

HAEMATOPOIETIC CELL TRANSPLANTATION (HCT) --- Annual/Unscheduled 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)

| | |
|--|--|
| <input type="checkbox"/> Relapse or progression/persistent disease | |
| <input type="checkbox"/> Secondary malignancy | |
| <input type="checkbox"/> CT-related | Select treatment related cause: <i>(select all that apply)</i> <input type="checkbox"/> Graft versus Host Disease <input type="checkbox"/> Non-infectious complication <input type="checkbox"/> Infectious complication: <i>(select all that apply)</i> <input type="checkbox"/> Bacterial infection <input type="checkbox"/> Viral infection <input type="checkbox"/> Fungal infection <input type="checkbox"/> Parasitic infection <input type="checkbox"/> Infection with unknown pathogen <input type="checkbox"/> Other treatment related cause of death; specify: _____ |
| <input type="checkbox"/> HCT-related | |
| <input type="checkbox"/> GT-related | |
| <input type="checkbox"/> IST-related | |
| <input type="checkbox"/> Unknown | |
| <input type="checkbox"/> Other cause of death; specify: _____ | |

BEST RESPONSE

*Complete only for the first annual follow-up
 Not applicable for Inborn Errors*

Best clinical/biological response after HCT* (observed before any subsequent treatment): _____

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

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

GRAFT FUNCTION

Poor graft function (defined as: frequent dependence on blood and/or platelet transfusions and/or growth factor support in the absence 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 since last follow-up:
(complete only if patient received an allogeneic HCT. This section is optional for malignant disorders)

Chimaerism test date: ____/____/____ (YYYY/MM/DD) Unknown

Source of cells tested: Peripheral blood
 Bone marrow

Select cell type and complete relevant test results:

- Global (%): _____ % donor Unknown
 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 allogeneic HCT)

Immunosuppression during this follow-up period:

- No
 Yes; **Immunosuppression stopped:**
 No
 Yes; **End date:** ____/____/____ (YYYY/MM/DD) Unknown
 Unknown
 Unknown



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

COMPLICATIONS SINCE THE LAST REPORT

-- GvHD --
Allogeneic HCT only

Did graft versus host disease (GvHD) occur during this follow-up period?

No (proceed to 'Complications since the last report - Non-infectious complications')

Yes: **Did the patient receive a systemic/immunosuppressive treatment for GvHD during this follow-up period?**

No

Yes: Started in this follow-up period; **Date treatment started:** ____/____/____ (YYYY/MM/DD) Unknown

Ongoing since previous follow-up

Treatment stopped: No

Yes; **Stop date of treatment:** ____/____/____ (YYYY/MM/DD) Unknown

Unknown

Unknown (proceed to 'Complications since the last report - Non-infectious complications')

Did acute GvHD occur during this follow-up period?

No

Yes: Started in this follow-up period; **Date of onset:** ____/____/____ (YYYY/MM/DD) Unknown

Ongoing since previous follow-up

Maximum observed organ severity score during this period:

| | | | | | | | |
|----------------------|-----------------------------------|----------------------------|--|----------------------------|----------------------------------|--|----------------------------------|
| Skin: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> 4 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Liver: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> 4 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Lower GI tract: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> 4 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Upper GI tract: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> Not evaluated | | <input type="checkbox"/> Unknown | | |
| Other site affected: | <input type="checkbox"/> No | | <input type="checkbox"/> Yes; specify: _____ | | | | |

Overall maximum grade observed: 1 2 3 4 Unknown Not evaluated

Steroid-refractory acute GvHD: No

Yes: Started in this follow-up period; **Date of onset:** ____/____/____ (YYYY/MM/DD) Unknown

Ongoing since previous follow-up

Unknown

Unknown

COMPLICATIONS SINCE THE LAST REPORT continued

-- GvHD --

Allogeneic HCT only

Did chronic GvHD occur during this follow-up period?

