

<h1>DAY 0</h1>	<h1>MED-B</h1> <h1>GENERAL INFORMATION</h1>
----------------	--

**TEAM**

EBMT Centre Identification Code (CIC) .....

Hospital ..... Unit .....

Contact person: .....

e-mail .....

Date of this report ..... - ..... - .....  
yyyy mm dd

**STUDY/TRIAL**

Patient following national / international study / trial:  No  Yes  Unknown

Name of study / trial .....

**PATIENT**

Unique Identification Code (UIC) ..... (to be entered only if patient previously reported)

**Hospital Unique Patient Number or Code (UPN):** .....

**Compulsory, registrations will not be accepted without this item.**

*All transplants performed in the same patient must be registered with the same patient identification number or code as this belongs to the patient and not to the transplant.*

Initials ..... (first name(s) – surname(s))

Date of birth ..... - ..... - ..... Sex:  Male  Female  
yyyy mm dd (at birth)

ABO Group ..... Rh factor:  Absent  Present  Not evaluated

**DISEASE**

Date of diagnosis : ..... - ..... - .....  
yyyy mm dd

**PRIMARY DISEASE DIAGNOSIS** (CHECK THE DISEASE FOR WHICH THIS TRANSPLANT WAS PERFORMED)

- |  |  |  |
|--|--|--|
| <input type="checkbox"/> Primary Acute Leukaemia<br><input type="checkbox"/> Acute Myelogenous Leukaemia (AML) & related Precursor Neoplasms<br><input type="checkbox"/> Precursor Lymphoid Neoplasms (old ALL)<br><input type="checkbox"/> Therapy related myeloid neoplasms (old Secondary Acute Leukaemia)<br><input type="checkbox"/> Chronic Leukaemia<br><input type="checkbox"/> Chronic Myeloid Leukaemia (CML)<br><input type="checkbox"/> Chronic Lymphocytic Leukaemia (CLL)<br><input type="checkbox"/> Lymphoma<br><input type="checkbox"/> Non Hodgkin<br><input type="checkbox"/> Hodgkin's Disease | <input type="checkbox"/> Myeloma /Plasma cell disorder<br><input type="checkbox"/> Solid Tumour<br><input type="checkbox"/> Myelodysplastic syndromes / Myeloproliferative neoplasm<br><input type="checkbox"/> MDS<br><input type="checkbox"/> MDS/MPN<br><input type="checkbox"/> Myeloproliferative neoplasm<br><input type="checkbox"/> Bone marrow failure including Aplastic anaemia<br><input type="checkbox"/> Inherited disorders<br><input type="checkbox"/> Primary immune deficiencies<br><input type="checkbox"/> Metabolic disorders | <input type="checkbox"/> Histiocytic disorders<br><input type="checkbox"/> Autoimmune disease<br><input type="checkbox"/> Juvenile Idiopathic Arthritis (JIA)<br><input type="checkbox"/> Multiple Sclerosis<br><input type="checkbox"/> Systemic Lupus<br><input type="checkbox"/> Systemic Sclerosis<br><input type="checkbox"/> Haemoglobinopathy |
|--|--|--|

Other diagnosis, specify: \_\_\_\_\_



## FIRST LINE THERAPIES

**DISEASE MODIFYING DRUGS AND IMMUNOSUPPRESSANTS**

No – Proceed to "Date of HSCT"

Yes:

**Date started** ..... - ..... - .....  
yyyy mm dd

- Yes, mark appropriate box(es)
- |  |  |
|--|--|
| <input type="checkbox"/> Cyclophosphamide                                  | <input type="checkbox"/> Cyclosporin-A   |
| <input type="checkbox"/> Methotrexate                                      | <input type="checkbox"/> Corticosteroids |
| <input type="checkbox"/> Non-steroidal anti-inflammatory (NSAIDS)          |  |
| <input type="checkbox"/> Anti tumour necrosis factor ( <i>Etanercept</i> ) |  |
| <input type="checkbox"/> Other drug or agent _____                         |  |

Unknown

**Other treatment**     No     Yes: \_\_\_\_\_     Unknown

**COMPLICATIONS DUE TO TOXICITY FROM CONVENTIONAL TREATMENT**

No complications

Yes:

- |  |                             |                              |                                  |
|--|-----------------------------|------------------------------|----------------------------------|
| Cataracts  | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Avascular necrosis of femoral head                             | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Severe hypertension  | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Renal insufficiency (>30% increase in creatinine)              | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Severe gastrointestinal (GI) toxicity, specify: _____          | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Hepatic dysfunction (≥3 fold increase in liver function tests) | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Severe gastrointestinal (GI) toxicity, specify: _____          | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Growth delay   | <input type="checkbox"/> No | <input type="checkbox"/> Yes | <input type="checkbox"/> Unknown |
| Other, specify: _____  |                             |                              |                                  |

Unknown

Did severe myelosuppression occur?     No     Yes     Not evaluated     Unknown

## DATE OF HSCT

**DATE OF TRANSPLANT :** ..... - ..... - .....  
yyyy mm dd

**TRANSPLANT TYPE**

Allogeneic: Proceed to STATUS OF DISEASE AT HSCT on page 6

Autologous: Mobilised     No: Proceed to STATUS OF DISEASE AT HSCT on page 6

Yes: Date of 1<sup>st</sup> pheresis/collection: ..... - ..... - .....  
yyyy mm dd

## STATUS OF DISEASE AT MOBILISATION

*Evaluation should be performed <4 weeks prior to mobilisation for stem cell collection.*

### DISEASE STATUS

Number of painful/tender joints : .....  Not evaluated  Unknown  
*(Eular/ACR 28 joint count, which comprises bilateral shoulders, elbows, wrists, MCPs, PIPs and knees. Appendix B.2)*

Number of swollen/effused joints : .....  Not evaluated  Unknown  
*(Eular/ACR 28 joint count, see above)*

Pediatric EPM-Range of motion final score (0-3) : ..... - .....  Not evaluated  Unknown  
*(Appendix B.3)*

Was morning stiffness present?  
 Yes, specify duration: ..... hours ..... minutes  No  Not evaluated  Unknown

Patient's weight: ..... Kg  Not evaluated  Unknown

Patient's height: ..... cm  Not evaluated  Unknown

Patient's weight **one year** prior to time of mobilisation: ..... Kg  Not evaluated  Unknown

Patient's height **one year** prior to time of mobilisation: ..... cm  Not evaluated  Unknown

### HAEMATOLOGICAL VALUES

Haemoglobin ..... g/dL  Not evaluated  Unknown

Erythrocyte sedimentation rate ..... mm/hr  Not evaluated  Unknown

Platelets: ..... (10<sup>9</sup>/l)  Not evaluated  Unknown

WBC ..... (10<sup>9</sup>/l)  Not evaluated  Unknown

DIFFERENTIAL:

Segs: % ..... - .....  Not evaluated  Unknown

Bands: % ..... - .....  Not evaluated  Unknown

Lymphocytes: % ..... - .....  Not evaluated  Unknown

Basophils: % ..... - .....  Not evaluated  Unknown

Monocytes: % ..... - .....  Not evaluated  Unknown

Eosinophils: % ..... - .....  Not evaluated  Unknown

### CLINICAL AND LABORATORY DATA

Serum creatinine ..... - ..... μmol/l  Not evaluated  Unknown

Serum AST ..... (IU/l)  Not evaluated  Unknown

Serum ALT ..... (IU/l)  Not evaluated  Unknown

Serum albumin ..... (g/dl)  Not evaluated  Unknown

Serum alkaline phosphatase ..... (IU/l)  Not evaluated  Unknown

Total serum bilirubin ..... (mg/dl)  Not evaluated  Unknown

C-reactive protein  Normal  Elevated  Not evaluated  Unknown

**AUTOANTIBODIES**

Were tests for autoantibodies done between diagnosis and mobilisation/transplant?  
 No  Yes  Unknown

**Specify antibody:**

Anti-nuclear (ANA)  Negative  Positive  Not evaluated  Unknown  
 Rheumatoid factor  Negative  Positive  Not evaluated  Unknown  
 Other, specify: \_\_\_\_\_  Negative  Positive

**RADIOGRAPHIC EVALUATION**

Were radiographic bone erosions present?  Negative  Positive  Not evaluated  Unknown

Was advanced skeletal age of affected joints noted radiographically?  
 No  Yes  Not evaluated  Unknown

Presence of osteoporotic fractures  Never  Previously but not now  Currently  
 Not evaluated  Unknown

**HEALTH ASSESSMENT QUESTIONNAIRE OR SURVEY COMPLETED**

No  Yes  unknown

**PATIENT'S SELF ASSESSMENT**

**Done Not done Unknown**

Childhood Health Assessment Questionnaire (CHAQ) completed?     
 (see Appendix B.4)

If yes: Specify range of possible scores for the **CHAQ pain** sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the **CHAQ disability** sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the **CHAQ severity** sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**PHYSICIAN'S ASSESSMENT**

**Done Not done Unknown**

Did the physician complete a **Global** Assessment of the patient's state?

