



Fludarabine/Busulfan/ATG Grafalon® - RIC - MUD

INDICATIONS FOR USE:

INDICATION	ICD10	Regimen Code	Reimbursement Status
Reduced intensity conditioning for matched unrelated donor allogeneic stem cell transplant in patients with myeloid disorders.	C92	00635a	Hospital

TREATMENT:

Conditioning chemotherapy is administered over 9 days. Stem cells are infused on day 0.

Facilities to treat anaphylaxis MUST be present when conditioning therapy and stem cells are administered.

D	B	D t.	Dilyand & Data
Drug	Dose	Route	Diluent & Rate
Fludarabine ^a	30mg/m ²	IV infusion	100ml sodium chloride 0.9% over 30 minutes
Busulfan ^{b,c}	0.8mg/kg	IV infusion	(See note ^d) ml sodium chloride 0.9% over 2 hours
Busulfan ^{b,c}	0.8mg/kg	IV infusion	(See note ^d) ml sodium chloride 0.9% over 2 hours
Busulfan ^{b,c}	0.8mg/kg	IV infusion	(See note ^d) ml sodium chloride 0.9% over 2 hours
Busulfan ^{b,c}	0.8mg/kg	IV infusion	(See note ^d) ml sodium chloride 0.9% over 2 hours
er 15 hours, infusion	must begin at t	time specified	
e,f,g ATG Grafalon®	20mg/kg	IV infusion	(See noteh) ml sodium chloride 0.9% over 12 hours
e,f,g ATG Grafalon®	20mg/kg	IV infusion	(See noteh) ml of sodium chloride 0.9% over 10 hours
Stem cell infusion			
Methotrexate ⁱ	10mg/m ²	IV infusion	50mls of sodium chloride 0.9% over 10 minutes
	Busulfan ^{b,c} Busulfan ^{b,c} Busulfan ^{b,c} Busulfan ^{b,c} Busulfan ^{b,c} ter 15 hours, infusion e,f,g ATG Grafalon* e,f,g ATG Grafalon* Stem cell infusion	Fludarabine ^a 30mg/m ² Busulfan ^{b,c} 0.8mg/kg er 15 hours, infusion must begin at a ce,f,g ATG Grafalon 20mg/kg Stem cell infusion	Fludarabine ^a 30mg/m ² IV infusion Busulfan ^{b,c} 0.8mg/kg IV infusion er 15 hours, infusion must begin at time specified e.f.g ATG Grafalon [®] 20mg/kg IV infusion e.f.g ATG Grafalon [®] 20mg/kg IV infusion Stem cell infusion

Dose rounding:

Fludarabine doses ≤50mg to the nearest 2.5mg and doses >50mg to the nearest 5mg

Busulfan to the nearest 1.2mg if <60mg, to nearest 6mg if >60mg. Oral busulfan available as 2mg and 25mg tablets.

ATG Grafalon® to the nearest 20mg

Methotrexate to the nearest 2.5mg

^aAll patients who have received fludarabine should receive irradiated blood products (lifetime recommendation).

bIV busulfan may be replaced with oral busulfan at the discretion of the haematology consultant An oral dose of 1mg/kg is equivalent to the 0.8mg/kg IV dose The dosing schedule for oral busulfan is 06:00, 12:00, 18:00, 23:59

'If a problem with an infusion bag (i.e. leaking bag, short expiry) is discovered outside of 8.30am-5pm, an oral dose of busulfan 1mg/kg equivalent to the intravenous dose will be available from the MDA press on Denis Burkitt Ward. This can only be used after discussion with a haematology consultant and must be prescribed by haematology registrar or consultant on a chemotherapy prescription/NCIS

d Calculation of busulfan infusion solution: [(busulfan dose (mg) divided by 6) x 10] [to the nearest 10ml] NaCl 0.9% - concentration to be as close to 0.5mg/ml as possible.

ePatient monitoring is required during the ATG Grafalon® infusion: BP, pulse, respiration and temperature at 15, 30 and then 60 minute intervals for the duration of the infusion.

'If an infusion reaction occurs during the administration of ATG Grafalon®, the infusion should be slowed. Chills and fever generally respond to antihistamines, antipyretics or corticosteroids. If the patient becomes hypotensive or experiences chest or back pain, indicating anaphylaxis, the infusion should be stopped and the medical team contacted immediately.

