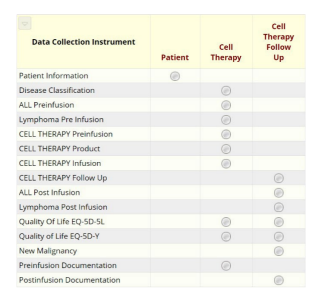


ANZTCT Cell Therapy – Critical Fields

The following list of data fields are the minimum requirement for reporting to the Therapeutic Goods Administration (TGA). Reporting is mandatory based on the approved usage of the Car-T commercial product.

A separate mandatory / critical field list, for FACT accreditation centres will be available in a separate document and will be more comprehensive list.

The critical fields are grouped into Data Collection instruments:



Reports in REDCap

You can access the TGA Critical Fields Reports in REDCap.



Detailed documentation and PDF forms can be found on the ANZTCT website: https://anztct.org.au/registry/data-management-resources/



| Field | Guideline |
|----------------------------------------------------------|------------------------------------------------------------------|
| Patient Information | |
| Current Hospital | |
| Patient UPN | |
| Sex | |
| Date of birth | |
| Country of residence, State (Aust), Postcode | Australian and New Zealand patients |
| Patient consent | / tustialian and non-zoalana patients |
| Survival Status | |
| Date of latest contact | |
| Primary cause of death | Displays if deceased |
| Contributing causes of death | Displays ii deceased |
| Disease Classification | |
| | |
| Date of diagnosis | A |
| Primary Disease for HCT / Cellular Therapy | Associated questions will display when option is selected |
| ACUTE LYMPHOBLASTIC LEUKAEMIA | |
| ALL Classification | |
| CNS disease: | |
| Did recipient have CNS leukaemia | |
| Disease Status at Infusion | |
| In CR by flow cytometry: Y N | if in CR |
| Date assessed | |
| LYMPHOMA | |
| NHL B-cell classification | |
| Transformed from CLL Transformed from different lymphoma | |
| Prior histology | if transformed |
| PET (or PET/CT) scan performed Y N | prior to preparative regimen / infusion |
| PET (or PET/CT) scan positive? | Positive lymphoma involvement at any disease site |
| Deauville Scale | Report the highest score if there are multiple values. |
| Disease Status prior Infusion | |
| Date assessed | |
| Total number of lines of therapy received | |
| ALL Pre Infusion | |
| Treatment was given: Y N | |
| Treatment 1, 2, 3 given: Y N | each line reported separately |
| Therapy type | e.g. induction, bridging to cell infusion |
| Intrathecal therapy Y N | |
| Systemic therapy Y N | |
| Specify systemic agents | |
| Radiation therapy Y N | |
| Cell therapy Y N | |
| Flow cytometry performed: YIN | |
| Tiow cytometry periorinea. The | |



| Field | Guideline |
|-------------------------------------------------------------------------|---------------------------------------|
| BM / blood: date, %disease detected | |
| Lymphoma Pre Infusion | |
| Section 2 – Laboratory values at Diagnosis - IPU Score | |
| LDH | |
| LDH upper normal limit | |
| Section 3 – Nodal and Organ Involvement at Diagnosis - IPI S | Score |
| Extranodal or splenic involvement: Y N Sites of involvement | |
| | |
| Stage of organ involvement ECOG Score | |
| | |
| Section 7 – Disease Treatment Prior to Infusion | |
| Was therapy given: Y N | |
| 1 st (2 nd , 3 rd) line therapy given | each line reported separately |
| Systemic therapy Y N | |
| Specify regimen | given as part of this line of therapy |
| Intrathecal therapy Y N | |
| Radiation therapy Y N | |
| Surgery Y N | |
| Cellular therapy (e.g. CAR-T cells) Y N | |
| This line of therapy given as bridging to cell infusion? | |
| CELL THERAPY Pre Infusion | |
| Referral Centre | |
| Referring doctor | |
| Date of first referral | |
| Section 2 - Cell Therapy | |
| Product Funding | |
| Section 3 – Prior Cell Therapy | |
| First cell therapy for this patient | |
| If "No", number, date of prior cell therapies | |
| Section 4 – Prior Transplant HCT | |
| Received prior HCT | |
| Prior HCT date | |
| Section 5 – Product Identification | |
| Name of product | Required for field branching in form |
| Date of product request | Date of order placed in system |
| Date of manufacturing start | |
| Final product ready for shipping | Gilead |
| Final product shipped | Gilead |
| Date receipt of product | Date of product arrival at centre |
| Actual setting of infusion | Gilead |
| Section 6 – Planned HCT | |
| Subsequent HCT planned part of protocol | |
| Subsequent HCT type | |
| Section 8 – Disease Assessment | |
| | |



