) Contributory cause of death:	No ☐ Yes ☐ Unknown
	ctions, copy and fill-in this table as many times as necessary.

 $^{^{\}star}$ Indicate the pathogen and sub-type (if applicable) by choosing from the list of pathogens provided in Appendix 2

^{**} Indicate CTCAE term by choosing from the list provided in Appendix 3 $\,$

^{***} If intravascular catheter-related infection, specify it by choosing from the list provided in Appendix 5



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Fungal infection: No Yes
1) Start date://(YYYY/MM/DD) Yeasts
Infection with clinical implications: No Yes: (select all that apply during this period) Symptoms/signs of disease
☐ Administration of pathogen-directed therapy ☐ Unknown Indicate at least 1 location involved during this period: Localisation 1 (CTCAE term)**:
Localisation 2 (CTCAE term)**:
Localisation 3 (CTCAE term)**:
Intravascular catheter-related infection No Yes; specify***:
Unknown Resolved: No Yes Unknown (if patient died) Contributory cause of death: No Yes Unknown
2) Start date://(YYYY/MM/DD) Yeasts Moulds Pathogen*:
Infection with clinical implications: \square No
Yes: (select all that apply during this period)
Symptoms/signs or disease
☐ Administration of pathogen-directed therapy ☐ Unknown
Indicate at least 1 location involved during this period:
Localisation 1 (CTCAE term)**:
Localisation 2 (CTCAE term)**:
Localisation 3 (CTCAE term)**:
Intravascular catheter-related infection: No Yes; specify***:
☐ Unknown
Resolved: No Yes Unknown (if patient died) Contributory cause of death: No Yes Unknown
If more than 2 fungal infections, copy and fill-in this table as many times as necessary.
* Indicate the pathogen and sub-type (if applicable) by choosing from the list of pathogens provided in Appendix 2

^{**} Indicate CTCAE term by choosing from the list provided in Appendix 3 $\,$

^{***} If intravascular catheter-related infection, specify it by choosing from the list provided in Appendix 5 HCT_FU_D100_v2.3



EBMT Centre Identification Code (CIC):	Treatment Type	☐ HCT	
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Parasitic infection: ☐ No ☐ Yes
Parasitic infection:
1) Start date:/ (YYYY/MM/DD)
Protozoa Helminths Pathogen*:
Infection with clinical implications:
☐ Yes: <i>(select all that apply during this period)</i> ☐ Symptoms/signs or disease
Administration of pathogen-directed therapy
Unknown Indicate at least 1 location involved during this period:
Localisation 1 (CTCAE term)**:
Localisation 2 (CTCAE term)**:
Localisation 3 (CTCAE term)**:
Resolved: No Yes Unknown
Tresolved. — Tres — Officiowii
(if patient died)
Contributory cause of death: No Yes Unknown
2) Start date: / / (YYYY/MM/DD) Protozoa Helminths
Pathogen*:
Infection with clinical implications: No
Yes: (select all that apply during this period)
Symptoms/signs or disease
Administration of pathogen-directed therapy
□ Unknown
Indicate at least 1 location involved during this period:
Localisation 1 (CTCAE term)**:
Localisation 3 (CTCAE term)**:
Resolved: No Yes Unknown
(if patient died)
Contributory cause of death: No Yes Unknown
If many than 2 manathis infactions are said fill in this table as the said and the
If more than 2 parasitic infections, copy and fill-in this table as many times as necessary.
* Indicate the pathogen and sub-type (if applicable) by choosing from the list of pathogens provided in Appendix 2

^{**} Indicate CTCAE term by choosing from the list provided in Appendix 3

^{***} If intravascular catheter-related infection, specify it by choosing from the list provided in Appendix 5 $\,$



EBMT Centre Identification Code (CIC):	Treatment Type	☐ HCT	
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	al documentation, like pneumonia, cellulitis, etc.)
1) Start date://(YYYY) Infection with clinical implications:	□ No □ Yes: (select all that apply)
	Symptoms/signs or disease
	Administration of pathogen-directed therapy
	Unknown
Indicate at least 1 location: Localisation 1 (CTCAE term)*:	
Localisation 2 (CTCAE term)*:	
Localisation 3 (CTCAE term)*:	
Intravascular catheter-related infect	tion: No
mitavassalai sametei relatea miest	Yes; specify**:
	☐ Unknown
Resolved: No Yes	Unknown
(if patient died) Contributory cause of death: No	o 🔲 Yes 🔲 Unknown
2) Start date : / / (YYYY/	/MM/DD)
Infection with clinical implications:	□ No
	Yes: (select all that apply)
	Symptoms/signs or disease
	☐ Administration of pathogen-directed therapy ☐ Unknown
Indicate at least 1 location:	CHKHOWH
Localisation 1 (CTCAE term)*:	
Localisation 2 (CTCAE term)*:	
Localisation 3 (CTCAE term)*:	
Intravascular catheter-related infecti	ion: No
	Yes; specify**:
	Unknown
Resolved: No Yes [☐ Unknown
Resolved: No Yes [(if patient died) Contributory cause of death: No	

