

PATIENT INFORMATION

1. PATIENT IDENTIFICATION

Hospital:	Name id (optional):
UPN:	DOB: __/__/__
Usual residence:	Sex:
Postcode:	CIBMTR ID (CRID):
Race:	AID:
Indigenous status:	
Patient consent:	

Cell Therapy PRE-INFUSION

1. PATIENT IDENTIFICATION

Referral centre:
 Referring doctor:
 Date of first referral for cell therapy: __/__/__

2. CELL THERAPY

Participating in CT clinical trial: Y | N
 If yes: Corporate Investigator initiated other _____
 Study id number:
Complete copies of above questions if on multiple trials
 If no, reason why not in clinical trial:
 Institutional guidelines Hospital exemption
 Compassionate use
 Product funding: Clinical Trial MBS MTOP Self-funded

3. PRIOR CELL THERAPY (CT)

This is first course of cell therapy (non HCT): Y | N | Unk
 If no: reported to: ANZTCT CIBMTR EBMT
 Number prior CTs: ____ Date of CT: __/__/__
 Where performed: Indication:
 Cell source(s): Auto Allo-unrelated Allo-related
Complete copies of above questions if >1 prior CT

4. PRIOR TRANSPLANT (HCT)

Received prior HCT: Y | N | Unk
 If yes, reported to: ANZTCT CIBMTR EBMT
 Prior HCT date: __/__/__ Where performed:
 HCT type: Auto Allo-unrelated Allo-related
Complete copies of above questions if >1 prior HCT

5. PRODUCT IDENTIFICATION

Product/s (this course) genetically modified: Y | N
 Donor type: Autologous Allo-unrelated Allo-related
 If related, donor relation:
 Same donor used for prior CT/HCT: Y | N | Unk | NA
 Donor age: Donor sex:
 Unrelated donor: GRID:
 Donor Registry:
 Donor country:
 Number of products: (per protocol) as part of this course of CT: ____
Complete copies of above questions if > 1 donor used
 Product name: _____
 Date of product request: __/__/__
 Date manufacturing started: __/__/__
 Final product ready for shipping: __/__/__
 Final product shipped: __/__/__
 Date receipt of product: __/__/__
 Planned setting of infusion: Inpatient Outpatient
 Actual setting of infusion: Inpatient Outpatient

Cell Therapy PRE-INFUSION (continued)
6. PLANNED HCT

Subsequent HCT planned as part of protocol: Y | N

 Subsequent HCT type: Autologous Allogeneic

Circumstance for subsequent HCT:

- Regardless of response to cell therapy
 Only if responds to cell therapy
 Only if fails or incomplete response

7. INDICATION

Indication for cell therapy:

-
- Malignant haematological disorder -> complete disease forms

-
- Other, specify:

If other indication, Date of diagnosis: __/__/__

8. BRIDGING THERAPY

Bridging therapy was given prior to CT infusion: Y | N

If yes, Date started: __/__/__

9. LYMPHODEPLETING THERAPY

Lymphodepleting therapy given prior infusion: Y | N

Drug	Total dose/mg*	Date started	Dose reduction, % and reason
		__/__/__	
		__/__/__	
		__/__/__	

* Total dose = daily dose x number of days

10. TOXICITY PROPHYLAXIS

CRS prophylaxis agents given:

Neurotoxicity prophylaxis agents given:

11. LAB ASSESSMENTS PRIOR TO LYMPHODEPLETION

Date of complete blood count: __/__/__

	Value	Units
WBC		x 10 ⁹ /L
Neutrophils x 10 ⁹ /L		x 10 ⁹ /L
Lymphocytes x 10 ⁹ /L		x 10 ⁹ /L
Haemoglobin g/L		g/L
Haematocrit %		%

RBC transfused ≤ 30 days prior: Y | N

Platelets x 10 ⁹ /L		x 10 ⁹ /L
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Platelets transfused ≤ 7 days prior: Y | N | Unk

Growth factor given within 7 days prior (or long-acting growth factors within 14 days): Y | N

	Value	Units	Date of sample
LDH		U/L	
LDH ULN		U/L	
Total serum ferritin		ug/L	
C-reactive protein		mg/L	
C-reactive protein ULN		mg/L	
Serum Creatinine		umol/L	