No

Yes: Started in this follow-up period; **Date of onset:** ____/____/____ (YYYY/MM/DD) Unknown

Ongoing since previous follow-up

Maximum NIH score during this period: Mild
 Moderate
 Severe
 Unknown
 Not evaluated

Date of maximum NIH score: ____/____/____ (YYYY/MM/DD) Unknown

Maximum observed organ severity score during this period:

| | | | | | | |
|----------------------|-----------------------------------|--|----------------------------|----------------------------|--|----------------------------------|
| Skin: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Oral: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Gastrointestinal: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Eyes: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Liver: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Joints and fascia: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Lungs: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Genitalia: | <input type="checkbox"/> 0 (none) | <input type="checkbox"/> 1 | <input type="checkbox"/> 2 | <input type="checkbox"/> 3 | <input type="checkbox"/> Not evaluated | <input type="checkbox"/> Unknown |
| Other site affected: | <input type="checkbox"/> No | <input type="checkbox"/> Yes; specify: _____ | | | | |

Steroid-refractory chronic GvHD: No

Yes: Started in this follow-up period; **Date of onset:** ____/____/____ (YYYY/MM/DD) Unknown

Ongoing since previous follow-up

Unknown

Was overlap syndrome observed: No Yes Unknown
(features of both chronic and acute GvHD)

Unknown



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

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)
 Unknown

Secondary graft failure

Complication observed during this follow-up period? No
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum grade observed during this period: Non-fatal Fatal

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Cardiac event

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Central nervous system (CNS) toxicity

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Gastrointestinal (GI) Toxicity (non-GvHD and non-infectious related)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

* Grade 0-2



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

COMPLICATIONS SINCE THE LAST REPORT

-- Non-infectious complications --

Liver disorder

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Renal failure (chronic kidney disease, acute kidney injury)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Respiratory disorders

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Skin Toxicity (non-GvHD and non-infectious related)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

* Grade 0-2



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

COMPLICATIONS SINCE THE LAST REPORT

-- Non-infectious complications --

Vascular event

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Avascular necrosis (AVN)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Cerebral haemorrhage

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Haemorrhage (other than cerebral haemorrhage)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

* Grade 0-2



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

COMPLICATIONS SINCE THE LAST REPORT

-- Non-infectious complications --

Cerebral thrombosis

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Cytokine release syndrome (CRS)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Haemophagocytic lymphohistiocytosis (HLH)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Pure red cell aplasia (PRCA)

Complication observed during this follow-up period? No
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum grade observed during this period: Non-fatal Fatal

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

* Grade 0-2



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

COMPLICATIONS SINCE THE LAST REPORT

-- Non-infectious complications --

Posterior reversible encephalopathy syndrome (PRES)

Complication observed during this follow-up period? No
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum grade observed during **this period**: Non-severe Severe Fatal Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Transplant-associated microangiopathy (TMA)

Complication observed during this follow-up period? No*
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum grade observed during **this period**: Non-severe Severe Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Sinusoidal obstruction syndrome/Veno-occlusive disease (SOS/VOD)

Complication observed during this follow-up period? No
 Yes: Newly developed Ongoing since previous assessment
 Unknown

Maximum grade observed during **this period**: Mild Moderate Severe Very severe Fatal Unknown

Onset date (YYYY/MM/DD): ____/____/____ Unknown *Only if newly developed*

Other complication observed during this follow-up period? No*

Yes: Newly developed Ongoing since previous assessment
 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 *Only if newly developed*

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

* Grade 0-2

COMPLICATIONS SINCE THE LAST REPORT

-- 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)
 Unknown

Bacterial infection: No Yes Unknown

1) **New or ongoing:** Newly developed Ongoing since previous assessment

Start date: ____/____/____ (YYYY/MM/DD) *only if newly developed* Unknown

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):** _____

(if patient died)

Contributory cause of death: No Yes Unknown

If more than 1 bacterial 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

COMPLICATIONS SINCE THE LAST REPORT

-- Infectious complications -- continued

Viral infection: No Yes Unknown

1) **New or ongoing:** Newly developed Ongoing since previous assessment

Start date: ____/____/____ (YYYY/MM/DD) *only if newly developed* Unknown

Pathogen*: _____

Infection with clinical implications: No

Yes***: (select all that apply during this period)

Symptoms/signs of disease

Administration of pathogen-directed therapy (including pre-emptive therapy)

Unknown

Indicate at least 1 location involved during this period:

Localisation 1 (CTCAE term):** _____

Localisation 2 (CTCAE term):** _____

Localisation 3 (CTCAE term):** _____

(if patient died)