Indicate CTCAE term by choosing from the list provided in Appendix 3 at page 25

^{**} If intravascular catheter-related infection, specify it by choosing from the list provided in Appendix 5 at page 25



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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Patient Number in EBMT Registry:	Treatment Date	//	(YYYY/MM/DD)

SECONDARY MALIGNANCIES AND AUTOIMMUNE DISORDERS

Did a secondary malignancy or autoimmune disorder occur after HCT? ☐ No
☐ Yes; Was this disease an indication for a subsequent HCT/CT/IST/GT?
☐ No (complete the non-indication diagnosis form)
Yes (complete the relevant indication diagnosis form)
☐ Unknown



☐ Yes

EBMT Centre Identification Code (CIC): $___$

Hospital Unique Patient Number (UPN): _____

	Patient Number in EBMT Registry:	Treatment Date / / (YYYY/MM/DD)
	ADDITIONAL TREATM	ENTS
Did the p	patient receive any additional disease treatment?	
☐ Yes:	complete the "Treatment — non-HCT/CT/GT/IST" form	
☐ Unkn	own	
	ADDITIONAL CELL INFU	JSIONS
-	patient receive additional cell infusions during this period? g a new HCT and CT)	
☐ Yes;	Is this cell infusion an allogeneic boost*? 🔲 No	☐ Yes
	* An allogeneic boost is an infusion of cells from the same don graft rejection.	or without conditioning, with no evidence of
	Date of the allogeneic boost: / _ / _ (YYYY//	MM/DD)
	Is this cell infusion an autologous boost?	☐ Yes
	Date of the autologous boost://(YYYY/	/MM/DD)
	nfusion is not a boost, attach the Cell Infusion (CI) sheet availab episodes of cell infusion that took place during this interval; then	
Did the pa □ No	tient receive subsequent HCT/CT (either at your or another ce	entre)?

Treatment Type HCT

If the patient had a subsequent HCT/CT, please, make sure that this subsequent treatment is registered using the appropriate treatment form before proceeding.

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RELAPSE, PROGRESSION, RECURRENCE OF DISEASE OR SIGNIFICANT WORSENING

(not relevant for Inborn errors)

e a relapse, progression	· · · · · · · · · · · · · · · · · · ·			
disease after HCT? (dete			se or significant worsening of organ function related	to the
; for every relapse, progr	ession, recu	urrence, sigi	nificant worsening complete the questions below	
Type: ☐ Relapse / Re	currence of	f disease		
(Continuous)	progressio	n / Significa	nt worsening	
Date of relapse/progre	ession/recu	ırrence/wor	rsening: / / (YYYY/MM/DD)	า
Malignant disorders o	nly:			
Malignant disorders of Type of relapse/pro				
		☐ Yes	☐ Unknown	
Type of relapse/pro	ogression:		☐ Unknown	
Type of relapse/pro Medullary: Extramedullary:	ogression:	☐ Yes	_	
Type of relapse/pro Medullary: Extramedullary:	ogression: No No ression was	☐ Yes☐ Yes☐ Yes	☐ Unknown [lary or both medullary and extramedullary:	
Type of relapse/pro Medullary: Extramedullary: If the relapse/progr Involvement at tim Skin:	ogression: No No ression was	☐ Yes☐ Yes☐ Yes	☐ Unknown [lary or both medullary and extramedullary:	
Type of relapse/pro Medullary: Extramedullary: If the relapse/progr Involvement at tim Skin: CNS:	ogression: No No No ression was	☐ Yes☐ Yes☐ Yes☐ extramedulese/progress	☐ Unknown Unknown Un	
Type of relapse/pro Medullary: Extramedullary: If the relapse/progr Involvement at tim Skin:	ogression: No No ression was e of relaps	☐ Yes☐ Yes ☐ Yes ☐ extramedula ☐ Yes☐ Yes☐ Yes☐ Yes☐ Yes☐ Yes☐ Yes☐ Yes	☐ Unknown Unknown	

copy and fill-in this table as many times as necessary.



