



ID: 347 v.5 Endorsed

Patients with leukaemia should be considered for inclusion into clinical trials. Link to ALLG website and ANZCTR website.

The anticancer drug(s) in this protocol <u>may</u> have been included in the ADDIKD guideline. Dose recommendations in kidney dysfunction have yet to be updated to align with the ADDIKD guideline. Recommendations will be updated once the individual protocol has been evaluated by the reference committee. For further information refer to the ADDIKD guideline. To assist with calculations, use the <u>eviQ Estimated Glomerular Filtration Rate (eGFR) calculator</u>.

International Consensus Guideline for Anticancer Drug Dosing in Kidney Dysfunction (ADDIKD)

2022

Click here



Related pages:

· Acute myeloid leukaemia FLAG (fludarabine cytarabine and filgrastim)

Treatment schedule - Overview

Drug	Dose	Route	Day
Filgrastim	5 micrograms/kg	Subcut	0 to 5 and continue daily until neutrophil recovery
iDArubicin	10 mg/m ²	IV	1 to 3
Fludarabine	30 mg/m ²	IV infusion	1 to 5
Cytarabine (Ara-C)	2,000 mg/m ²	IV infusion	1 to 5

Cycles: 1 or 2. Upon recovery, a second cycle may be given (generally only if the patient has responded to cycle 1).

Notes

The scheduling of the drugs in this protocol is important; commence cytarabine administration 4 hours after the start of the fludarabine infusion.¹

Drug status: Fludarabine and Cytarabine: PBS General Schedule

Idarubicin: Restricted Benefit

Filgrastim: PBS authority

Cost: ~ \$3,560 per cycle

Treatment schedule - Detail

The supportive therapies (e.g. antiemetics, premedications, etc.), infusion times, diluents, volumes and routes of administration, if included, are listed as defaults. They may vary between institutions and can be substituted to reflect individual institutional policy.

Antiemetics if included in the treatment schedule are based upon recommendations from national and international guidelines. These are defaults only and may be substituted to reflect individual institutional policy. Select here for recommended doses of alternative antiemetics.

Day 0			
Filgrastim	5 micrograms/kg (Subcut)	inject subcutaneously once daily from day 0 and continue until neutrophil recovery. To be given before chemotherapy on days 1 to 5.	
Day 1 to 3			
Filgrastim	5 micrograms/kg (Subcut)	inject subcutaneously once daily from day 0 and continue until neutrophil recovery. To be given before chemotherapy on days 1 to 5.	
iDArubicin	10 mg/m ² (IV)	over 10 to 15 minutes	
Fludarabine	30 mg/m ² (IV infusion)	in 100 mL sodium chloride 0.9% over 30 minutes	
Cytarabine (Ara-C)	2,000 mg/m ² (IV infusion)	in 500 mL sodium chloride 0.9% over 4 hours. Administer 4 hours after commencing the fludarabine infusion.	
Day 4 and 5			
Filgrastim	5 micrograms/kg (Subcut)	inject subcutaneously once daily from day 0 and continue until neutrophil recovery. To be given before chemotherapy on days 1 to 5.	
Fludarabine	30 mg/m ² (IV infusion)	in 100 mL sodium chloride 0.9% over 30 minutes	
Cytarabine (Ara-C)	2,000 mg/m ² (IV infusion)	in 500 mL sodium chloride 0.9% over 4 hours. Administer 4 hours after commencing the fludarabine infusion.	

Cycles:

1 or 2. Upon recovery, a second cycle may be given (generally only if the patient has responded to cycle 1).

Indications and patient population

• Relapsed or refractory acute myeloid leukaemia in patients younger than 60 years of age (predominant use).

Clinical information

Venous access required	IV cannula (IVC) or central venous access device (CVAD) is required to administer this treatment. Read more about central venous access device line selection
Antiemetics for multi-day protocols	Antiemetic therapy should be administered throughout the duration of the chemotherapy protocol and to cover delayed nausea. The acute and delayed emetic risk of multi-day chemotherapy protocols will overlap depending on the individual drugs and their sequence of administration. More or less antiemetic cover may be required.
	Ensure that patients also have sufficient antiemetics for breakthrough emesis:
	Metoclopramide 10 mg three times a day when necessary (maximum of 30 mg/24 hours, up to 5 days) OR
	Prochlorperazine 10 mg PO every 6 hours when necessary.
	Read more about preventing anti-cancer therapy induced nausea and vomiting

Cumulative lifetime dose of anthracyclines	Cumulative doses should take into account all previous anthracyclines received during a patient's lifetime (i.e. daunorubicin, doxorubicin, epirubicin, idarubicin and mitoxantrone). Criteria for reducing the total anthracycline cumulative lifetime dose include: • patient is elderly • prior mediastinal radiation • hypertensive cardiomegaly • concurrent therapy with high dose cyclophosphamide and some other cytotoxic drugs (e.g. bleomycin, dacarbazine, dactinomycin, etoposide, melphalan, mitomycin and vincristine). Baseline clinical assessments include echocardiogram (ECHO) or gated heart pool scan (GHPS) and electrocardiogram (ECG) evaluation. Patients with normal baseline cardiac function (left ventricular ejection fraction (LVEF) > 50%) and low risk patients require LVEF monitoring when greater than 70% of the anthracycline threshold is reached or if the patient displays symptoms of cardiac impairment. Post-treatment cardiac monitoring is recommended for patients who have received high levels of total cumulative doses of anthracyclines at the clinician's discretion. Read more about cardiac toxicity associated with anthracyclines
Ocular toxicities	Administer corticosteroid eye drops to minimise corneal toxicity from high dose cytarabine. Commence on the day of first dose of cytarabine and continue for at least 72 hours after completion of final cytarabine dose. Read more about ocular toxicities associated with high dose cytarabine
Cytarabine syndrome	Treatment with cytarabine may cause a "cytarabine syndrome" characterised by flu-like symptoms, skin rash and occasionally chest pain.
Cytarabine-induced neurotoxicity	This may occur in patients treated with high-dose cytarabine. Assess cerebellar function prior to each cytarabine dose. Note: an increased risk of cytarabine-induced neurotoxicity has been associated with kidney dysfunction. Read more about neurotoxicity associated with high-dose cytarabine and access the cytarabine cerebellar neurotoxicity assessment chart
Tumour lysis risk	Assess patient for risk of developing tumour lysis syndrome. Read more about prevention and management of tumour lysis syndrome.
Pneumocystis jirovecii pneumonia (PJP) prophylaxis	Read more about prophylaxis of pneumocystis jiroveci (carinii) in cancer patients
Antiviral prophylaxis	Read more about antiviral prophylaxis drugs and doses
Antifungal prophylaxis	Antifungal prophylaxis is recommended. Note: The concurrent use of azole antifungals and anthracyclines should be avoided due to a potential drug interaction and increased toxicities. Commence prophylaxis with an azole antifungal 24 hours after the last dose of anthracycline. ² Read more about antifungal prophylaxis drugs and doses.
Biosimilar drug	Read more about biosimilar drugs on the Biosimilar Awareness Initiative page
Growth factor support	G-CSF (short or long-acting) is available on the PBS for chemotherapy induced neutropenia depending on clinical indication and/or febrile neutropenia risk. Access the PBS website
Irradiated blood components	The use of irradiated of blood components is recommended for patients receiving this treatment. Read more about the indications for the use of irradiated blood components
Blood tests	FBC, EUC, eGFR and LFTs at baseline and prior to each treatment.
Hepatitis B screening and prophylaxis	Routine screening for HBsAg and anti-HBc is recommended prior to initiation of treatment. Prophylaxis should be determined according to individual institutional policy. Read more about hepatitis B screening and prophylaxis in cancer patients requiring cytotoxic and/or immunosuppressive therapy