Contributory cause of death: No Yes Unknown

If more than 1 viral 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

*** Also answer 'Yes' in case of pre-emptive therapy

COMPLICATIONS SINCE THE LAST REPORT

-- Infectious complications -- continued

Invasive fungal infection: No Yes Unknown

1) **New or ongoing:** Newly developed Ongoing since previous assessment

Start date: ____/____/____ (YYYY/MM/DD) *only if newly developed* Unknown

Yeasts Moulds

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):** _____

(if patient died)

Contributory cause of death: No Yes Unknown

If more than 1 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

COMPLICATIONS SINCE THE LAST REPORT

-- Infectious complications -- continued

Parasitic infection: No Yes Unknown

1) **New or ongoing:** Newly developed Ongoing since previous assessment

Start date: ____/____/____ (YYYY/MM/DD) *only if newly developed* Unknown

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 2 (CTCAE term):** _____

Localisation 3 (CTCAE term):** _____

(if patient died)

Contributory cause of death: No Yes Unknown

If more than 1 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

COMPLICATIONS SINCE THE LAST REPORT

-- Infectious complications -- continued

Infection with unknown pathogen: No Yes Unknown

(for clinical infections without microbiological documentation, like pneumonia, cellulitis, etc.)

1) **New or ongoing:** Newly developed Ongoing since previous assessment

Start date: ____/____/____ (YYYY/MM/DD) *only if newly developed* Unknown

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)*: _____

(if patient died)

Contributory cause of death: No Yes Unknown

If more than 1 infections with unknown pathogen, copy and fill-in this table as many times as necessary.

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



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

SECONDARY MALIGNANCIES AND AUTOIMMUNE DISORDERS

Did secondary malignancy or autoimmune disorder occur since the last follow-up?

- No
- Yes; **Was it a secondary malignancy or autoimmune disorder?**
- Secondary malignancy
 - Autoimmune disorder

Date of diagnosis: ____/____/____ (YYYY/MM/DD)

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

ADDITIONAL TREATMENTS

Did the patient receive any additional disease treatment since the last follow-up?

- No
- Yes; Started in this follow-up period;
 Ongoing since previous follow-up
- Unknown

complete the "Treatment — non-HCT/CT/GT/IST form and /or the Cell infusion sheet"

ADDITIONAL CELL INFUSIONS

Did the patient receive additional cell infusions (excluding a new HCT and CT) since the last follow-up?

No

Yes: **Is this cell infusion an allogeneic boost* ?** No Yes

** An allogeneic boost is an infusion of cells from the same donor without conditioning, with no evidence of graft rejection.*

Date of the allogeneic boost: ____/____/____ (YYYY/MM/DD)

Is this cell infusion an autologous boost? No Yes

Date of the autologous boost: ____/____/____ (YYYY/MM/DD)

Unknown

If this cell infusion is not a boost, attach the Cell Infusion (CI) sheet available in Appendix 6, completing as many sheets as episodes of cell infusion that took place during this interval; then continue below.

Did the patient receive subsequent HCT/CT (either at your or another centre)?

No

Yes

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

RELAPSE, PROGRESSION, RECURRENCE OF DISEASE OR SIGNIFICANT WORSENING
(not relevant for Inborn errors)

MRD detectable (with any method): (only for Acute leukaemia)

- No
 Yes: **Date of first detectable MRD:** ____/____/____ (YYYY/MM/DD) Unknown
 Unknown

Was there a relapse, progression, recurrence of disease or significant worsening of organ function related to the primary disease since last follow-up? *(detected by any method)*

- No
 Yes; *for every relapse, progression, recurrence, significant worsening complete the questions below*

Type: Relapse / Recurrence of disease

(Continuous) progression / Significant worsening

Date of relapse/progression/recurrence/worsening: ____/____/____ (YYYY/MM/DD) Unknown

Malignant disorders only:

Type of relapse/progression:

Medullary: No Yes Unknown

Extramedullary: No Yes Unknown

If the relapse/progression was extramedullary or both medullary and extramedullary:

Involvement at time of relapse/progression:

Skin: No Yes Not evaluated

CNS: No Yes Not evaluated

Testes/Ovaries: No Yes Not evaluated

Other: No Yes; specify: _____

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



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

DISEASE STATUS
Disease specific

Disease status at this follow-up or at time of death*: _____

* 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

PREGNANCY AFTER HCT

Has patient become pregnant or impregnated another person since last follow-up?