12. PATIENT ASSESSMENT

Karnofsky/Lansky Score: ____ ECOG: ____

13. COMORBID CONDITIONS

Prior viral exposure/infections: (select from checklist)

Comorbidities (Sorrer et al)

- | | |
|--|--|
| <input type="checkbox"/> Arrhythmia | <input type="checkbox"/> Obesity |
| <input type="checkbox"/> Cardiac | <input type="checkbox"/> Peptic Ulcer |
| <input type="checkbox"/> Cerebrovascular | <input type="checkbox"/> Psychiatric |
| <input type="checkbox"/> Diabetes | <input type="checkbox"/> Pulmonary, mod |
| <input type="checkbox"/> Heart valve dis | <input type="checkbox"/> Pulmonary, severe |
| <input type="checkbox"/> Hepatic, mild | <input type="checkbox"/> Renal, mod/severe -> on dialysis: Y N |
| <input type="checkbox"/> Hepatic, mod/sev | <input type="checkbox"/> Rheumatologic |
| <input type="checkbox"/> Infection | <input type="checkbox"/> Prior malignancy, specify: _____ |
| <input type="checkbox"/> Inflammatory bowel disorder | <input type="checkbox"/> Other comorbidity: _____ |

Cell Therapy PRODUCT
1. PRODUCT SOURCE

Date product collected (leukapheresis): __/__/__
 Tissue source: marrow | peripheral blood | other: _____
 Cell type:
 Lymphocytes unselected | CD4+ | CD8+ | other: _____
 Where manufactured / processed:
 Novartis | Kite pharma | Celgene | Janssen |
 Cell processing lab on site | other: _____

2. AUTOLOGOUS PRODUCT

Method of collection:
 BM aspirate | Leukapheresis | other: _____
 Number of collections: _____

3. CELL MANIPULATION -

Not required for commercial products

Cells selected /modified/engineered: Y | N
 Portion manipulated: Entire product | Portion
 If portion, unmanipulated portion also infused: Y | N
 Same manipulation method on entire/all portions: Y | N
 Method used: _____

Complete following if genetically manipulated:

Transfection -> Viral transduction | Non-viral transfection
 Gene editing -> specify gene
 Cells engineered to express a non-native protein: Y | N
 ->T-cell receptor | CAR | Suicide gene, specify
 Other genetic manipulation

Manipulated to recognize specific target/antigen:
 If yes, specify target: _____

4. CELL PRODUCT ANALYSIS

Not required for commercial products

Transfection efficiency performed (genetically engineered cells):
 If yes: Date performed: __/__/__
 Transfection efficiency % _____ target achieved: Y | N
 Viability of cells performed: Y | N | Unk
 If yes: Date performed: __/__/__
 Viability of cells % _____
 Method: 7-AAD | Propidium iodide | Other

5. OUT OF SPECIFICATION

Commercial products only

Product is out of specification: Y | N | Unk
 If yes, reason: _____

6. PRODUCT INFUSION

Total number planned infusions of this product: _____
 (number of infusions specified in the protocol)

Cell Therapy INFUSION
1. CELL PRODUCT IDENTIFIERS

Cell product ID _____ (e.g. Kite Konnect)
 ISBT DIN number _____
 Batch number _____ (Kymriah, Carvykti)
 Lot number _____ (Yescarta, Tecartus)

If Product was not infused

Reason why not infused:
 Disease progression | Comorbidities | Other: _____

2. INFUSION

Date of infusion: __/__/__
 Entire product volume infused: Y | N
 If no, reserved portion fate:
 Discarded | Cryopreserved | other, specify
 Route of infusion: IV | other, specify

3. CELL DOSES

Recipient weight /kg _____
 Recipient height /cm _____
 Report total number of cells given (not cells per kg)
 Total number of cells _____ x 10 ____
 Lymphocytes unselected _____ x 10 ____
 CD4+ lymphocytes _____ x 10 ____
 CD8+ lymphocytes _____ x 10 ____
 Other, specify cell type and dose _____

4. CONCOMITANT THERAPY

Recipient receive concomitant therapy: Y | N
 If yes, specify drugs: _____
 When given: Simultaneous | Post cell therapy | Unknown