Vaccinations	Live vaccines are contraindicated in cancer patients receiving immunosuppressive therapy and/or who have poorly controlled malignant disease. Refer to the recommended schedule of vaccination for immunocompromised patients, as outlined in the Australian Immunisation Handbook. Read more about COVID-19 vaccines and cancer.
Fertility, pregnancy and lactation	Cancer treatment can have harmful effects on fertility and this should be discussed with all patients of reproductive potential prior to commencing treatment. There is a risk of foetal harm in pregnant women. A pregnancy test should be considered prior to initiating treatment in females of reproductive potential if sexually active. It is important that all patients of reproductive potential use effective contraception whilst on therapy and after treatment finishes. Effective contraception methods and adequate contraception timeframe should be discussed with all patients of reproductive potential. Possibility of infant risk should be discussed with breastfeeding patients. Read more about the effect of cancer treatment on fertility

Dose modifications

Evidence for dose modifications is limited, and the recommendations made on eviQ are intended as a guide only. They are generally conservative with an emphasis on safety. Any dose modification should be based on clinical judgement, and the individual patient's situation including but not limited to treatment intent (curative vs palliative), the anti-cancer regimen (single versus combination therapy versus chemotherapy versus immunotherapy), biology of the cancer (site, size, mutations, metastases), other treatment related side effects, additional co-morbidities, performance status and patient preferences. Suggested dose modifications are based on clinical trial findings, product information, published guidelines and reference committee consensus. The dose reduction applies to each individual dose and not to the total number of days or duration of treatment cycle unless stated otherwise. Non-haematological gradings are based on Common Terminology Criteria for Adverse Events (CTCAE) unless otherwise specified. Renal and hepatic dose modifications have been standardised where possible. For more information see dosing considerations & disclaimer.

The dose recommendations in kidney dysfunction (i.e.renal impairment) displayed may not reflect those in the ADDIKD guideline and have been included for historical reference only. Recommendations will be updated once the individual protocol has been evaluated by the reference committee, with this version of the protocol then being archived. Clinicians are expected to refer to the ADDIKD guideline prior to prescribing in kidney dysfunction.

International Consensus Guideline for Anticancer Drug Dosing in Kidney Dysfunction (ADDIKD).

Note: all dose reductions are calculated as a percentage of the starting dose.

Haematological toxicity

Dose reductions for haematological toxicity not usually recommended. Discuss with Haematologist.

Renal impairment		
Creatinine clearance (mL/min)		
30 to 50	Reduce fludarabine by 50%	
less than 30	Fludarabine contraindicated	
No specific dose modifications are recommended for cytarabine in renal impairment.		

Note: an increased risk of neurotoxicity has been associated with high-dose cytarabine and a creatinine clearance of less than 60 mL/min.

Hepatic impairment		
Hepatic dysfunction		
Mild Reduce idarubicin by 25%		

Hepatic impairment	
Moderate	Reduce idarubicin by 50%
Severe	Omit idarubicin. Consider alternative regimen
Elevations in liver function tests occur with both standard and high dose cytarabine. Significant liver function abnormalities may require discontinuation or a dose reduction. Interruption of azole antifungal use should also be considered.	

Interactions

Drug interactions in eviQ protocols are under review and being updated to align with current literature. Further site-wide updates and changes will occur in due course. References & Disclaimer

The drug interactions shown below are not an exhaustive list. For a more comprehensive list and for detailed information on specific drug interactions and clinical management, please refer to the specific drug product information and the following key resources:

- MIMS interactions tab (includes link to a CYP-450 table) (login required)
- Australian Medicines Handbook (AMH) interactions tab (login required)
- Micromedex Drug Interactions (login required)
- Cancer Drug Interactions
- Cytochrome P450 Drug Interactions

Cytarabine		
	Interaction	Clinical management
Cytidine deaminase (CDA) inhibitors (e.g. cedazuridine)	Potential increased effect/toxicity of cytarabine due to reduced clearance	Avoid combination or monitor for increased cytarabine effect/toxicity

Fludarabine		
	Interaction	Clinical management
Dipyridamole	Reduced efficacy of fludarabine possible due to inhibition of adenosine uptake	Avoid combination or monitor for decreased clinical response to fludarabine

Idarubicin		
	Interaction	Clinical management
Cardiotoxic drugs (eg. calcium channel blockers, propranolol)	Increased risk of idarubicin-induced cardiotoxicity	Avoid combination or monitor closely for cardiotoxicity