If yes: Specify range of possible scores for Physician Rated Global Assessment:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**DISEASE RESPONSE TO THE MOBILISATION**

Response  Transient  No response  Not evaluated

## STATUS OF DISEASE AT HSCT

**Evaluation should be performed <2 weeks prior to conditioning**

### DISEASE STATUS

Number of painful/tender joints : .....  Not evaluated  Unknown  
*(Eular/ACR 28 joint count, which comprises bilateral shoulders, elbows, wrists, MCPs, PIPs and knees. Appendix B.2)*

Number of swollen/effused joints : .....  Not evaluated  Unknown  
*(Eular/ACR 28 joint count, see above)*

Pediatric EPM-Range of motion final score (0-3) : ..... - .....  Not evaluated  Unknown  
*(Appendix B.3)*

Was morning stiffness present?  
 Yes, specify duration: ..... hours ..... minutes  No  Not evaluated  Unknown

Patient's weight **one year** prior to time of transplant: ..... Kg  Not evaluated  Unknown

Patient's height **one year** prior to time of transplant: ..... cm  Not evaluated  Unknown

### HAEMATOLOGICAL VALUES

	Units	Not evaluated	Unknown
Haemoglobin	g/dL	<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Erythrocyte sedimentation rate	mm/hr	<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Platelets:	(10 <sup>9</sup> /l)	<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
WBC	(10 <sup>9</sup> /l)	<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
DIFFERENTIAL:			
Segs: %		<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Bands: %		<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Lymphocytes: %		<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Basophils: %		<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Monocytes: %		<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Eosinophils: %		<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown

### CLINICAL AND LABORATORY DATA

Serum albumin	(g/dl)	<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
Serum alkaline phosphatase	(IU/l)	<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown
C-reactive protein	<input type="checkbox"/> Normal <input type="checkbox"/> Elevated	<input type="checkbox"/> Not evaluated	<input type="checkbox"/> Unknown

### AUTOANTIBODIES

Were tests for autoantibodies done between diagnosis and mobilisation/transplant?  
 No  Yes  Unknown

#### Specify antibody:

Anti-nuclear (ANA)  Negative  Positive  Not evaluated  Unknown  
 Rheumatoid factor  Negative  Positive  Not evaluated  Unknown  
 Other, specify: \_\_\_\_\_  Negative  Positive

**RADIOGRAPHIC EVALUATION**

Were radiographic bone erosions present?  Negative  Positive  Not evaluated  Unknown  
 Was advanced skeletal age of affected joints noted radiographically?  
 No  Yes  Not evaluated  Unknown  
 Presence of osteoporotic fractures  Never  Previously but not now  Currently  
 Not evaluated  Unknown

**HEALTH ASSESSMENT QUESTIONNAIRE OR SURVEY COMPLETED**

No  Yes  unknown

**PATIENT'S SELF ASSESSMENT**

**Done Not done Unknown**

Childhood Health Assessment Questionnaire (**CHAQ**) completed?     
*(see Appendix B.4)*

If yes: Specify range of possible scores for the **CHAQ pain** sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the **CHAQ disability** sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the **CHAQ severity** sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**PHYSICIAN'S ASSESSMENT**

**Done Not done Unknown**

Did the physician complete a **Global** Assessment of the patient's state?

If yes: Specify range of possible scores for Physician Rated Global Assessment:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**FORMS TO BE FILLED IN**

**TYPE OF HSCT**

- AUTOgraft, **proceed to Autograft day 0 form**
- ALLOgraft or Syngeneic graft, **proceed to Allograft day 0 form**
- If  Other : ....., contact the EBMT Central Registry Office for instructions

<h1>DAY 100</h1>	<h1>MED-B</h1> <h2>JUVENILE IDIOPATHIC ARTHRITIS (JIA)</h2>
------------------	---

Unique Identification Code (UIC) ..... (if known)

Date of this report .....  
yyyy mm dd

Hospital Unique Patient Number .....