| Field | Guideline | |
|------------------------------------------------------------|---------------------------------------------------------------------------------------------------|--|
| Bridging therapy given prior to cell therapy | | |
| Section 9 – Lymphodepleting Therapy | | |
| Lymphodepleting therapy given Y N | | |
| Drug, total dose and units, date started | e.g. NHL: Flu 75mg/m2 + Cy 750mg/m2 ALL: Flu 120mg/m2 + Cy 1000mg/m2 (unless dose modified) | |
| Dose reduction | If "Yes", % dose reduction & Reason dose reduction | |
| Section 10 – Patient Assessment | | |
| Karnofsky / Lansky Score | | |
| ECOG prior to cell therapy | | |
| Section 11 - Comorbid Conditions | | |
| Co-existing diseases or organ impairment present (HCT-CI)? | If "Yes", comorbidities | |
| CELL THERAPY Product | | |
| Name of product | Required for field branching in form | |
| Date product collected | e.g. apheresis date | |
| Tissue source | | |
| Cell type | | |
| Product is out of specification | | |
| Reason out of spec | | |
| Section 6 - Product Infusion | | |
| Total number planned infusions of this product | | |
| CELL THERAPY Infusion | | |
| Name of product | Required for field branching in form | |
| Product was infused | | |
| Reason why not infused | | |
| Section 1 – Cell Product Identifiers | | |
| Batch number | Kymriah | |
| Lot number | Yescarta | |
| Section 2 – Infusion | | |
| Date of infusion | | |
| Section 3 – Cell Doses | | |
| Recipient weight | | |
| Total number of cells / x10^_ | | |
| CELL THERAPY Follow Up | | |
| Hospital | | |
| UPN | | |
| Date of Cell Therapy infusion | | |
| Follow up period | 30 d / 100 d / 6 mth / annual | |
| Section 2 – Survival | | |
| Date of actual contact | | |
| Survival status | | |
| Cause of death | Report on the Patient Information Page | |
| Section 3 – Subsequent Cellular Infusions | | |



| Field | Guideline |
|------------------------------------------------------------------|---------------------------------------------|
| New course of cell therapy given since last report: Y N | complete new Pre-infusion form |
| Reason given | |
| HCT given since last report | |
| Date of HCT | |
| Initial neutrophil recovery | and associated questions |
| Initial platelet recovery | and associated questions |
| Section 7 – Current Haematology Values | |
| Neutrophils x10^9/L | |
| Haemoglobin g/L | |
| RBC transfused 30 days prior | |
| Platelets x10^9/L | |
| Platelets transfused 7 days prior | |
| Section 8 – New Malignancy | |
| New malignancy diagnosed: Y N | |
| Section 11 – CRS | |
| Cytokine Release Syndrome (CRS) Y N | If 'Yes', all associated questions required |
| Section 12 – Toxicities - Neurotoxicity | |
| Neurotoxicity Y N | If 'Yes', all associated questions required |
| Section 13 – Other Toxicities | |
| Hypogammaglobulinemia Y N | If 'Yes', all associated questions required |
| Tumour lysis syndrome: Y N | |
| Section 14 – Grade 3 or 4 Toxicities | |
| Developed grade 3 organ toxicity Y N | If 'Yes', all associated questions required |
| Developed grade 4 organ toxicity Y N | if 'Yes', all associated questions required |
| Section 16 – Infection | |
| Developed clinically significant infection since last report Y N | If 'Yes', all associated questions required |
| Section 17 – Hospitalisation | |
| Hospital admission Y N | If 'Yes', all associated questions required |
| Section 18 – High Cost Medication Usage | |
| High cost medications used | |
| Section 19 – Functional Status | |
| Recipient/female partner pregnant in this reporting period Y N | If 'Yes', all associated questions required |
| ALL Post Infusion | |
| Hospital | |
| UPN | |
| Date of Cell therapy infusion | |
| Follow up period | 30 d / 100 d / 6 mth / annual |
| Section 1 – Best response to HCT or Cell Therapy | |
| Best response to Cellular Therapy | |
| Date of best response or previously reported | |
| Tests performed at time of best response | |
| Molecular testing | If 'Yes', all associated questions required |
| Flow cytometry | If 'Yes', all associated questions required |



| Field | Guideline |
|-------------------------------------------------------------|-----------------------------------------------|
| Cytogenetics tested | If 'Yes', all associated questions required |
| Section 3 – Disease detection since last report | |
| Disease was detected by any assessment method: Y N If 'Yes' | |
| Molecular testing | If 'Yes', all associated questions required |
| Flow cytometry | If 'Yes', all associated questions required |
| Cytogenetic testing | If 'Yes', all associated questions required |
| Lymphoma Post Infusion | |
| Hospital | |
| UPN | |
| Date of Cell therapy infusion | |
| Follow up period | 30 d / 100 d / 6 mth / annual |
| Section 1 – Best response to HCT / Cell Therapy | |
| Best response by PET (metabolic) criteria since last report | |
| Date assessed | |
| Section 3 - Disease Relapse or Progression since last repor | t |
| Relapse or progression occurred | |
| Disease detected by: | |
| Molecular testing | |
| Date sample | |
| Cytogenetic testing: | |
| FISH, date | |
| Karyotyping, date | |
| Radiological assessment Date assessed | |
| Section 5 – Disease status at time of evaluation | |
| Current disease status by PET (metabolic) criteria | |
| Deauville Score at 12 months | |
| Date assessed | |
| If 'Not assessed', Reason not assessed | |
| New Malignancy | |
| Hospital | |
| UPN | |
| Date of Cell therapy infusion | |
| Follow up period | 30 d / 100 d / 6 mth / annual |
| New Malignancy details | |
| New malignancy diagnosis | |
| Date of diagnosis | |
| Quality of Life EQ-5D / EQ-5D-Y | |
| Quality Of Life EQ-5D-5L | At Infusion / 30 d / 100 day / 6 mth / annual |