No;

Yes: **Did the pregnancy result in a live birth?**

No; **Date of spontaneous or induced termination:** ____/____/____ (YYYY/MM/DD) Unknown

Yes; **Year of birth:** ____ (YYYY) **Month of birth:** __ (MM) Unknown

Still pregnant at time of follow-up

Unknown

Unknown

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 20 |
| CHRONIC LEUKAEMIAS | Go to page 20 |
| PLASMA CELL NEOPLASMS (PCN) | Go to page 21 |
| MPN, MDS, MDS / MPN OVERLAP SYNDROMES | Go to page 23 |
| AUTOIMMUNE DISORDERS | Go to page 24 |
| HAEMOGLOBINOPATHIES | Go to page 24 |
| LYMPHOMAS | Go to page 25 |
| SOLID TUMOURS | Go to page 25 |
| BONE MARROW FAILURE SYNDROMES (BMF) including APLASTIC ANAEMIA (AA) | Go to page 25 |
| OTHER DIAGNOSIS | Go to page 26 |
| | |

Appendix 1
 Best Response and Disease Status (Disease Specific)

Acute leukaemias (AML, PLN, Other)

| |
|--|
| <input type="checkbox"/> Complete remission (CR) |
| <input type="checkbox"/> Not in complete remission |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

Proceed to next page for Diseases Status section

Chronic leukaemias (CML, CLL, PLL, Other)
Chronic Myeloid Leukaemia (CML):

| |
|--|
| <input type="checkbox"/> Chronic phase (CP); Number: <input type="checkbox"/> 1 st <input type="checkbox"/> 2 nd <input type="checkbox"/> 3 rd or higher <input type="checkbox"/> Unknown Haematological remission: <input type="checkbox"/> No <input type="checkbox"/> Yes <input type="checkbox"/> Not evaluated <input type="checkbox"/> Unknown Cytogenetic remission: <input type="checkbox"/> No <input type="checkbox"/> Yes <input type="checkbox"/> Not evaluated <input type="checkbox"/> Unknown Molecular remission: <input type="checkbox"/> No <input type="checkbox"/> Yes <input type="checkbox"/> Not evaluated <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Accelerated phase; Number: <input type="checkbox"/> 1 st <input type="checkbox"/> 2 nd <input type="checkbox"/> 3 rd or higher <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Blast crisis; Number: <input type="checkbox"/> 1 st <input type="checkbox"/> 2 nd <input type="checkbox"/> 3 rd or higher <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

Proceed to next page for Diseases Status section

Appendix 1
Best Response and Disease Status (Disease Specific)

Chronic Lymphocytic Leukaemia (CLL), Prolymphocytic Leukaemia (PLL) and other chronic leukaemias:

| |
|--|
| <input type="checkbox"/> Complete remission (CR) |
| <input type="checkbox"/> Partial remission (PR) |
| <input type="checkbox"/> Progression: <input type="checkbox"/> Resistant to last regimen <input type="checkbox"/> Sensitive to last regimen <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Stable disease (no change, no response/loss of response) |
| <input type="checkbox"/> Relapse |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

Proceed to next page for Diseases Status section

Plasma cell neoplasms (PCN)

| | |
|---|---|
| <input type="checkbox"/> Complete remission (CR) | Number: <input type="checkbox"/> 1st <input type="checkbox"/> 2nd <input type="checkbox"/> 3rd or higher <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Stringent complete remission (sCR) | |
| <input type="checkbox"/> Very good partial remission (VGPR) | |
| <input type="checkbox"/> Partial remission (PR) | |
| <input type="checkbox"/> Relapse | |
| <input type="checkbox"/> Progression | |
| <input type="checkbox"/> Stable disease (no change, no response/loss of response) | |
| <input type="checkbox"/> Not evaluated | |
| <input type="checkbox"/> Unknown | |

Proceed to next page for Diseases Status section



EBMT Centre Identification Code (CIC): _____

Treatment Type HCT

Hospital Unique Patient Number (UPN): _____

Patient Number in EBMT Registry: _____

Treatment Date ____/____/____ (YYYY/MM/DD)

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

Complete only for PCN Disease Status

Was the patient on dialysis during this follow-up period?