Initials: ..... (first name(s)\_surname(s))

Date of birth .....  
yyyy mm dd

Sex:  Male  Female  
 (at birth)

Date of the most recent transplant before this follow up: ..... - ..... - .....  
yyyy mm dd

**BEST DISEASE STATUS AT 100 DAYS AFTER TRANSPLANTATION**

*To be completed 100 days post transplant, or at time of death if death occurred <100 days post transplant, or immediately prior to start of high-dose therapy (conditioning) for second transplant if second transplant done <100 days after first transplant.*

Response  Transient  No response  Not evaluated

Date of evaluation .....  
yyyy mm dd

**DISEASE STATUS**

Number of painful/tender joints : .....  Not evaluated  Unknown  
 (Eular/ACR 28 joint count, which comprises bilateral shoulders, elbows, wrists, MCPs, PIPs and knees. Appendix B.2)

Number of swollen/effused joints : .....  Not evaluated  Unknown  
 (Eular/ACR 28 joint count, see above)

Pediatric EPM-Range of motion final score (0-3) : .....  Not evaluated  Unknown  
 (Appendix B.3)

Was morning stiffness present?  
 Yes, specify duration: ..... hours ..... minutes  No  Not evaluated  Unknown

Patient's weight: ..... Kg  Not evaluated  Unknown

Patient's height: ..... cm  Not evaluated  Unknown



**CLINICAL AND LABORATORY DATA**

Erythrocyte sedimentation rate ..... mm/hr    
 C-reactive protein  Normal  Elevated

**RADIOGRAPHIC EVALUATION**

Were radiographic bone erosions present?  Negative  Positive  Not evaluated  Unknown  
 Was advanced skeletal age of affected joints noted radiographically?  
 No  Yes  Not evaluated  Unknown  
 Presence of osteoporotic fractures  Never  Previously but not now  Currently  
 Not evaluated  Unknown

**HEALTH ASSESSMENT QUESTIONNAIRE OR SURVEY COMPLETED**

**PATIENT'S SELF ASSESSMENT**

**Done Not done Unknown**

Childhood HEALTH ASSESSMENT QUESTIONNAIRE (CHAQ) completed?     
 (see Appendix B.4)

If yes: Specify range of possible scores for the CHAQ PAIN sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the CHAQ DISABILITY sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the CHAQ SEVERITY sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**PHYSICIAN'S ASSESSMENT**

**Done Not done Unknown**

Did the physician complete a GLOBAL ASSESSMENT of the patient's state?

If yes: Specify range of possible scores for PHYSICIAN RATED GLOBAL ASSESSMENT:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**FORMS TO BE FILLED IN**

**TYPE OF TRANSPLANT**

- AUTOgraft, **proceed to Autograft day 100 form**
- ALLOgraft or Syngeneic graft, **proceed to Allograft day 100 form**

<b>FOLLOW UP</b>	<b>MED-B JUVENILE IDIOPATHIC ARTHRITIS (JIA)</b>
------------------	--

Unique Identification Code (UIC) ..... (if known)  
 Date of this report .....  
yyyy mm dd  
 Patient following national / international study / trial:  No  Yes  Unknown  
 Name of study / trial .....  
 Hospital Unique Patient Number .....  
 Initials: ..... (first name(s)\_surname(s))  
 Date of birth .....  
yyyy mm dd  
 Sex:  Male  Female  
 (at birth)  
 Date of the most recent transplant before this follow up: .....  
yyyy mm dd

**PATIENT LAST SEEN**

**DATE OF LAST CONTACT OR DEATH:** .....  
yyyy mm dd

**GRAFT VERSUS HOST DISEASE (GvHD) SINCE LAST REPORT**

ANSWER IF PATIENT HAS HAD AN ALLOGRAFT AT ANY TIME

**ACUTE GRAFT VERSUS HOST DISEASE (AGvHD)**

**Maximum grade**  grade 0 (Absent)  grade I  grade II  grade III  grade IV  Not evaluated

If present:  New onset  Recurrent  Persistent

Reason:  Tapering  DLI  Unexplained

Date onset of this episode: .....  Not applicable  
 (if new or recurrent) yyyy mm dd

Stage skin	<input type="checkbox"/> 0	<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
Stage liver	<input type="checkbox"/> 0	<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
Stage gut	<input type="checkbox"/> 0	<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

**Resolution**

No  Yes: Date of resolution: .....  
yyyy mm dd

CIC: ..... Hospital UPN: ..... HSCT Date..... - ..... - .....  
yyyy mm dd  
 Patient Number in EBMT database (if known): .....