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Disease status after HCT or at time of death*:

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^{*} Indicate the disease status at this follow-up or at time of death corresponding to indication diagnosis by selecting from the list provided in Appendix 1



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Appendix 1 Best Response and Disease Status (Disease Specific)

Complete only one section with the main indication diagnosis for which HCT was given.

ACUTE LEUKAEMIAS	Go to page 23
CHRONIC LEUKAEMIAS	Go to page 23
PLASMA CELL NEOPLASMS (PCN)	Go to page 24
MPN, MDS, MDS / MPN OVERLAP SYNDROMES	Go to page 26
LYMPHOMAS	Go to page 27
SOLID TUMOURS	Go to page 27
BONE MARROW FAILURE SYNDROMES (BMF) including APLASTIC ANAEMIA (AA)	Go to page 27
AUTOIMMUNE DISORDERS	Go to page 28
HAEMOGLOBINOPATHIES	Go to page 28
OTHER DIAGNOSIS	Go to page 29



☐ Not in complete remission

□ Not evaluated

	Appendix 1	
	_ Patient Number in EBMT Registry:	Treatment Date / / (YYYY/MM/DD)
EBMT	EBMT Centre Identification Code (CIC): Hospital Unique Patient Number (UPN):	Treatment Type HCT

Best Response and Disease Status (Disease Specific) Acute leukaemias (AML, PLN, Other) ☐ Complete remission (CR)

Proceed to next page for Diseases Status section					
Chronic leukaemias (CML, CLL, PLL, Other)					
Chronic Myeloid Leukaemia (CML):	_				
\square Chronic phase (CP); Number : \square 1 st \square 2 nd \square 3 rd or higher \square Unknown					
Haematological remission: ☐ No ☐ Yes ☐ Not evaluated ☐ Unknown					
Cytogenetic remission: 🔲 No 🔲 Yes 🔲 Not evaluated 🔲 Unknown					
Molecular remission: □ No □ Yes □ Not evaluated □ Unknown					
\square Accelerated phase; Number : \square 1 st \square 2 nd \square 3 rd or higher \square Unknown					
☐ Blast crisis; Number : ☐ 1 st ☐ 2 nd ☐ 3 rd or higher ☐ Unknown					
☐ Not evaluated	\dashv				
	\dashv				

Proceed to next page for Diseases Status section



☐ Unknown

EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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Appendix 1 Best Response and Disease Status (Disease Specific)

Best Response and Disease Status (Disease Specific) Chronic Lymphocytic Leukaemia (CLL), Prolymphocytic Leukaemia (PLL) and other chronic leukaemias: ☐ Complete remission (CR) ☐ Partial remission (PR) Sensitive to last regimen ☐ Unknown Progression: Resistant to last regimen ☐ Stable disease (no change, no response/loss of response) ☐ Relapse □ Not evaluated Unknown Proceed to next page for Diseases Status section Plasma cell neoplasms (PCN) ☐ Complete remission (CR) Number: 1st ☐ Stringent complete remission (sCR) ☐ 2nd ☐ Very good partial remission (VGPR) ☐ 3rd or higher ☐ Partial remission (PR) Unknown ☐ Relapse ☐ Progression ☐ Stable disease (no change, no response/loss of response) ☐ Not evaluated

Proceed to next page for Diseases Status section



☐ Unknown

EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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Appendix 1 Best Response and Disease Status (Disease Specific) continued

Complete only for PCN Disease Status Was the patient on dialysis after HCT? ☐ No ☐ Yes; Start date: _ _ _ / _ _ (YYYY/MM/DD) ☐ Unknown Did dialysis stop? ☐ No ☐ Yes; ☐ Unknown ☐ Unknown Complete only for leukaemias (AL, CLL) and PCN Disease Status Leukaemias (AL, CLL) and PCN (complete only for patient in CR or sCR) Minimal residual disease (MRD): □ Negative ☐ Positive; ☐ Increasing (>1log10 change) ☐ Stable (<1log10 change) ☐ Decreasing (>1log10 change) ☐ Unknown □ Not evaluated ☐ Unknown Date MRD status evaluated: _ _ _ / _ _ (YYYY/MM/DD) ☐ Unknown Sensitivity of MRD assay: Method used: **10**-6 (select all that apply) ☐ PCR _ ≤10-4 ☐ Flow cytometry **□** ≤10⁻³ ☐ NGS Other; specify: _ ☐ Other; specify:

Unknown



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст		
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Appendix 1 Best Response and Disease Status (Disease Specific) continued

Myeloproliferative neoplasms (MPN), Myelodysplastic neoplasms (MDS), MDS/MPN overlap syndromes

☐ Complete remission (CR)	Number:
	☐ 2nd
	☐ 3rd or higher
	Unknown
☐ Improvement but no CR	
Primary refractory phase (no change)	
Relapse	Number: 1st
	2nd
	☐ 3rd or higher
	Unknown
☐ Progression/Worsening	
☐ Not evaluated	
Unknown	



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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Patient Number in EBMT Registry:	Treatment Date	1 1	(YYYY/MM/DD)

Appendix 1 Best Response and Disease Status (Disease Specific) continued



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст		
Hospital Unique Patient Number (UPN):				
Patient Number in EBMT Registry:	Treatment Date	1	1	(YYYY/MM/DD)

Appendix 1 Best Response and Disease Status (Disease Specific) continued

	continued	
Autoimmune disorders		
☐ No evidence of disease		
☐ Improved		
Unchanged		
Worse		
☐ Not evaluated		
Unknown		
laemoglobinopathies		
<u>Thalassaemia:</u> Complete only for Thalassen	nia Past Daspansa	
☐ Transfusion independent;	Date of last transfusion: / / (YYYY/MM/DD) Unknown (after HCT)	
☐ Transfusions required;	Date of first transfusion: / / (YYYY/MM/DD) Unknown (after HCT)	
☐ Not evaluated		
Unknown		
Complete only for Thalassemia Patient requires transfusion No		
Yes; Date of first transfus (after HCT)	sion: / / (<i>YYYY/MM/DD</i>)	
Number of units: (during follow-up peri		
Did transfusions sto	PP? No Yes; Date of last transfusion:/_/_(YYYY/MM/DD) Unknown Unknown	
Unknown		



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
Hospital Unique Patient Number (UPN):		_	
Patient Number in EBMT Registry:	Treatment Date	1 1	(YYYY/MM/DD)

Appendix 1 Best Response and Disease Status (Disease Specific)

Unknown

continued
laemoglobinopathies
Sickle cell disease:
Complete only for Sickle cell disease Best Response
☐ No return of sickling episodes
Return of sickling episodes; Date of first episode://(YYYY/MM/DD) Unknown (after HCT)
☐ Not evaluated
Unknown
Complete only for Sickle cell disease Disease Status Sickling episodes occur during follow-up period:
No
Yes; First return of sickling episodes after HCT Ongoing presence of sickling episodes episodes Date of first episode://(YYYY/MM/DD) Unknown (after HCT)
Number of SCD episodes: Unknown (after HCT)
☐ Unknown
Other diagnosis No evidence of disease
☐ Improved
□ No response
□ Worse
☐ Not evaluated



EBMT Centre Identification Code (CIC):
Hospital Unique Patient Number (UPN):
Patient Number in EBMT Registry:

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

Appendix 2

-- Pathogens as per EBMT Registry database --

*As defined by the IDSA (Mermel LA, Allon M, Bouza E, Craven DE, Flynn P, O'Grady NP, et al. Clinical practice guidelines for the diagnosis and management of intravascular catheter-related infection: 2009 Update by the Infectious Diseases Society of America. Clin Infect Dis. 2009;49(1):1-45)

Bacterial infections

Gram-positive:

- · Clostridioides difficile
- · Enterococcus faecalis (vancomycin-susceptible)
- · Enterococcus faecalis (vancomycin-resistant)
- · Enterococcus faecium (vancomycin-susceptible)
- · Enterococcus faecium (vancomycin-resistant)
- · Listeria monocytogenes
- · Nocardia spp (specify)
- · Staphylococcus aureus MSSA (methicillin-susceptible)
- · Staphylococcus aureus MRSA (methicillin-resistant) vancomycin-susceptible
- · Staphylococcus aureus MRSA (methicillin-resistant) vancomycin not tested
- · Staphylococcus aureus MRSA and VISA (vancomycin-intermediate, MIC 4-8 µg/ml)
- \cdot Staphylococcus aureus MRSA and VRSA (vancomycin-resistant, MIC \geq 16 $\mu g/ml)$
- · Staphylococcus coagulase-negative spp (at least two positive blood cultures)
- · Streptococcus pneumoniae
- · Streptococcus viridans
- · Streptococcus other spp (specify)
- · Gram-positive bacteria other spp (specify)