- No
- Yes; Started in this follow-up period: **Start date:** ____/____/____ (YYYY/MM/DD) Unknown
- Ongoing since previous follow-up
- Did dialysis stop?** No
- Yes; **End date:** ____/____/____ (YYYY/MM/DD) Unknown
- Unknown

Complete only for AL, CLL and PCN Disease Status

Leukaemias (AL, CLL) and PCN (complete only for patient in CR or sCR)

Minimal residual disease (MRD):

- Positive
 - Negative
 - Not evaluated
 - Unknown
- Date MRD status evaluated:** ____/____/____ (YYYY/MM/DD) Unknown



EBMT Centre Identification Code (CIC): _____

Treatment Type HCT

Hospital Unique Patient Number (UPN): _____

Patient Number in EBMT Registry: _____

Treatment Date ____/____/____ (YYYY/MM/DD)

Appendix 1**Best Response and Disease Status (Disease Specific)
continued****Myeloproliferative neoplasms (MPN), Myelodysplastic neoplasms (MDS), MDS/MPN overlap syndromes**

| | |
|---|---|
| <input type="checkbox"/> Complete remission (CR) | <u>Number:</u> <input type="checkbox"/> 1st <input type="checkbox"/> 2nd <input type="checkbox"/> 3rd or higher <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Improvement but no CR | |
| <input type="checkbox"/> Primary refractory phase (no change) | |
| <input type="checkbox"/> Relapse | <u>Number:</u> <input type="checkbox"/> 1st <input type="checkbox"/> 2nd <input type="checkbox"/> 3rd or higher <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Progression/Worsening | |
| <input type="checkbox"/> Not evaluated | |
| <input type="checkbox"/> Unknown | |

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

Autoimmune disorders

| |
|---|
| <input type="checkbox"/> No evidence of disease |
| <input type="checkbox"/> Improved |
| <input type="checkbox"/> Unchanged |
| <input type="checkbox"/> Worse |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

Haemoglobinopathies

Thalassaemia:

Complete only for Thalassemia Best Response

| | |
|--|--|
| <input type="checkbox"/> Transfusion independent | Date of last transfusion: ____/____/____ (YYYY/MM/DD) <input type="checkbox"/> Unknown <i>(after HCT)</i> |
| <input type="checkbox"/> Transfusions required; | Date of first transfusion: ____/____/____ (YYYY/MM/DD) <input type="checkbox"/> Unknown <i>(after HCT)</i> |
| <input type="checkbox"/> Not evaluated | |
| <input type="checkbox"/> Unknown | |

Complete only for Thalassemia Disease Status

Patient requires transfusions during follow-up period:

No

Yes; Return to transfusion dependence after HCT or transfusion free period; **Date of first transfusion:** ____/____/____ (YYYY/MM/DD) Unknown
(after HCT or transfusion free period)

Ongoing transfusion dependence since previous assessment

Number of units: ____ Unknown
(during follow-up period)

Did transfusions stop? No

Yes; **Date of last transfusion:** ____/____/____ (YYYY/MM/DD) Unknown

Unknown

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

| |
|---|
| <input type="checkbox"/> Chemorefractory relapse or progression, including primary refractory disease |
| <input type="checkbox"/> Complete remission (CR) |
| <input type="checkbox"/> Partial remission (PR) |
| <input type="checkbox"/> Stable disease (no change, no response/loss of response) |
| <input type="checkbox"/> Untreated relapse (from a previous CR) or progression (from a previous PR) |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

Technique used for disease assessment: CT scan
 PET
 MRI
 Unknown

Solid tumours

| |
|--|
| <input type="checkbox"/> Complete remission (CR): <input type="checkbox"/> Confirmed <input type="checkbox"/> Unconfirmed <input type="checkbox"/> Unknown |
| <input type="checkbox"/> First partial remission |
| <input type="checkbox"/> Partial remission (PR) |
| <input type="checkbox"/> Progressive disease |
| <input type="checkbox"/> Relapse: <input type="checkbox"/> Resistant <input type="checkbox"/> Sensitive <input type="checkbox"/> Unknown |
| <input type="checkbox"/> Stable disease (no change, no response/loss of response) |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

Bone marrow failures (incl. AA)

| |
|---|
| <input type="checkbox"/> Complete remission (CR) |
| <input type="checkbox"/> Partial remission (PR) |
| <input type="checkbox"/> Haematological improvement (HI); <i>NIH partial response</i> |
| <input type="checkbox"/> Stable disease (no change, no response/loss of response) |
| <input type="checkbox"/> Relapse / Progression |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