ANSWER IF PATIENT HAS HAD AN ALLOGRAFT AT ANY TIME  
**CHRONIC GRAFT VERSUS HOST DISEASE (cGVHD)**

**Presence of cGVHD**

- No  
 Yes:  First episode  
 Recurrence

Date of onset ..... - ..... - .....  
yyyy mm dd

- Present continuously since last reported episode  
 Maximum extent during this period  
 Limited  Extensive  Unknown

Maximum NIH score during this period \_\_\_\_\_  
 Mild  Moderate  Severe  Not evaluated

- Organs affected  Skin  Gut  Liver  Mouth  
 Eyes  Lung  Other, specify .....  Unknown

Resolved: Date of resolution: ..... - ..... - .....  
yyyy mm dd

**LATE GRAFT FAILURE**  No  Yes

**OTHER COMPLICATIONS SINCE LAST REPORT**

PLEASE USE THE DOCUMENT "[DEFINITIONS OF INFECTIOUS DISEASES AND COMPLICATIONS AFTER STEM CELL TRANSPLANTATION](#)" TO FILL THESE ITEMS.

**INFECTION RELATED COMPLICATIONS**

- No complications  
 Yes

Type	Pathogen <i>Use the list of pathogens listed after this table for guidance. Use "unknown" if necessary.</i>	Date <i>Provide different dates for different episodes of the same complication if applicable.</i>
Bacteremia / fungemia / viremia / parasites		
<b>SYSTEMIC SYMPTOMS OF INFECTION</b>		
Septic shock		
ARDS		
Multiorgan failure due to infection		
<b>ENDORGAN DISEASES</b>		
Pneumonia		

Type	Pathogen <i>Use the list of pathogens listed after this table for guidance. Use "unknown" if necessary.</i>	Date <i>Provide different dates for different episodes of the same complication if applicable.</i>
Hepatitis		
CNS infection		
Gut infection		
Skin infection		
Cystitis		
Retinitis		
Other: ..... VOTINCOM		
		yyyy mm dd

**DOCUMENTED PATHOGENS** (Use this table for guidance on the pathogens of interest)

Type	Pathogen	Type	Pathogen
Bacteria	S. pneumoniae	Viruses	HSV
	Other gram positive (i.e.: other streptococci, staphylococci, listeria ...)		VZV
	Haemophilus influenzae		EBV
	Other gram negative (i.e.: E. coli klebsiella, proteus, serratia, pseudomonas ...)		CMV
	Legionella sp		HHV-6
	Mycobacteria sp		RSV
	Other: .....		Other respiratory virus (influenza, parainfluenza, rhinovirus)
			Adenovirus
Fungi	Candida sp	HBV	
	Aspergillus sp	HCV	
	Pneumocystis carinii	HIV	
	Other: .....	Papovavirus	
		Parvovirus	
Parasites	Toxoplasma gondii	Other: .....	
	Other: .....		

**NON INFECTION RELATED COMPLICATIONS**

- No complications
- Yes

<b>Type</b> <i>(Check all that are applicable for this period)</i>	<b>Yes</b>	<b>No</b>	<b>Unknown</b>	<b>Date</b>
Idiopathic pneumonia syndrome	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
VOD	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cataract	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Haemorrhagic cystitis, non infectious	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
ARDS, non infectious	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Multiorgan failure, non infectious	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
HSCT-associated microangiopathy	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Renal failure requiring dialysis	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Haemolytic anaemia due to blood group	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Aseptic bone necrosis	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Other: ..... VOTCOMPS	<input type="checkbox"/>			

*yyyy mm dd*

**GRAFT ASSESSMENT AND HAEMOPOIETIC CHIMAERISM**  
 (ALLOS ONLY)

**Graft loss**  
 No  Yes  Not evaluated

**Overall chimaerism**  Full (*donor* ≥95 %)  Mixed (*partial*)  
 Autologous reconstitution (*recipient* ≥95 %)  Aplasia  
 Not evaluated

INDICATE THE DATE(S) AND RESULTS OF ALL TESTS DONE FOR ALL DONORS.  
 SPLIT THE RESULTS BY DONOR AND BY THE CELL TYPE ON WHICH THE TEST WAS PERFORMED IF APPLICABLE.  
 COPY THIS TABLE AS MANY TIMES AS NECESSARY.