General		
	Interaction	Clinical management
Warfarin	Anti-cancer drugs may alter the anticoagulant effect of warfarin.	Monitor INR regularly and adjust warfarin dosage as appropriate; consider alternative anticoagulant.
Direct oral anticoagulants (DOACs) e.g. apixaban, rivaroxaban, dabigatran	Interaction with both CYP3A4 and P-gp inhibitors /inducers. DOAC and anti-cancer drug levels may both be altered, possibly leading to loss of efficacy or toxicity (i.e. increased bleeding).	Apixaban: avoid concurrent use with strong CYP3A4 and P-gp inhibitors. If treating VTE, avoid use with strong CYP3A4 and P-gp inducers. Rivaroxaban: avoid concurrent use with strong CYP3A4 and P-gp inhibitors. Dabigatran: avoid combination with strong P-gp inducers and inhibitors. If concurrent use is unavoidable, monitor closely for efficacy/toxicity of both drugs.
Digoxin	Anti-cancer drugs can damage the lining of the intestine; affecting the absorption of digoxin.	Monitor digoxin serum levels; adjust digoxin dosage as appropriate.
Antiepileptics	Both altered antiepileptic and anti- cancer drug levels may occur, possibly leading to loss of efficacy or toxicity.	Where concurrent use of an enzyme-inducing antiepileptic cannot be avoided, monitor antiepileptic serum levels for toxicity, as well as seizure frequency for efficacy; adjust dosage as appropriate. Also monitor closely for efficacy of the anti-cancer therapy.
Antiplatelet agents and NSAIDs	Increased risk of bleeding due to treatment related thrombocytopenia.	Avoid or minimise combination. If combination deemed essential, (e.g. low dose aspirin for ischaemic heart disease) monitor for signs of bleeding.
Serotonergic drugs, including selective serotonin reuptake inhibitors (SSRIs e.g. paroxetine) and serotonin noradrenaline reuptake inhibitors (SNRIs e.g. venlafaxine)	Increased risk of serotonin syndrome with concurrent use of 5-HT3 receptor antagonists (e.g. palonosetron, ondansetron, granisetron, tropisetron, dolasetron, etc.)	Avoid combination. If combination is clinically warranted, monitor for signs and symptoms of serotonin syndrome (e.g. confusion, agitation, tachycardia, hyperreflexia). For more information link to TGA Medicines Safety Update
Vaccines	Diminished response to vaccines and increased risk of infection with live vaccines.	Live vaccines (e.g. BCG, MMR, zoster and varicella) are contraindicated in patients on immunosuppressive therapy. Use with caution in patients on non-immunosuppressive therapy. For more information; refer to the recommended schedule of vaccination for cancer patients, as outlined in the Australian Immunisation Handbook

Administration

eviQ provides safe and effective instructions on how to administer cancer treatments. However, eviQ does not provide every treatment delivery option, and is unable to provide a comprehensive list of cancer treatment agents and their required IV line giving set/filter. There may be alternative methods of treatment administration, and alternative supportive treatments that are also appropriate. Please refer to the individual

Day 0

Subcutaneous injection

General patient assessment prior to each day of treatment.

Commence corticosteroid eye drops 24 hours before starting cytarabine. Continue for 72 hours after completion of the last dose of cytarabine.

Filgrastim

Inject subcutaneously ONCE daily, starting on day 0 and continuing until neutrophil recovery.

Days 1 to 3

Safe handling and waste management

Safe administration

General patient assessment prior to each treatment.

Any toxicity grade 2 or greater may require dose reduction, delay or omission of treatment and review by medical officer before recommencing treatment.

Prime IV line(s).

Insert IV cannula or access TIVAD or CVAD.

- · baseline weight
- baseline neurological assessment e.g. daily handwriting assessment that includes the date, time and location.

Pre treatment medication

Verify antiemetics taken or administer as prescribed.

Continue corticosteroid eye drops until 72 hours after completion of the last dose of cytarabine.

Filgrastim

Inject subcutaneously ONCE daily, starting on day 0 and continuing until neutrophil recovery.

Ochemotherapy - Time out

Idarubicin

Administer idarubicin (vesicant):

- over 10 to 15 minutes
 - o via a minibag OR
 - $\circ~$ by IV bolus via a side port of a freely flowing IV infusion
- · ensure vein is patent and monitor for signs of extravasation throughout administration
- flush with ~150 mL of sodium chloride 0.9%
- potential for flare reaction during administration of idarubicin (facial flushing and red streaking along the vein) stop infusion exclude extravasation before continuing at a slower rate of infusion.

Although rare, cardiac arrhythmias may occur during or immediately after idarubicin administration. If sudden onset of dyspnoea, palpitations or irregular pulse occurs, stop administration immediately and obtain urgent medical officer review.

Fludarabine

- administer via IV infusion over 30 minutes
- flush with ~ 50 mL of sodium chloride 0.9%

Administer cytarabine 4 hours AFTER commencing the fludarabine infusion

Cytarabine

Prior to administration:

Ensure corticosteroid eye drops have been administered before starting cytarabine.

Verify that cytarabine neurological assessment has been performed prior to administration of cytarabine:

- if the patient scores 0 then administer cytarabine as charted
- if the patient scores 1 or above, do not administer the cytarabine and immediately notify medical officer.

Administer cytarabine:

- · via IV infusion over 4 hours
- flush with ~50 mL of sodium chloride 0.9%.

Continue safe handling precautions until 7 days after completion of drug(s)

Days 4 and 5

Safe handling and waste management

Safe administration

General patient assessment prior to each treatment.

Any toxicity grade 2 or greater may require dose reduction, delay or omission of treatment and review by medical officer before recommencing treatment.

· daily weight

Pre treatment medication

Verify antiemetics taken or administer as prescribed.

Continue corticosteroid eye drops until 72 hours after completion of the last dose of cytarabine.

Filgrastim

Inject subcutaneously ONCE daily, starting on day 0 and continuing until neutrophil recovery.

Ochemotherapy - Time out

Fludarabine

- administer via IV infusion over 30 minutes
- flush with ~ 50 mL of sodium chloride 0.9%

Administer cytarabine 4 hours AFTER commencing the fludarabine infusion

Cytarabine

Prior to administration:

Ensure corticosteroid eye drops have been administered before starting cytarabine.

Verify that cytarabine neurological assessment has been performed prior to administration of cytarabine:

- if the patient scores 0 then administer cytarabine as charted
- if the patient scores 1 or above, do not administer the cytarabine and immediately notify medical officer.

Administer cytarabine:

- via IV infusion over 4 hours
- flush with ~50 mL of sodium chloride 0.9%.

Remove IV cannula and/or deaccess TIVAD or CVAD.

Continue safe handling precautions until 7 days after completion of drug(s)

Discharge information

Antiemetics

· Antiemetics as prescribed.

Corticosteroid eye drops

• Continue corticosteroid eye drops for at least 72 hours after completion of final cytarabine dose.

Growth factor support

• Arrangements for administration if prescribed.

Prophylaxis medications

• Prophylaxis medications (if prescribed) i.e. tumour lysis prophylaxis, PJP prophylaxis, antifungals, antivirals.

Patient information

• Ensure patient receives patient information sheet.

Side effects

The side effects listed below are not a complete list of all possible side effects for this treatment. Side effects are categorised into the approximate onset of presentation and should only be used as a guide.