Complete only for Bone marrow failures (incl. AA) Disease Status

Did transfusions stop during the follow-up period? Patient was never transfusion dependent
 No
 Yes; **Did the patient return to transfusion dependency afterwards?**
 No
 Yes; **First transfusion date:** ____/____/____ (YYYY/MM/DD) Unknown (after transfusion free period)
 Unknown
 Unknown

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

Haemoglobinopathies

Sickle cell disease:

Complete only for Sickle cell disease Best Response

| | |
|---|---|
| <input type="checkbox"/> No return of sickling episodes | |
| <input type="checkbox"/> Return of sickling episodes; | Date of first episode: ____/____/____ (YYYY/MM/DD) <input type="checkbox"/> Unknown (after HCT) |
| <input type="checkbox"/> Not evaluated | |
| <input type="checkbox"/> Unknown | |

Complete only for Sickle cell disease Disease Status

Sickling episodes occur during follow-up period:

| | |
|--|--|
| <input type="checkbox"/> No | |
| <input type="checkbox"/> Yes; <input type="checkbox"/> First return of sickling episodes after HCT | Date of first episode : ____/____/____ (YYYY/MM/DD) <input type="checkbox"/> Unknown (after HCT) |
| <input type="checkbox"/> Ongoing presence of sickling episodes | |
| Number of SCD episodes: ____ <input type="checkbox"/> Unknown (during follow-up) | |
| <input type="checkbox"/> Unknown | |

Other diagnosis

| |
|---|
| <input type="checkbox"/> No evidence of disease |
| <input type="checkbox"/> Improved |
| <input type="checkbox"/> No response |
| <input type="checkbox"/> Worse |
| <input type="checkbox"/> Not evaluated |
| <input type="checkbox"/> Unknown |

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:

- . Bacillus (in blood: report only if ≥2 positive separately taken cultures)
- . Clostridioides difficile (c difficile/CDT/CDI)
- . Clostridium other (NOT difficile)
- . Corynebacterium jeikeium
- . Corynebacterium other (NOT jeikeium) (in blood: report only if ≥2 positive separately taken cultures)
- . Enterococcus faecalis (vancomycin-susceptible)
- . Enterococcus faecalis (vancomycin-resistant)
- . Enterococcus faecium (vancomycin-susceptible)
- . Enterococcus faecium (vancomycin-resistant)
- . Listeria monocytogenes
- . Nocardia (specify)
- . Propionibacterium (in blood: report only if ≥2 positive separately taken cultures)
- . Rothia
- . Staphylococcus aureus (s aureus/staph aureus) MSSA (methicillin-susceptible)
- . Staphylococcus aureus (s aureus/staph aureus) MRSA (methicillin-resistant vancomycin-susceptible)
- . Staphylococcus coagulase-negative (in blood: report only if ≥2 positive separately taken cultures)
- . Staphylococcus lugdunensis
- . Streptococcus pneumoniae (pneumococcus)
- . Streptococcus viridans
- . Streptococcus other (specify)
- . Gram-positive bacteria other (specify)

Gram-negative:

- . Acinetobacter baumannii
- . Acinetobacter lwoffii (in blood: report only if ≥2 positive separately taken cultures)
- . Acinetobacter other (NOT baumannii, NOT lwoffii)
- . Bacteroides fragilis
- . Bordetella pertussis
- . Borrelia
- . Brucella
- . Campylobacter jejuni
- . Citrobacter freundii
- . Coxiella burnetii (Q fever)
- . Enterobacter cloacae
- . Enterobacter other
- . Escherichia coli (e coli)
- . Fusobacterium
- . Haemophilus influenzae (haemophilus influenzae type B/Hib/hemophilus influenzae)
- . Haemophilus other (hemophilus)
- . Helicobacter pylori
- . Klebsiella pneumoniae (carbapenem-susceptible)
- . Klebsiella other (NOT pneumoniae) (carbapenem-susceptible)
- . Klebsiella (any species) (carbapenem-resistant)
- . Klebsiella (any species) (carbapenem-susceptibility not checked)
- . Legionella pneumophila
- . Morganella morganii
- . Micrococcus (in blood: report only if ≥2 positive separately taken cultures)
- . Moraxella catarrhalis
- . Neisseria gonorrhoeae (gonococcus)
- . Neisseria meningitidis (meningococcus)
- . Proteus vulgaris
- . Providencia
- . Pseudomonas aeruginosa (PSA) (carbapenem-susceptible)
- . Pseudomonas aeruginosa (PSA) (carbapenem-resistant)
- . Pseudomonas aeruginosa (PSA) (carbapenem-susceptibility not checked)
- . Pseudomonas other (NOT aeruginosa)
- . Raoultella
- . Salmonella (specify)
- . Serratia marcescens
- . Shigella
- . Stenotrophomonas maltophilia
- . Treponema pallidum (syphilis/lues)
- . Yersinia
- . Gram-negative bacteria other (specify)