Date of test	Identification of donor or Cord Blood Unit given by the centre	Number in the infusion order (if applicable)	Cell type on which test was performed	% Donor cells	Test used
..... - ..... - ..... yyyy mm dd	.....	<input type="checkbox"/> N/A	<input type="checkbox"/> BM ..... % <input type="checkbox"/> PB mononuclear cells (PBMC) ..... % <input type="checkbox"/> T-cell ..... % <input type="checkbox"/> B-cells ..... % <input type="checkbox"/> Red blood cells ..... % <input type="checkbox"/> Monocytes ..... % <input type="checkbox"/> PMNs (neutrophils) ..... % <input type="checkbox"/> Lymphocytes, NOS ..... % <input type="checkbox"/> Myeloid cells, NOS ..... % <input type="checkbox"/> Other, specify: ..... %		<input type="checkbox"/> FISH <input type="checkbox"/> Molecular <input type="checkbox"/> Cytogenetic <input type="checkbox"/> ABO group <input type="checkbox"/> Other: ..... <input type="checkbox"/> unknown
..... - ..... - ..... yyyy mm dd	.....	<input type="checkbox"/> N/A	<input type="checkbox"/> BM ..... % <input type="checkbox"/> PB mononuclear cells (PBMC) ..... % <input type="checkbox"/> T-cell ..... % <input type="checkbox"/> B-cells ..... % <input type="checkbox"/> Red blood cells ..... % <input type="checkbox"/> Monocytes ..... % <input type="checkbox"/> PMNs (neutrophils) ..... % <input type="checkbox"/> Lymphocytes, NOS ..... % <input type="checkbox"/> Myeloid cells, NOS ..... % <input type="checkbox"/> Other, specify: ..... %		<input type="checkbox"/> FISH <input type="checkbox"/> Molecular <input type="checkbox"/> Cytogenetic <input type="checkbox"/> ABO group <input type="checkbox"/> Other: ..... <input type="checkbox"/> unknown
..... - ..... - ..... yyyy mm dd	.....	<input type="checkbox"/> N/A	<input type="checkbox"/> BM ..... % <input type="checkbox"/> PB mononuclear cells (PBMC) ..... % <input type="checkbox"/> T-cell ..... % <input type="checkbox"/> B-cells ..... % <input type="checkbox"/> Red blood cells ..... % <input type="checkbox"/> Monocytes ..... % <input type="checkbox"/> PMNs (neutrophils) ..... % <input type="checkbox"/> Lymphocytes, NOS ..... % <input type="checkbox"/> Myeloid cells, NOS ..... % <input type="checkbox"/> Other, specify: ..... %		<input type="checkbox"/> FISH <input type="checkbox"/> Molecular <input type="checkbox"/> Cytogenetic <input type="checkbox"/> ABO group <input type="checkbox"/> Other: ..... <input type="checkbox"/> unknown

CIC: ..... Hospital UPN: ..... HSCT Date..... - ..... - .....  
Patient Number in EBMT database (if known): .....  
yyyy mm dd

**SECONDARY MALIGNANCY, LYMPHOPROLIFERATIVE OR MYELOPROLIFRATIVE DISORDER DIAGNOSED**

- Previously reported
  - Yes, date of diagnosis: ..... - ..... - .....  
yyyy mm dd
- Diagnosis:  AML  MDS  Lymphoproliferative disorder  Other .....