^g Platelets should be >50x10⁹/L pre day 1 ATG Grafalon® treatment. If the patient has no reaction to ATG Grafalon®, platelets can be maintained at >30x10⁹/L for the remaining days of ATG Grafalon® administration. Platelets should be maintained at >50x10⁹/L in the setting of clinically symptomatic bleeding.

hEach ml of ATG Grafalon® should be diluted with 6ml of sodium chloride 0.9% in accordance with SPC. Pharmacy to complete volume.

Day +1 methotrexate should be administered at least 24 hours after the stem cells have infused. In the event where this timing results in methotrexate being infused during the night, it is reasonable to reschedule the administration time of the day +3 methotrexate to the next morning, to avoid administration during the night. The amended administration timing can then be maintained for subsequent methotrexate doses.

*denotes recommended administration time

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ELIGIBILITY:

- Indications as above
- Medical assessment as per SJH BMT assessment form

EXCLUSIONS:

- Hypersensitivity to fludarabine, busulfan, ATG Grafalon®, methotrexate or any of the excipients.
- Pregnancy and lactation

PRESCRIPTIVE AUTHORITY:

• The treatment plan must be initiated by a Haematology Consultant working in the area of stem cell transplantation in a unit suitable for carrying out this treatment.

TESTS:

 Baseline and regular tests in accordance with SJH Haematopoietic Stem Cell Transplant work-up protocols

Disease monitoring:

Disease monitoring should be in line with the patient's treatment plan and any other test/s as directed by the supervising Consultant.

DOSE MODIFICATIONS:

- Any dose modification should be discussed with a Haematology Consultant.
- Chemotherapy dosing in obese adult patients: For patients with a BMI > 30kg/m² please refer to 'Chemotherapy Dosing in Obese Adult Stem Cell Transplant Recipients – Guidelines' for guidance on individual drug dosing as per SJH policy available on the SJH intranet.
- Renal and Hepatic Impairment:
 - Dose modifications are generally not undertaken in conditioning regimens.
 - Discuss with the consultant if hepatic impairment or if creatinine clearance is <70ml/min for advice on fludarabine dosing. Guidance to inform this discussion available at:
 U:\PHARMCOMP\Clinical\haematology\Haematology Drugs\Fludarabine
 - Consult the following resources to inform any renal or hepatic dose modification discussions:
 - o Summary of product characteristics (SPC) available at http://www.hpra.ie
 - Krens et al Lancet Oncol 2019;20(4) e200-e207 "Dose Recommendations for anticancer drugs in patients with renal or hepatic impairment" available at https://pubmed.ncbi.nlm.nih.gov/30942181/
 - UCHL renal impairment guidelines and hepatic impairment guidelines available on SJH intranet

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SUPPORTIVE CARE:

Antiemetics

Table 1: Recommended SJH Regimen Specific Antiemetics

Prevention of a	revention of acute emesis		Prevention of delayed emesis		Comments	
Drug	Dose	Admin day	Drug	Dose	Admin day	No additional
Ondansetron	8mg PO/IV TDS	-5, -4, -3	No delayed cover required	N/A	N/A	dexamethasone is required due to steroid cover with ATG Grafalon® supportive medication

ATG Grafalon supportive medications:

- Methylprednisolone 2mg/kg once daily IV 90mins before commencing ATG on Day -3 to Day -1
- Chlorphenamine 10mg IV 30mins before commencing ATG on Day -3 to Day -1
- Prednisolone 1mg/kg once daily PO (or and equivalent IV alternative starting on Day 0 and continuing for 5 days
- Taper to zero over next 5 days to prevent serum sickness

Busulfan conditioning seizure prophylaxis:

 Phenytoin 600mg STAT orally at midnight the night before busulfan treatment, then 300mg once daily PO on day -5 to day -3

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OTHER SUPPORTIVE CARE:

Table 2: Recommended SJH regimen specific supportive care

GvHD prophylaxis:	Ciclosporin	Tacrolimus
Refer to signed off BMT assessment form for confirmed choice and target level of immunosuppression	 Ciclosporin 5mg/kg once daily IV over 6 hours from day -1 The equivalent oral dose is: (Total IV dose x 0.67) twice daily PO Target levels: 100-150microgram/Litre 	 0.03mg/kg once daily IV over 22 hours, starting from day -1 The equivalent oral dose is: (Total IV dose) twice daily PO Target levels: 5-10 nanogram/ml
GvHD and VOD prophylaxis	Ursodeoxycholic acid 250rContinue until day +90	
HSV prophylaxis	Valaciclovir 500mg once d Or	oral route not available or ANC <
CMV prophylaxis Prescribe for all CMV seropositive recipients	+1 if patient is receiving cide Letermovir 480mg once date +1 if patient is receiving tate Letermovir via the oral rout Letermovir IV at the same	aily PO/IV, as appropriate, starting Day closporin immunosuppression aily PO/IV, as appropriate, starting Day crolimus immunosuppression ate is first line. oral dose should be prescribed only tolerate oral or where there are on.
	Patients should bring their oral lete admission. High tech prescription we their counselling appointment prepharmacist if any supply issues arise. When ANC>1.0 x 10 ⁹ /L, pre-emptive tube] (Tuesday and Fridays) should reactivation/infection in <u>all</u> patients.	vill have been provided to patient at admission. Liaise with transplant e. e monitoring (9mls in EDTA [purple be carried out for CMV

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Antifungal prophylaxis	When ANC<0.5 x 10 ⁹ /L or if patient on high dose steroids • Liposomal amphotericin 1mg/kg once daily IV Mon/Wed/I	Fri
Refer to signed off BMT assessment form for confirmed choice of antifungal prophylaxis	OrCaspofungin 70mg/kg once daily IV Mon/Wed/Fri	
	If at higher risk due to prior possible/probable fungal infection: • Liposomal amphotericin 1mg/kg once daily IV Or	
	 Caspofungin 70mg once daily IV if >80kg Or Caspofungin 70mg once daily IV on day 1 of treatment and 	d
	50mg once daily IV thereafter if <80kg	
PJP prophylaxis	First line therapy	
	 Co-trimoxazole 960mg BD Mon/Wed/Fri PO Commence only on engraftment when ANC > 1.0x10⁹/L if appropriate 	
	Second line therapy (if allergic to co-trimoxazole or contraindicated PJP Prophylaxis and T. gondii IgG NEGATIVE	<u>d):</u>
	 Pentamidine 300mg nebule and salbutamol 2.5mg nebule pentamidine, every 4 weeks plus 	pre-
	Phenoxymethylpenicillin 333mg BD daily PO	
	Continue the phenoxymethylpenicillin until patients have been revaccinated and have adequate pneumococcal/haemophilus titres	S
	PJP prophylaxis and T.gondii IgG POSITIVE	
	 Atovaquone 750mg BD PO plus 	
	 Pyrimethamine 25mg once daily PO plus 	
	Folinic acid 15mg once daily PO plus	
	Phenoxymethylpenicillin 333mg BD daily PO	
	Continue the phenoxymethylpenicillin until patients have been	
	revaccinated and have adequate pneumococcal/haemophilus titre	S
	Please note: If a patient is to be discharged on atovaquone, pyrimethamine or folinic acid, please contact pharmacy in advance	to
Mouthcare:	arrange supply and funding through a community drugs scheme Mucositis WHO grade < 2: • Sodium chloride 0.9% 10ml QDS mouthwash	
	Nystatin 1ml QDS PO (use 15 minutes after sodium chloric 0.9% mouthwash)	de
	 Mucositis WHO grade ≥2: Chlorhexidine digluconate 0.12% (Kin® mouthwash) 10mls mouthwash Nystatin 1ml QDS PO (use 15 minutes after Kin® mouthwash) 	
Gastro protection:	Lansoprazole 30mg /omeprazole 40mg once daily PO Or	-
	Esomeprazole 40mg once daily IV (if oral route not available)	ole)
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Folate supplementation:	Methotrexate is included as GvHD prophylaxis. Folinic acid should not		
	be administered on the same days as methotrexate		
	The first dose of folinic acid must be administered at a minimum of 24		
	hours post completion of methotrexate. Prescribe as outlined below:		
	 Folinic acid 15mg once daily IV on days +2,+4,+5, and +7 onwards 		
	Switch to folic acid 5mg once daily PO when oral route is		
	available		
Vitamin K supplementation	Beginning on day + 2 post stem cell transplant		
	 Vitamin K (phytomenadione) 10mg once weekly IV 		
Prevention of vaginal bleeding;	If required for menstruating female patients until platelets > 50 x10 ⁹ /L		
	 Norethisterone 5mg TDS PO if >55Kg 		
	 Norethisterone 5mg BD PO if <55kg 		
Tumour Lysis syndrome	Consider allopurinol in active disease pre transplant		
	 Allopurinol 300mg once daily PO for 5-7 days and review 		
Hepatitis B prophylaxis/treatment	A virology screen is completed as part of transplant workup. Hepatitis B		
	prophylaxis or treatment may be initiated in consultation with a Virology		
	Consultant or Hepatology Consultant if required.		
	Options may include:		
	 Lamivudine 100mg once daily PO 		
	Or		
	Entecavir 500mcg once daily PO		
Prevention of constipation	Consider laxatives if appropriate e.g.		
	 Senna two tablets (15mg) nocte PO while on ondansetron. 		
Antibiotic standing order	Antibiotic standing order should be prescribed for neutropenic		
	sepsis/neutropenic fever based on previous microbiology and renal		
	function		
	Piptazobactam 4.5g QDS IV		
	Plus		
	 Amikacin* 15mg/kg once daily IV 		
	*Ciprofloxacin 400mg BD IV may be considered instead of amikacin in		
	cases of renal impairment		
	Refer to local Antimicrobial Guidelines in the Prescriber's Capsule for		
	antibiotic choice where a patient is allergic to any of the above		
Magnesium and Potassium Standing	Magnesium and Potassium Standing order: Magnesium and potassium		
order:	standing orders should be prescribed for all transplant patients in		
	accordance with stem cell unit practice as indicated on EPMAR.		
VTE prophylaxis	Consider VTE prophylaxis in accordance with local SJH policy		
Bone Health	Consider calcium and vitamin D supplementation prior to discharge for		
	patients who are on high dose steroids. Other medications for		
	maintenance of bone health may need to be considered as appropriate.		
	Calcium carbonate and colecalciferol (Caltrate® 600mg/400unit)		
	one tablet BD		