Gram-negative:

- · Acinetobacter baumannii
- · Campylobacter jejuni
- · Citrobacter freundii
- · Enterobacter cloacae
- · Enterobacter other spp (specify)
- · Escherichia coli
- · Haemophilus influenzae
- · Helicobacter pylori
- · Klebsiella aerogenes (carbapenem-susceptible)
- · Klebsiella pneumoniae (carbapenem-susceptible)
- · Klebsiella (any species) (carbapenem-resistant) (specify)
- · Legionella pneumophila
- · Morganella morganii
- · Neisseria gonorrhoeae
- · Neisseria meningitidis
- · Proteus vulgaris
- \cdot Providencia spp
- · Pseudomonas aeruginosa (carbapenem-susceptible)
- · Pseudomonas aeruginosa (carbapenem-resistant)
- · Salmonella spp (specify)
- · Serratia marcescens
- · Shigella spp
- · Stenotrophomonas maltophilia
- · Treponema pallidum
- · Gram-negative bacteria other spp (specify)

Other bacteria:

- · Chlamydia spp
- · Chlamydophila
- · Mycobacterium other spp (specify)
- · Mycobacterium tuberculosis
- · Mycoplasma pneumoniae
- · Rickettsia spp
- · Bacteria other (specify)

Viral infections:

- · Adenovirus
- · Gastrointestinal viruses:
 - o Norovirus
 - o Rotavirus
- · Hepatotropic viruses:
 - o HAV
 - o HBV
 - o HCV
 - o HEV
- Herpes group: o CMV
 - o EBV
 - o HHV6
 - o HHV7
 - o HHV8
 - o HS
 - o VZ
- · HIV
- · Human papilloma viruses (HPV)
- · Parvovirus
- · Polyomaviruses:
 - o BK
 - o JC
 - o Merkel cell
 - o Other polyomavirus (specify)
- · Respiratory viruses:
 - o Enterovirus
 - o Human coronavirus
 - o Influenza A
 - o Influenza B
 - o Metapneumovirus
 - o Parainfluenza
 - o Rhinovirus
 - o RSV
 - o SARS-CoV-2
 - o Respiratory virus other (specify)
- · Viruses other (specify)



EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
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Appendix 2

-- Pathogens as per EBMT Registry database -- continued

*As defined by the IDSA (Mermel LA, Allon M, Bouza E, Craven DE, Flynn P, O'Grady NP, et al. Clinical practice guidelines for the diagnosis and management of intravascular catheter-related infection: 2009 Update by the Infectious Diseases Society of America. Clin Infect Dis. 2009;49(1):1-45)

Fungal infections:

Yeasts:

- · Candida albicans
- · Candida auris
- · Candida other (specify)
- · Cryptococcus neoformans
- · Trichosporon (specify)
- · Pneumocytis jiroveci
- · Yeasts other (specify)

Moulds:

- · Aspergillus flavus
- · Aspergillus fumigatus
- · Aspergillus other spp (specify)
- Aspergillus terreus
- · Fusarium other spp (specify)
- · Fusarium solani
- · Lomentospora prolificans (formerly Scedosporium prolificans)
- · Order Mucorales (specify)
- · Dematiaceous fungi (Phaeohyphomycosis) (specify)
- · Scedosporium spp (specify)
- · Moulds other spp (specify)
- \cdot Mould infection diagnosed based on positive galactomannan only, without microbiological confirmation
- · Blastomyces spp
- · Histoplasma spp (specify)
- · Coccidioides spp
- · Paracoccidioides spp

Parasitic infections:

Protozoa:

- · Babesia spp (specify)
- · Cryptosporidium
- · Giardia spp
- · Leishmania spp (specify)
- · Plasmodium spp (specify)
- · Toxoplasma gondii
- Trypanosoma cruziProtozoa other spp (specify)

Helminths:

- · Strongyloides stercoralis
- · Other helminths



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Appendix 3	
 CTCAE term	

CTCAE terms related to infections and infestations (version 5.0.) https://ctep.cancer.gov/protocoldevelopment/electronic_applications/ctc.htm#ctc_50