Other bacteria:

- . Chlamydia
- . Chlamydoxiphila
- . Mycobacterium other (specify)
- . Mycobacterium tuberculosis (TB)
- . Mycoplasma pneumoniae
- . Rickettsia
- . Ureoplasma
- . Bacteria other (specify)

Viral infections:

- . Adenovirus (ADV)
- . Chikungunya virus
- . Crimean-Congo haemorrhagic fever virus (CCHFV)
- . Dengue virus
- . Gastrointestinal viruses:
 - o Norovirus
 - o Rotavirus
 - o Sapovirus
 - o Astrovirus
- . Hepatotropic viruses:
 - o Hepatitis A virus (HAV)
 - o Hepatitis B virus (HBV)
 - o Hepatitis C virus (HCV)
 - o Hepatitis D virus (HDV)
 - o Hepatitis E virus (HEV)
- . Herpes group:
 - o Cytomegalovirus (CMV)
 - o Epstein-Barr virus (EBV)
 - o Herpes simplex virus (HS/HSV)
 - o Herpesvirus 6 (HHV6)
 - o Herpesvirus (HHV7)
 - o Herpesvirus 8 (HHV8/Kaposi's sarcoma-associated herpesvirus/KSHV/Kaposi)
 - o Varicella zoster virus (VZV/VZV/HSV/shingles/zoster/chickenpox)
- . Human immunodeficiency virus (HIV)
- . Human papilloma viruses (HPV)
- . Human T-lymphotropic virus 1 (HTLV-1)
- . Human T-lymphotropic virus 2 (HTLV-2)
- . Measles virus
- . Mumps virus
- . Parechovirus
- . Parvovirus (parvovirus B-19/B-19)
- . Poliovirus
- . Polyomaviruses:
 - o BK polyomavirus (BKV)
 - o JC virus (JCV)
 - o Merkel cell virus
- . Respiratory viruses:
 - o Bocavirus
 - o Enterovirus
 - o Human coronavirus (excluding SARS-CoV-2 or COVID-19)
 - o Influenza A virus (including birdflu)
 - o Influenza B virus
 - o Human metapneumovirus (hMPV)
 - o Parainfluenza
 - o Rhinovirus
 - o Respiratory syncytial virus (RSV)
 - o SARS-CoV-2 virus (COVID-19)
- . Rubella virus
- . Sandfly viruses (Naples virus/Sicilian virus/Toscana virus/Cyprus virus/Turkey virus/Tehran virus/phlebovirus)
- . Tick-borne encephalitis virus (TBE)
- . West Nile virus (WNV)
- . Yellow fever virus
- . Zika virus (ZIKV)

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
- Geotrichum
- Magnusiomyces
- Pneumocystis jirovecii
- Saccharomyces
- Saprochaete
- Trichosporon
- Yeasts other (specify)

Moulds:

- Aspergillus flavus
- Aspergillus fumigatus
- Aspergillus other (NOT flavus, NOT fumigatus, NOT terreus)
- Aspergillus terreus
- Blastomyces
- Coccidioides
- Dematiaceous fungi / phaeohyphomycosis (specify)
- Fusarium solani
- Fusarium other (NOT solani)
- Galactomannan positive in blood or BAL, without microbiological confirmation of fungal infection
- Histoplasma
- Lomentospora prolificans / scedosporium prolificans
- Mucorales (mucor/rhizomucor/rhizopus/lichteinia) (specify)
- Paracoccidioides
- Scedosporium other (NOT prolificans) (specify)
- Moulds other (specify)