Immediate (onset hours to days)				
Bone pain	Bone pain, usually in the lower back or pelvis, associated with G-CSF.			
Extravasation, tissue or vein injury	The unintentional instillation or leakage of a drug or substance out of a blood vessel into surrounding tissue. This has the potential to cause damage to affected tissue. Read more about extravasation management			
Flare reaction	Anthracycline flare reaction is caused by a localised allergic reaction. It is characterised by erythematous vein streaking, urticaria and pruritus which may occur during drug administration and is often associated with too rapid an infusion. Extravasation must be ruled out if flare occurs.			
Nausea and vomiting	Read more about prevention of treatment induced nausea and vomiting			
Neurotoxicity	High dose cytarabine has been associated with acute cerebellar syndrome and diffuse cerebral dysfunction. Read more about neurotoxicity associated with high dose cytarabine			
Ocular toxicities	Reversible corneal toxicity (keratitis), haemorrhagic conjunctivitis, vision loss and other ocular side effects can occur with high dose cytarabine. Corticosteroid eye drops must be administered concurrently with treatment. Read more about ocular toxicities associated with cytarabine			
Red-orange discolouration of urine	Pink/red/orange discolouration of the urine. This can last for up to 48 hours after some anthracycline drugs.			
Taste and smell alteration	Read more about taste and smell changes			

Early (onset days to weeks)	
Neutropenia	Abnormally low levels of neutrophils in the blood. This increases the risk of infection. Any fever or suspicion of infection should be investigated immediately and managed aggressively. Read more about immediate management of neutropenic fever
Thrombocytopenia	A reduction in the normal levels of functional platelets, increasing the risk of abnormal bleeding. Read more about thrombocytopenia
Cytarabine (Ara-C) syndrome	Flu-like symptoms including fever, myalgia and malaise can occur 6 to 12 hours after cytarabine administration. Symptoms generally resolve within 24 hours of completing therapy.
Diarrhoea	Read more about treatment induced diarrhoea
Fatigue	Read more about fatigue
Oral mucositis	Erythematous and ulcerative lesions of the gastrointestinal tract (GIT). It commonly develops following chemotherapy, radiation therapy to the head, neck or oesophagus, and high dose chemotherapy followed by a blood and marrow transplant (BMT). Read more about oral mucositis
Photosensitivity	Increased sensitivity to ultraviolet (UV) light resulting in an exaggerated sunburn-like reaction accompanied by stinging sensations and urticaria.
Skin rash	Anti-cancer drugs can cause a number of changes in the skin with maculo-papular rash the most common type of drug-induced skin reaction. Read more about skin rash

Late (onset weeks to months)				
Anaemia	Abnormally low levels of red blood cells (RBCs) or haemoglobin in the blood. Read more about anaemia			
Alopecia	Hair loss may occur from all parts of the body. Patients can also experience mild to moderate discomfort of the hair follicles, and rarely pain as the hair is falling out. Read more about alopecia and scalp cooling			
Cognitive changes (chemo fog)	Changes in cognition characterised by memory loss, forgetfulness and feeling vague. This is also referred to as 'chemo brain' or 'chemo fog'. Read more about cognitive changes (chemo fog)			
Neurotoxicity	Neurotoxicity related to fludarabine is a rare but potentially serious adverse event characterised by visual disturbances, altered mental state and CNS toxicity. Seizures leading to paralysis or coma have been reported in the literature. Periodic neurologic assessments are recommended.			
Pulmonary toxicity	Pulmonary toxicity may include damage to the lungs, airways, pleura and pulmonary circulation. Read more about pulmonary toxicity associated with anti-cancer drugs			

Delayed (onset months to year	rs)
Cardiotoxicity	Anthracyclines are the most frequently implicated anti-cancer drugs associated with cardiotoxicity, which typically manifests as a reduction in left ventricular ejection fraction (LVEF), cardiomyopathy, or symptomatic CHF. Anthracycline induced cardiotoxicity has been categorised into acute, early-onset chronic progressive and late-onset chronic progressive and is usually not reversible. The risk of clinical cardiotoxicity increases with a number of risk factors including higher total cumulative doses. Read more about cardiac toxicity associated with anthracyclines

Evidence

Cytarabine is one of the most effective drugs used for treatment of acute myeloid leukaemia (AML) and myelodysplastic syndromes (MDS). The administration of fludarabine prior to high dose cytarabine has been shown to increase intracellular

accumulation of the active metabolite, ara-CTP.¹ In addition, G-CSF prior to fludarabine increases the fraction of cells in cycle when they are most vulnerable to ara-C and enhances the incorporation of ara-C into DNA.³ These two aspects provide the rationale for the FLAG protocols. Anthracyclines are known to improve response rates when added to other cytarabine-based regimens. On this basis, Parker and colleagues performed a pilot study adding idarubicin to FLAG for treatment of high-risk AML and MDS patients.⁴ The daily dose of idarubicin given varies from 8 to 12 mg/m² in subsequent reported series.⁴, 5, 6, 7, 8, 9

There have not been any randomised studies comparing FLAG-Ida with FLAG. Virchis and colleagues studied FLAG ± Ida in high-risk AML patients and showed no significant difference in response. In this study, selection of FLAG or FLAG-Ida for each patient was at clinician discretion and FLAG-Ida was restricted to patients less than 65 years. It should be noted that the majority of patients receiving FLAG-Ida in reported series have been less than 60 years old and the protocol is not generally recommended for older patients.

Efficacu

Currently, FLAG-Ida and FLAG protocols are used most commonly as salvage therapy for relapsed and refractory AML patients. There have been several studies of FLAG-Ida in this setting with reported complete response (CR) rates ranging from 52 to 63% range (see table). If complete or partial remission is achieved a further cycle of FLAG-Ida is given^{4, 5, 6} followed by allogeneic stem cell transplantation if possible.

Reference	N =	Age* (years)	CR (%)	TRM (%)	Comments
Parker et al 1997 ⁴	19	44 (range 18 - 72 years)	63	0	- both AML & "high risk MDS" patients - disease de novo (9), relapsed/refractory (7) or secondary (3)
Steinmetz et al 1999 ⁶	57	52	52	23	- relapsed/refractory or secondary AML
Pastore et al 2003 ⁷	46	41	52	7	- relapsed/refractory AML
Virchis et al 2004 ⁵	105	35	59	17	 high risk acute leukaemia or MDS (66 patients were MDS or AML) either FLAG or FLAG-Ida given but no information re: numbers in each group FLAG-Ida restricted to <65 yrs (median age 28 vs 57 in FLAG group)
Yavuz et al 2006 ⁸	34	37	54	18	- relapsed/refractory AML - 22 ALL patients also treated with FLAG-Ida (data not summarised here)
Martin et al 2009 ⁹	23	48	52	35	- a second cohort of 48 pts given FLAG-Ida + gemtuzumab with no improvement in outcomes - G-CSF concurrent with chemotherapy better than sequential (CR 62% vs 29%)
Bergua et al 2016 ¹⁰	221	54	50	N/A	- a second retrospective cohort of 38 pts given FLAG-lda + gemtuzumab was also analysed with no significant difference in outcomes
Westhus et al 2019 ¹¹	132	52	56	16	- relapsed/refractory AML