Respiratory tract infections

- · Pneumonia
- · Other respiratory tract infections

Intra-abdominal infections

- · Esophagus or gastric infection
- · Liver site infection (including biliary tract and gallbladder)
- · Lower gastrointestinal infection
- · Other intra-abdominal infection

Skin, soft tissue and muscle infections

- . Lymph gland infection
- . Skin, soft tissue or muscle infection

Blood infections

- · Bacteremia
- · Fungemia
- · Viremia (including DNAemia)
- . DNAemia for parasitic infection

Other infections

. Device-related infection (other than intravascular catheter)

Uro-genital tract infections

- · Genital infection
- · Urinary tract infection

Nervous system infection

- · Central nervous system infection
- · Other nervous system infection

Cardiovascular infections

- . Endocarditis infective
- . Other cardiovascular infection

Head and neck infections (excluding lymph gland)

- · Conjunctivitis infective
- · Corneal infection
- . Ear infection
- · Endophthalmitis infective
- · Oral cavity infection
- · Retinitis infective
- · Sinusitis infective

Osteoarticular infections

- · Joint infection
- · Bone infection



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Appendix 4

-- Non-infectious Complications CTCAE term -- No Reporting Required

Non-infectious complications

- · Allergic reaction
- · All laboratory abnormalities
- · All types of pain · Alopecia
- Gastritis
- · Blurred vision
- · Hematoma · Diarrhoea (enteropathy) · Hypertension

· Hematologic toxicities

- · Dry mouth
- · Injection site reaction
- · Dyspepsia
- · Malaise
- Dysphagia \cdot Edema
- · Mucositis · Sore throat
- · Esophageal stenosis
- Tinnitus · Vertigo
- Fatigue · Flashes
- · Weight loss

Infectious complications

- Minor ophthalmologic bacterial infections
- External otitis treated topically
- Otitis media treated with oral antibiotics
- Isolated lip herpes simplex
- Bacterial tonsillitis or pharyngitis treated orally
- Laryngitis without viral identification managed at home by inhalations or without any intervention
- URTI without viral/bacterial identification managed at home
- Bilateral cervical lymph node enlargement concurrent with URTI that resolved without specific treatment, together with the resolution of URTI
- Local superficial wound infection resolved under topical antibiotics (incl. impetigo)
- Minor skin bacterial infections
- Minor fungal skin infection
- Diaper rash treated with local antifungals
- · Candidal balanitis treated topically

- · Vaginal candidiasis treated topically or with a single oral dose
- · Asymptomatic bacteriuria due to a pathogen not multi-resistant
- · Single low urinary tract infection treated orally without need for hospitalisation
- · Phlebitis following peripheral intravascular infusion that resolved after intravascular removal without treatment with antibiotics
- · Any isolate that is considered part of the normal flora of the place (oral cavity, vagina, skin, stools) except if it carries an antimicrobial resistance that has clinical implications (induce isolation precautions or a pathogen-directed therapy)
- · Positive culture without clinical implications

Appendix 5

-- Intravascular catheter-related infections --

CVC infections:

- · Catheter colonization · Tunnel infection
- · Phlebitis Pocket infection
- Exit site infection Bloodstream infection



☐ 3 ☐ 4

☐ Present but grade unknown

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Appendix 6 Cell Infusion Sheet Chronological number of CI episode for this patient: Date of the first infusion (after HCT): _ _ _ / _ _ (YYYY/MM/DD) Number of infusions within this episode (10 weeks): (Count only infusions that are part of the same regimen and given for the same indication.) Source of cells: □ Allogeneic ☐ Autologous Type of cells: □ Lymphocytes (DLI) ☐ Fibroblasts ☐ Dendritic cells ☐ NK cells ☐ Regulatory T-cells ☐ Gamma/delta cells ☐ Virus-specifc T-cells; specify virus: ____ ☐ Other; specify: __ Not applicable for Inborn Errors Disease status at time of this cell infusion*: _

* Indicate the disease status corresponding to indication diagnosis by selecting from the list provided in Appendix 1

Indication: (check all that apply)	Poor graft function Infection prophylaxis Other; specify:				
Acute GvHD maximum grade (after this infusion episode but before any subsequent cell infusion/HCT/CT): 0 (none) Date Acute GvHD onset after cell infusion: / / (YYYY/MM/DD)					

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☐ Unknown