IF THE PATIENT HAS RECEIVED AN ALLOGRAFT PRIOR TO THE DIAGNOSIS OF ACUTE LEUKAEMIA, ANSWER THE FOLLOWING QUESTION

Is this secondary malignancy a donor cell leukaemia?  No  Yes  Not applicable

**ADDITIONAL TREATMENT SINCE LAST FOLLOW UP INCLUDING CELL THERAPY**

**Was any additional treatment given for the disease indication for transplant**

- No
- Yes: Start date of the additional treatment since last report: ..... - ..... - .....  
yyyy mm dd
- Unknown

*-Cell therapy*

Did the disease treatment include additional cell infusions (*excluding a new HSCT*)

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

*A boost is an infusion of cells from the same donor without conditioning, in the presence of engraftment (neutrophils > 5 x 10e9), with the same donor being present in a proportion higher than 10%*

Is this cell infusion an autologous boost?  No  Yes

⇒ If cell infusion is not a boost, please complete **CELLULAR THERAPY** on the following page

CIC: ..... Hospital UPN: ..... HSCT Date..... - ..... - .....  
 Patient Number in EBMT database (if known): .....  
 yyyy mm dd

**CELLULAR THERAPY**

One cell therapy regimen is defined as any number of infusions given within 10 weeks for the same indication. If more than one regimen of cell therapy has been given since last report, copy this section and complete it as many times as necessary.

Date of first infusion: .....  
 yyyy mm dd

Disease status before this cellular therapy  CR  Not in CR  Not evaluated  Unknown

**Source of cells:**  Allo  Auto  
 (check all that apply)

**Type of cells** (check all that apply)

- Donor lymphocyte infusion (DLI)
- Mesenchymal cells
- Fibroblasts
- Dendritic cells
- NK cells
- Regulatory T-cells
- Gamma/delta cells
- Other .....
- Unknown

Number of cells infused by type	
Nucleated cells (/kg*) (DLI only)	..... x 10 <sup>8</sup> <input type="checkbox"/> Not evaluated <input type="checkbox"/> unknown
CD 34+ (cells/kg*) (DLI only)	..... x 10 <sup>6</sup> <input type="checkbox"/> Not evaluated <input type="checkbox"/> unknown
CD 3+ (cells/kg*) (DLI only)	..... x 10 <sup>6</sup> <input type="checkbox"/> Not evaluated <input type="checkbox"/> unknown
Total number of cells infused	
All cells (cells/kg*) (non DLI only)	..... x 10 <sup>6</sup> <input type="checkbox"/> Not evaluated <input type="checkbox"/> unknown

Chronological number of this cell therapy for this patient .....

**Indication** (check all that apply)

- Planned/protocol
- Prophylactic
- Treatment of GvHD
- Loss/decreased chimaerism
- Other, specify .....
- Treatment for disease
- Mixed chimaerism
- Treatment viral infection
- Treatment PTLD, EBV lymphoma

**Number of infusions within 10 weeks** .....  
 (count only infusions that are part of same regimen and given for the same indication)

**Acute Graft Versus Host Disease** (after this infusion but before any further infusion / HSCT):

- Maximum grade  grade 0 (absent)  grade 1  grade 2  
 grade 3  grade 4  present, grade unknown



**ADDITIONAL DISEASE TREATMENT FOR JIA**

- No Proceed to FIRST EVIDENCE OF DISEASE WORSENING SINCE LAST HSCT  
 Yes:  Preemptive / preventive (*planned before the transplant took place*)  
 For relapse / progression or persistent disease (*not planned*)

**Date started** ..... - ..... - .....  
yyyy mm dd

**Drugs or agents:**

- No  
 Yes, mark appropriate box(es)  
     Cyclophosphamide   
     Cyclosporin-A   
     Methotrexate   
     Corticosteroids   
 Non-steroidal anti-inflammatory (NSAIDS)   
 Anti tumour necrosis factor (*Etanercept*)   
 Other drug or agent  \_\_\_\_\_  
 Unknown

**Other treatment**  No  Yes: \_\_\_\_\_  Unknown

**FIRST EVIDENCE OF DISEASE WORSENING SINCE LAST HSCT**

**EVIDENCE OF DISEASE ACTIVITY**

- Previously reported  
 No  
 Yes; date first noted: ..... - ..... - .....  
yyyy mm dd  
 Continuous worsening since HSCT

**LAST DISEASE AND PATIENT STATUS**

**DISEASE STATUS**

*Fill in this section only if the evaluation has been performed less than 2 weeks prior to the DATE OF LAST CONTACT OR DEATH in this form.*