^{*} Median age in years; CR = complete remission; TRM = treatment related mortality; AML = acute myeloid leukaemia; MDS = myelodysplastic syndrome

Toxicity

FLAG-Ida, like FLAG, is generally well tolerated with myelosuppression the major toxicity. The median time to neutrophil recovery (>0.5 x 10⁹/L) ranges from 16 to 28 days^{4, 5, 7, 8} and median time to platelet count > 20 x 10⁹/L is 18 to 31 days.^{4, 5} Virchis and colleagues did not note any significant difference in recovery times when FLAG-Ida was compared with FLAG.⁵ Febrile neutropenia was virtually universal and fungal infection rates varied from 6 to 17%, where reported.^{5, 8, 9} Non-haematological toxicities were generally mild, and included alopecia, nausea, vomiting and liver function test derangement. Two studies reported neurological adverse events in small numbers of patients.^{4, 5} Reported treatment related mortality rates varied greatly between studies, ranging from 0%⁴ to 35%,⁹ with infection being a leading cause of death.^{5, 6}

References

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- **4** Parker, J. E., A. Pagliuca, A. Mijovic, et al. 1997. "Fludarabine, cytarabine, G-CSF and idarubicin (FLAG-IDA) for the treatment of poor-risk myelodysplastic syndromes and acute myeloid leukaemia." Br.J.Haematol. 99(4):939-944.
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- 6 Steinmetz, H. T., A. Schulz, P. Staib, et al. 1999. "Phase-II trial of idarubicin, fludarabine, cytosine arabinoside, and filgrastim (Ida-FLAG) for treatment of refractory, relapsed, and secondary AML." Ann Hematol 78(9):418-425.
- **7** Pastore, D., G. Specchia, P. Carluccio, et al. 2003. "FLAG-IDA in the treatment of refractory/relapsed acute myeloid leukemia: single-center experience." Ann Hematol 82(4):231-235.
- **8** Yavuz, S., S. Paydas, U. Disel, et al. 2006. "IDA-FLAG regimen for the therapy of primary refractory and relapse acute leukemia: a single-center experience." Am J Ther 13(5):389-393.
- 9 Martin, M. G., K. M. Augustin, G. L. Uy, et al. 2009. "Salvage therapy for acute myeloid leukemia with fludarabine, cytarabine, and idarubicin with or without gemtuzumab ozogamicin and with concurrent or sequential G-CSF." Am J Hematol 84(11):733-737.
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- 11 Westhus, J., R. Noppeney, U. Dührsen, et al. 2019. "FLAG salvage therapy combined with idarubicin in relapsed/refractory acute myeloid leukemia." Leuk Lymphoma 60(4): 1014-1022.

History

Version 5

Date	Summary of changes
22/09/2020	Biosimilar drug added to clinical information. Version number changed to v.5
23/10/2020	Protocol reviewed at the Haematology Reference Committee meeting with further discussions continuing via email.
15/06/2022	Protocol reviewed electronically by Haematology Reference Committee. Evidence updated and antifungal prophylaxis clinical information block updated. For review in 2 years.
25/07/2023	Neurotoxicity added to "Late" category of Side effects section.

Version 4

Date	Summary of changes
23/05/2008	Protocol review and additional information added to increase comprehensiveness of protocol
08/03/2010	Review, new dose modifications and transferred to eviQ

Date	Summary of changes
17/09/2010	Full protocol review at Haematology Reference Committee meeting. Evidence section updated; indication specified for patients less than 60 years.
23/11/2011	New format to allow for export of protocol information Protocol version number changed to <i>V.2</i> Antiemetics and premedications added to the treatment schedule Additional Clinical Information, Key Prescribing table and Key Administration table combined into new section titled Clinical Considerations Drug specific information placed behind the drug name link
04/08/2014	Protocol reviewed by email survey. Added link to ALLG and ANZCTR with statement 'Patients with AML should be considered for inclusion into clinical trials'. Changed the wording in the drug schema for filgrastim to read 0 and continue until neutrophil recovery (as per the 2010 RC minutes). Next review in 2 years.
11/02/2016	Standard review, updated drug costs, review in 5 years
31/05/2017	Transferred to new eviQ website. Version number change to V.3.
14/12/2017	The days on which filgrastim is administered was adjusted in the overview and detail treatment schedules to mention days 1 to 5. Version number increased to V.4.
10/10/2019	Clinical information updated with PBS expanded indications for G-CSF.

The information contained in this protocol is based on the highest level of available evidence and consensus of the eviQ reference committee regarding their views of currently accepted approaches to treatment. Any clinician (medical oncologist, haematologist, radiation oncologist, medical physicist, radiation therapist, pharmacist or nurse) seeking to apply or consult this protocol is expected to use independent clinical judgement in the context of individual clinical circumstances to determine any patient's care or treatment. While eviQ endeavours to link to reliable sources that provide accurate information, eviQ and the Cancer Institute NSW do not endorse or accept responsibility for the accuracy, currency, reliability or correctness of the content of linked external information sources. Use is subject to eviQ's disclaimer available at www.eviQ.org.au

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19 Sep 2023

Patient information - Acute myeloid leukaemia (AML) - FLAG-Ida (fludarabine, cytarabine, idarubicin, filgrastim)



Patient's name:

Your treatment

The treatment schedule below explains how the drugs for this treatment are given.

FLAG - Ida (fludarabine, cytarabine, idarubicin and filgrastim)					
This treatment may be given for up to two cycles. Your doctor will advise you of the number of treatments you will have.					
Day	Treatment How it is given How long it takes				
0 (and continue until white cell count increases)	Granulocyte Colony Stimulating Factor (G-CSF)	By injection under the skin	About 5 minutes		
1 to 3	Idarubicin (EYE-da-ROO-bi-sin)	By a drip into a vein	About 15 minutes		
1 to 5	Fludarabine (Flu-dara-been)	By a drip into a vein	About 30 minutes		
	Cytarabine (sye-TARE-a-been)	By a drip into a vein	About 4 hours		

When to get help

Anticancer drugs (drugs used to treat cancer) can sometimes cause serious problems. It is important to get medical help immediately if you become unwell.