Parasitic infections:**Protozoa:**

- Amoeba
- Babesia
- Cryptosporidium
- Giardia
- Leishmania
- Plasmodium (malaria)
- Toxoplasma gondii
- Trypanosoma cruzi
- Protozoa other

Helminths:

- Schistosoma
- Strongyloides stercoralis
- Helminths other

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, please specify:

Intra-abdominal infections

- Esophagus or gastric infection
- Liver site infection (including biliary tract and gallbladder), please specify:
- Lower gastrointestinal infection, please specify:
 - . Enteritis infective, please specify:
- Other intra-abdominal infection, please specify:

Skin, soft tissue and muscle infections

- . Lymph gland infection
- . Skin, soft tissue or muscle infection, please specify:

Blood infections

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

Other infections

- . Device-related infection (other than intravascular catheter)
- . Post-transplant lymphoproliferative disorder (PTLD)**

Uro-genital tract infections

- Genital infection, please specify:
- Urinary tract infection, please specify:

Nervous system infection

- Central nervous system infection, please specify:
- Other nervous system infection, please specify:

Cardiovascular infections

- . Endocarditis infective
- . Other cardiovascular infection, please specify:

Head and neck infections (excluding lymph gland)

- . Ear infection
- Oral cavity infection, please specify:
- Retinitis infective
- Sinusitis infective
- Other eye infection, please specify:

Osteoarticular infections

- Joint infection
- Bone infection

* Only if pathogen 'JC virus' is selected

** Only if pathogen 'Epstein-Barr virus' is selected

Appendix 4

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

Non-infectious complications

Infectious complications

- Allergic reaction
- All laboratory abnormalities
- All types of pain
- Alopecia
- Blurred vision
- Dry mouth
- Dyspepsia
- Dysphagia
- Edema
- Esophageal stenosis
- Fatigue
- Flashes
- Gastritis
- Hematologic toxicities
- Hematoma
- Hypertension
- Injection site reaction
- Malaise
- Mucositis
- Sore throat
- Tinnitus
- Vertigo
- Weight loss

- 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
- Neutropenic fever and sepsis of unknown origin

Appendix 5

-- Intravascular catheter-related infections --

CVC infections:

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

Appendix 6
Cell Infusion Sheet

Chronological number of CI episode for this patient: _____

Date of the first infusion (within this episode): ____/____/____ (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)
- Mesenchymal
- Fibroblasts
- Dendritic cells
- NK cells
- Regulatory T-cells
- Gamma/delta cells
- Virus-specific T-cells (VST)
- Other; specify: _____

(Only if 'Type of cells: = Virus-specific T-cells (VST))'

Specificity of VST product: Single-virus specific Multi-virus specific

If single-virus specific, to which virus? *(Please register this virus in the Infectious complications - Viral infection part and fill in the VST in the 'Pre-emptive viral therapy' or 'Treatment of end-organ viral disease' section)*

- EBV
- CMV
- ADV
- BKV or JCV
- HHV-6

If multi-virus specific, to which viruses? *(Please register these viruses in the Infectious complications - Viral infection part and fill in the VST in the 'Pre-emptive viral therapy' or 'Treatment of end-organ viral disease' section)*
 (Select all that apply)



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

Treatment Type HCT
 Treatment Date ____/____/____ (YYYY/MM/DD)

Appendix 6
 Cell Infusion Sheet continued

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)

- | | |
|--|--|
| <input type="checkbox"/> Planned/protocol <input type="checkbox"/> Prophylactic <input type="checkbox"/> Treatment of acute GvHD <input type="checkbox"/> Treatment of chronic GvHD <input type="checkbox"/> Treatment PTLD, EBV lymphoma <input type="checkbox"/> Treatment for primary disease <input type="checkbox"/> Mixed chimaerism <input type="checkbox"/> Loss/decreased donor chimaerism <input type="checkbox"/> Treatment of viral infection other than EBV | <input type="checkbox"/> Poor graft function <input type="checkbox"/> Infection prophylaxis <input type="checkbox"/> Other; specify: _____ |
|--|--|

Only for Acute leukemia donor lymphocyte infusions:

Response to DLI:

- Complete remission (CR) **MRD status:** MRD negative MRD positive Not evaluated Unknown
 Not in CR
 Not evaluated
 Unknown