- Number of painful/tender joints : .....  Not evaluated  Unknown  
*(Eular/ACR 28 joint count, which comprises bilateral shoulders, elbows, wrists, MCPs, PIPs and knees. Appendix B.2)*
- Number of swollen/effused joints : .....  Not evaluated  Unknown  
*(Eular/ACR 28 joint count, see above)*
- Pediatric EPM-Range of motion final score (0-3) : .....  Not evaluated  Unknown  
*(Appendix B.3)*
- Was morning stiffness present?  
 Yes, specify duration: ..... hours ..... minutes  No  Not evaluated  Unknown
- Patient's weight: ..... Kg  Not evaluated  Unknown  
 Patient's height: ..... cm  Not evaluated  Unknown

CIC: ..... Hospital UPN: ..... HSCT Date..... - ..... - .....  
yyyy mm dd  
 Patient Number in EBMT database (if known): .....

**CLINICAL AND LABORATORY DATA**

	Units	Not evaluated	Unknown
Erythrocyte sedimentation rate .....	mm/hr	<input type="checkbox"/>	<input type="checkbox"/>
C-reactive protein	<input type="checkbox"/> Normal <input type="checkbox"/> Elevated	<input type="checkbox"/>	<input type="checkbox"/>

**RADIOGRAPHIC EVALUATION**

Were radiographic bone erosions present?  Negative  Positive  Not evaluated  Unknown  
 Was advanced skeletal age of affected joints noted radiographically?  
 No  Yes  Not evaluated  Unknown  
 Presence of osteoporotic fractures  Never  Previously but not now  Currently  
 Not evaluated  Unknown

**SURVEYS COMPLETED**

No  
 Yes *Only if the surveys have been performed less than 2 weeks prior to the DATE OF LAST CONTACT OR DEATH in this form.*  
 unknown

**PATIENT'S SELF ASSESSMENT**

	Done	Not done	Unknown
Childhood HEALTH ASSESSMENT QUESTIONNAIRE (CHAQ) completed? <i>(see Appendix B.4)</i>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If yes: Specify range of possible scores for the CHAQ PAIN sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the CHAQ DISABILITY sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

Specify range of possible scores for the CHAQ SEVERITY sub-scale:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**PHYSICIAN'S ASSESSMENT**

	Done	Not done	Unknown
Did the physician complete a GLOBAL ASSESSMENT of the patient's state?	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If yes: Specify range of possible scores for PHYSICIAN RATED GLOBAL ASSESSMENT:

Patient's score: ..... - .....  
 Worst possible score: ..... - .....  
 Best possible score: ..... - .....

**PREGNANCY AFTER HSCT**

Has patient or partner become pregnant after this HSCT?  
 No  
 Yes: Did the pregnancy result in a live birth?  No  Yes  Unknown  
 Unknown

**SURVIVAL STATUS**

- Alive
- Dead

**PERFORMANCE SCORE** (if alive)

- Type of score used**
- Karnofsky
  - Lansky
- SCORE**
- 100 (Normal, NED)
  - 90 (Normal activity)
  - 80 (Normal with effort)
  - 70 (Cares for self)
  - 60 (Requires occasional assistance)
  - 50 (Requires assistance)
  - 40 (Disabled)
  - 30 (Severely disabled)
  - 20 (Very sick)
  - 10 (Moribund)
- Not evaluated
  - Unknown

**MAIN CAUSE OF DEATH** (check only one main cause)

- Relapse or progression / persistent disease
- Secondary malignancy (including lymphoproliferative disease)
- HSCT related cause
- Cell therapy (non HSCT) Related Cause (if applicable)
- Other: .....
- Unknown

**Contributory Cause of Death** (check as many as appropriate):

	Yes	No	Unknown
GvHD (if previous allograft)	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Interstitial pneumonitis	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Pulmonary toxicity	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Infection	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
bacterial	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
viral	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
fungal	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
parasitic	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Rejection / poor graft function	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
History of severe Venous Occlusive disorder (VOD)	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Haemorrhage	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Cardiac toxicity	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Central nervous system toxicity	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Gastro intestinal toxicity	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Skin toxicity	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Renal failure	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Multiple organ failure	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

Other: .....

**ADDITIONAL NOTES IF APPLICABLE**

**COMMENTS** .....

.....

.....

**IDENTIFICATION & SIGNATURE**