IMMEDIATELY go to your nearest hospital Emergency Department, or contact your doctor or nurse if you have any of the following at any time:	Emergency contact details Ask your doctor or nurse from your treating team who to contact if you have a problem
 a temperature of 38°C or higher chills, sweats, shivers or shakes shortness of breath uncontrolled vomiting or diarrhoea pain, tingling or discomfort in your chest or arms you become unwell. 	Daytime: Night/weekend: Other instructions:

During your treatment immediately tell the doctor or nurse looking after you if you get any of the following problems:

- leaking from the area where the drugs are being given
- pain, stinging, swelling or redness in the area where the drugs are being given or at any injection sites
- a skin rash, itching, feeling short of breath, wheezing, fever, shivers, or feeling dizzy or unwell in any way (allergic reaction).

Other information about your treatment

Changes to your dose or treatment delays

Sometimes a treatment may be started at a lower dose or the dose needs to be changed during treatment. There may also be times when your treatment is delayed. This can happen if your doctor thinks you are likely to have severe side effects, if you get severe side effects, if your blood counts are affected and causing delays in treatment, or if you are finding it hard to cope with the treatment. This is called a dose reduction, dose change or treatment delay. Your doctor will explain if you need any changes or delays to your treatment and the reason why.

Blood tests and monitoring

You will need to have a blood test before you start treatment and regularly throughout your treatment. Your doctor or nurse will tell you when to have these blood tests.

Central venous access devices (CVADs)

This treatment may involve having chemotherapy through a central venous access device (CVAD). Your doctor or nurse will explain this to you. For more information, see the eviQ patient information sheets on CVADs.

Other medications given during this treatment

- Anti-sickness (anti-nausea) medication: you may be given some anti-sickness medication. Make sure you take this medication as your doctor or nurse tells you, even if you don't feel sick. This can help to prevent the sickness starting.
- **G-CSF**: you will be given injection(s) of a drug called G-CSF (also called filgrastim, lipegfilgrastim or pegfilgrastim) under your skin. This helps to boost your white blood cell count. Your white blood cells help to fight infection. Lipegfilgrastim and pegfilgrastim are given once. Filgrastim is given for several days until your white blood cells recover. Follow this link to read more information on how to give this injection.
- Eye drops: you will be given eye drops to help prevent sore eyes. You will start using the eye drops before you have your first dose of cytarabine and continue to use the eye drops until 72 hours after your last dose of cytarabine.
- **Prophylaxis medication:** you may need to take some medications to prevent infection and to help prevent or reduce some of the side effects of the chemotherapy. Your doctor or nurse will tell you how and when to take these medications.

Side effects

Cancer treatments can cause damage to normal cells in your body, which can cause side effects. Everyone gets different side effects, and some people will have more problems than others.

The table below shows some of the side effects you may get with this treatment. You are unlikely to get all of those listed and you may also get some side effects that have not been listed.

Tell your doctor or nurse about any side effects that worry you. Follow the instructions below and those given to you by your doctor or nurse.

Immediate (onset hours to days)				
Bone pain after G-CSF injection	 You may have discomfort or a dull ache in your pelvis, back, arms or legs. To reduce the pain, take paracetamol before each injection. Tell your doctor or nurse as soon as possible if your pain is not controlled. 			
Pain or swelling at injection site (extravasation)	 This treatment can cause serious injury if it leaks from the area where it is going into the vein. This can cause pain, stinging, swelling or redness at or near the site where the drug enters the vein. If not treated correctly, you may get blistering and ulceration. Tell your doctor or nurse immediately if you get any of the symptoms listed above during or after treatment. 			

You may get redness and itching along the vein where your chemotherapy is being infused. Redness and itching along • This will usually go away within 30 minutes of stopping the injection. vein Tell your doctor or nurse as soon as possible if you get any of the symptoms listed above. Your nurse will check to make sure the drug has not leaked out of the vein. • You may feel sick (nausea) or be sick (vomit). Nausea and vomiting • Take your anti-sickness medication as directed even if you don't feel sick. • Drink plenty of fluids (unless you are fluid restricted). · Eat small meals more frequently. • Try food that does not require much preparation. • Try bland foods like dry biscuits or toast. • Gentle exercise may help with nausea. · Ask your doctor or nurse for eviQ patient information - Nausea and vomiting during cancer Tell your doctor or nurse immediately, or go to the nearest hospital Emergency Department if you have uncontrolled vomiting or feel dizzy or light-headed. • High doses of cytarabine can affect the nervous system. Nervous system changes Tell your doctor or nurse immediately, or go to the nearest hospital Emergency from cytarabine Department if you get any of the following symptoms during or soon after your treatment: o dizziness, drowsiness or double vision agitation o difficulty walking in a straight line difficulty writing with a pen or pencil jerky movements o slow, slurred speech. · You may get: Eye problems from eye pain or irritation cytarabine blurred vision watery or gritty eyes sensitivity to light. • You will be given eye drops to help prevent and control these symptoms. It is important to use these eye drops as directed. Protect your eyes from the weather (sun and wind) by wearing sunglasses, especially if you have lost your eyelashes. • Tell your doctor or nurse if you get any of the symptoms listed above. • Your urine will turn an orange or red colour. Urine turning orange or red • This is not harmful and should only last for up to 48 hours after treatment. • You may find that food loses its taste or tastes different. Taste and smell changes These changes are likely to go away with time. · Do your mouth care regularly. • Chew on sugar-free gum or eat sugar-free mints. Add flavour to your food with sauces and herbs. Ask your doctor or nurse for eviQ patient information - Taste and smell changes during cancer treatment.

Early (onset days to weeks)

Infection risk (neutropenia)

- This treatment lowers the amount of white blood cells in your body. The type of white blood
 cells that help to fight infection are called neutrophils. Having low level of neutrophils is
 called neutropenia. If you have neutropenia, you are at greater risk of getting an infection. It
 also means that your body can't fight infections as well as usual. This is a serious side effect,
 and can be life threatening.
- · Wash your hands often.
- Keep a thermometer at home and take your temperature regularly, and if you feel unwell.
- Do your mouth care regularly.
- Inspect your central line site (if you have one) daily for any redness, pus or swelling.
- · Limit contact with people who are sick.
- Learn how to recognise the signs of infection.
- Ask your doctor or nurse for eviQ patient information Infection during cancer treatment.
- Tell your doctor or nurse immediately, or go to the nearest hospital Emergency Department if you get any of the following signs or symptoms:
 - a temperature of 38°C or higher
 - o chills, shivers, sweats or shakes
 - a sore throat or cough
 - uncontrolled diarrhoea
 - shortness of breath
 - o a fast heartbeat
 - become unwell even without a temperature.

Low platelets (thrombocytopenia)

- This treatment lowers the amount of platelets in your blood. Platelets help your blood to clot. When they are low, you are at an increased risk of bleeding and bruising.
- Try not to bruise or cut yourself.
- · Avoid contact sport or vigorous exercise.
- Clear your nose by blowing gently.
- · Avoid constipation.
- Brush your teeth with a soft toothbrush.
- Don't take aspirin, ibuprofen or other similar anti-inflammatory medications unless your doctor tells you to.
- Tell your doctor or nurse if you have any bruising or bleeding.
- Tell your doctor or nurse immediately, or go to your nearest hospital Emergency Department if you have any uncontrolled bleeding.

Flu-like symptoms from cytarabine

- You may get a fever, skin rash, aches and pains or increased sweating.
- These symptoms are caused by the drug cytarabine.
- Symptoms usually happen 6 to 12 hours after your dose, and may last until 24 hours after your treatment has finished.
- To reduce any pain or fever, take paracetamol, if needed.
- Don't take aspirin, ibuprofen or other similar anti-inflammatory medications unless your doctor tells you to.
- Tell your doctor or nurse if these symptoms do not get better after 24 hours.

Diarrhoea

- You may get bowel motions (stools, poo) that are more frequent or more liquid.
- You may also get bloating, cramping or pain.
- Take your antidiarrhoeal medication as directed by your doctor.
- Drink plenty of fluids (unless you are fluid restricted).
- Eat and drink small amounts more often.
- Avoid spicy foods, dairy products, high fibre foods, and coffee.
- Ask your doctor or nurse for eviQ patient information Diarrhoea during cancer treatment.
- Tell your doctor or nurse immediately, or go to your nearest hospital Emergency
 Department if your diarrhoea is not controlled, you have 4 or more loose bowel motions
 per day, and if you feel dizzy or light-headed.

Tiredness and lack of energy (fatigue)

- You may feel very tired, have no energy, sleep a lot, and not be able to do normal activities or things you enjoy.
- Do not drive or operate machinery if you are feeling tired.
- Nap for short periods (only 1 hour at a time)
- Prioritise your tasks to ensure the best use of your energy.
- Eat a well balanced diet and drink plenty of fluids (unless you are fluid restricted).
- Try some gentle exercise daily.
- Allow your friends and family to help.
- Tell your doctor or nurse if you get any of the symptoms listed above.

Mouth pain and soreness (mucositis)

- · You may have:
 - bleeding gums
 - o mouth ulcers
 - a white coating on your tongue
 - pain in the mouth or throat
 - o difficulty eating or swallowing.
- Avoid spicy, acidic or crunchy foods and very hot or cold food and drinks.
- Try bland and soft foods.
- Brush your teeth gently with a soft toothbrush after each meal and at bedtime. If you normally floss continue to do so.
- Rinse your mouth after you eat and brush your teeth, using either:
 - o 1/4 teaspoon of salt in 1 cup of warm water, or
 - 1/4 teaspoon of bicarbonate of soda in 1 cup of warm water
- Ask your doctor or nurse for eviQ patient information Mouth problems during cancer treatment.
- Tell your doctor or nurse if you get any of the symptoms listed above.

Skin that is more sensitive to the sun (photosensitivity)

- After being out in the sun you may develop a rash like a bad sunburn.
- Your skin may become red, swollen and blistered.
- · Avoid direct sunlight.
- Protect your skin from the sun by wearing sun-protective clothing, a wide-brimmed hat, sunglasses and a sunscreen of SPF 50 or higher.
- Tell your doctor or nurse if you get any of the symptoms listed above.

Skin rash

- You may get a red, bumpy rash and dry, itchy skin.
- Moisturise your skin with a gentle non-perfumed moisturising cream like sorbolene or aqueous cream.
- · Do not scratch your skin.
- Protect your skin from the sun by wearing sun-protective clothing, a wide-brimmed hat, sunglasses and sunscreen of SPF 50 or higher.
- Talk to your doctor or nurse about other ways to manage your skin rash.

Late (onset weeks to months) • You may feel dizzy, light-headed, tired and appear more pale than usual. Low red blood cells • Tell your doctor or nurse if you have any of these signs or symptoms. You might need a (anaemia) blood transfusion. Tell your doctor or nurse immediately, or go to the nearest hospital Emergency Department if you have any chest pain, trouble breathing, or feel like your heart is racing. • Your hair may start to fall out from your head and body. Hair loss (alopecia) • Hair loss usually starts 2 to 3 weeks after your first treatment. • You may become completely bald and your scalp might feel tender. • Use a gentle shampoo and a soft brush. • Take care with hair products like hairspray, hair dye, bleaches and perms. • Protect your scalp from the cold with a hat, scarf or wig. • Protect your scalp from the sun with a hat or sunscreen of SPF 50 or higher. • Moisturise your scalp to prevent itching. · Ask your doctor or nurse about the Look Good Feel Better program You may notice that you are unable to concentrate, feel unusually disorganised or tired Chemo brain (lethargic) and have trouble with your memory. (chemotherapy-related These symptoms usually improve once treatment is completed. cognitive impairment) Ask your doctor or nurse for eviQ patient information – Memory changes and chemotherapy (chemo brain). Tell your doctor or nurse if you get any of the symptoms listed above. Doses of fludarabine can affect the nervous system. **Nervous system changes** Tell your doctor or nurse immediately, or go to the nearest hospital Emergency from fludarabine Department if you get any of the following symptoms during or soon after your treatment: agitation or confusion o dizziness, drowsiness or double vision difficulty walking in a straight line difficulty writing with a pen or pencil jerky movements or seizures o slow, slurred speech. Lung problems are rare, but can be serious. They may occur throughout treatment or after Lung problems the completion of treatment. You may get: o shortness of breath fever dry cough wheezing fast heartbeat o chest pain. • Your doctor will monitor how well your lungs are working during your treatment. • Tell your doctor or nurse immediately, or go to the nearest hospital Emergency Department if you have chest pain or become short of breath.

Delayed (onset months to years)

Heart problems

- You may get:
 - o chest pain or tightness
 - o shortness of breath
 - swelling of your ankles
 - o an abnormal heartbeat.
- Heart problems can occur months to years after treatment.
- Tell your doctor if you have a history of heart problems or high blood pressure.
- Before or during treatment, you may be asked to have a test to see how well your heart is working.
- Tell your doctor or nurse immediately, or go to the nearest hospital Emergency Department if you get any of the symptoms listed above.

General advice for people having cancer treatment

Chemotherapy safety

- Learn how to keep you and your family safe while you are having anticancer drugs.
- See our patient information sheet Chemotherapy safety at home.

Blood clot risk

- Cancer and anticancer drugs can increase the risk of a blood clot (thrombosis).
- Tell your doctor if you have a family history of blood clots.
- A blood clot can cause pain, redness, swelling in your arms or legs, shortness of breath or chest pain.
- If you have any of these symptoms go to your nearest hospital Emergency Department.

Medications and vaccinations

- Before you start treatment, tell your doctor about any medications you are taking, including vitamins or herbal supplements.
- Don't stop or start any medications during treatment without talking to your doctor and pharmacist first.
- Paracetamol is safe to take if you have a headache or other mild aches and pains. It is recommended that you avoid taking aspirin, ibuprofen and other anti-inflammatory type medications for pain while you are having treatment. However, if these medications have been prescribed by your doctor, do not stop taking them without speaking with your doctor.
- Vaccinations such as flu and tetanus vaccines are safe to receive while having treatment. Do not have any live vaccines during your treatment or for 6 months after it finishes. If you are unsure, check with your doctor before you have any vaccinations.
- People you live with should be fully vaccinated, including having live vaccines according to the current vaccination schedule. Extra
 care needs to be taken with hand washing and careful disposal of soiled nappies for infants who have recently received the
 rotavirus vaccine.

Other medical and dental treatment

- If you go to hospital or any other medical appointment (including dental appointments), always tell the person treating you that you are receiving anticancer drugs.
- Before you have any dental treatment, talk to your doctor.

Diet and food safety

- While you are receiving this treatment it is important that you try to maintain a healthy diet.
- Speak to your doctor or nurse about whether drinking alcohol is safe with your treatment.
- If you have any concerns about recent weight loss or weight gain or questions about your diet, ask to speak to a dietitian.
- There are some foods that may cause infection in high risk individuals and should be avoided. For more information on foods to avoid and food hygiene please ask for a copy of the Listeria and food brochure.

Fertility

- Some cancer treatments can reduce your fertility. This can make it difficult or impossible to get pregnant or father a child.
- Talk to your doctor or nurse before you start any treatment. Depending on your situation there may be fertility sparing options available to you and/or your partner, discuss these with your doctor or nurse.

Pregnancy and breastfeeding

- Some cancer treatments can be dangerous to unborn babies. Talk to your doctor or nurse if you think there is any chance that you could be pregnant.
- Do not try to get pregnant or father a child during this treatment. Contraception should be used during treatment and after stopping treatment. Ask your doctor or nurse about what type of contraception you should use.
- If you are planning pregnancy/fatherhood after completing this treatment, talk to your doctor. Some doctors advise waiting between 6 months and 2 years after treatment.
- · Do not breastfeed if you are on this treatment, as anti-cancer medications can also pass into breast milk.

Sex life and sexuality

- The desire to have sex may decrease as a result of this treatment or its side effects.
- Your emotions and the way you feel about yourself may also be affected by this treatment.
- It may help to discuss your concerns with your partner and doctor or nurse.

Quitting smoking

- It is never too late to quit smoking. Quitting smoking is one of the best things you can do to help your treatment work better.
- · There are many effective tools to improve your chances of quitting.
- Talk to your treating team for more information and referral to a smoking cessation support service.

Staying active

- Research shows that exercise, no matter how small, has many benefits for people during and after cancer treatment.
- Talk to your doctor before starting an exercise program. Your doctor can advise whether you need a modified exercise program.

For more information about cancer treatment, side effects and side effect management see our Patient and carers section.

Where to get more information

Telephone support

- Call Cancer Council on 13 11 20 for cancer information and support
- Call the Leukaemia Foundation on 1800 620 420 (Mon to Fri 9am 5pm)
- Call the Lymphoma Nurse Support Line on 1800 953 081 (Mon to Fri 9am 5pm)
- Call the Myeloma Australia Support Line on 1800 693 566 (Mon to Fri 9am 5pm)

Haematology, transplant and cellular therapy information

- Arrow bone marrow transplant foundation arrow.org.au
- Australasian Menopause Society menopause.org.au
- Chris O'Brien Lifehouse Total Body Irradiation mylifehouse.org.au/departments/radiation-oncology/total-body-irradiation/
- Healthy Male Andrology Australia healthymale.org.au/
- International Myeloma Foundation myeloma.org
- Leukaemia Foundation leukaemia.org.au
- Lymphoma Australia lymphoma.org.au
- Myeloma Australia myeloma.org.au
- NSW Agency for Clinical Innovation, Blood & Marrow Transplant Network https://aci.health.nsw.gov.au/networks/bmtct
- NSW Agency for Clinical Innovation aci.health.nsw.gov.au/projects/immune-effector-cell-service
- NCCN Guidelines for Patients Immunotherapy Side Effects: CAR T-Cell Therapy nccn.org/patientresources/patient-resources/guidelines-for-patients
- Talk Blood Cancer cmlsupport.org.uk/organisation-type/social-media-groups

General cancer information and support

- Australian Rare Cancer (ARC) Portal arcportal.org.au/
- Beyondblue beyondblue.org.au
- Cancer Australia canceraustralia.gov.au
- Cancer Council Australia cancer.org.au
- Cancer Voices Australia cancervoicesaustralia.org
- CanTeen canteen.org.au
- Carers Australia carersaustralia.com.au

- Carer Help carerhelp.com.au
- eviQ Cancer Treatments Online eviQ.org.au
- Food Standards Australia New Zealand: Listeria & Food Safety foodstandards.gov.au/publications/pages/listeriabrochuretext.aspx
- LGBTQI+ People and Cancer cancercouncil.com.au/cancer-information/lgbtqi
- Look Good Feel Better lgfb.org.au
- · Patient Information patients.cancer.nsw.gov.au
- Radiation Oncology Targeting Cancer targetingcancer.com.au
- Redkite redkite.org.au
- Return Unwanted Medicines returnmed.com.au
- Staying active during cancer treatment patients.cancer.nsw.gov.au/coping-with-cancer/physical-wellbeing/staying-active

Quit smoking information and support

Quitting smoking is helpful even after you have been diagnosed with cancer. The following resources provide useful information and support to help you quit smoking. Talk to your treating team about any other questions you may have.

- Call Quitline on 13 QUIT (13 78 48)
- iCanQuit iCanQuit.com.au
- · Patient Information patients.cancer.nsw.gov.au/coping-with-cancer/physical-wellbeing/quitting-smoking
- Quitnow quitnow.gov.au

Additional notes:		

This document is a guide only and cannot cover every possible situation. The health professionals caring for you should always consider your individual situation when making decisions about your care. Contact your cancer clinic staff or doctor if you have any questions or concerns about your treatment, or you are having problems coping with side effects. While eviQ endeavours to link to reliable sources that provide accurate information, eviQ and the Cancer Institute NSW do not endorse or accept responsibility for the accuracy, currency, reliability or correctness of the content of linked external information sources. Use of this document is subject to eviQ's disclaimer available at www.eviQ.org.au

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