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Hepatic veno occlusive disease (VOD):

- Defibrotide may be prescribed for the treatment of hepatic veno-occlusive disease (VOD) in consultation with the haematology consultant
- Dosing of intravenous Defibrotide :
 - o The recommended dose is 6.25mg/kg IV every 6 hours (25mg/kg/day)
 - Calculate the total daily dose. Divide by 200 to calculate the total number of vials needed and split
 the dose such that the minimum amount of wastage can be achieved. Defibrotide should be
 administered for a minimum of 21 days and continued until the signs and symptoms of VOD resolve.
 - IV infusion is given over 2 hours (maximum concentration 400mg/100ml NaCl 0.9%)

ADVERSE EFFECTS / REGIMEN SPECIFIC COMPLICATIONS:

 Please refer to the relevant Summary of Product Characteristics and SJH Stem Cell Transplant Programme PPGs for full details.

DRUG INTERACTIONS:

• The relevant Summary of Product Characteristics and current drug interaction databases should be consulted.

REFERENCES:

- 1. Kröger N et al. Allogeneic stem cell transplantation after reduced-intensity conditioning in patients with myelofibrosis: a prospective, multicentre study of the Chronic Leukemia Working Party of the European Group for Blood and Marrow Transplantation. Blood. 2009;114:5264-5270
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- 7. NCCP Classification Document for Systemic Anti-Cancer Therapy (SACT) Induced Nausea and Vomiting. V3 2021. Available at: https://www.hse.ie/eng/services/list/5/cancer/profinfo/chemoprotocols/nccp-classification-document-for-systemic-anti-cancer-therapy-sact-induced-nausea-and-vomiting.pdf
- 8. Fludara® summary of product characteristics accessed Oct 2020 available at https://www.hpra.ie/img/uploaded/swedocuments/Licence PA0611-004-001 11112019115658.pdf

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- 10. Grafalon ATG * summary of product characteristics accessed Oct 2020 Available at : https://www.hpra.ie/img/uploaded/swedocuments/Licence PA1015-001-001 19032020152832.pdf
- 11. Methotrexate 1g/10ml Summary of Product Characteristics. Accessed October 2020. Available at https://www.hpra.ie/img/uploaded/swedocuments/Licence PA0822-206-006 19052021104201.pdf

Version	Date	Amendment	Approved By
1	06/08/2021		SJH Stem Cell Transplant Group
1a	09/07/2024	Extension of review date as agreed with	NCCP
		clinical reviewer	1100.

Comments and feedback welcome at oncologydrugs@cancercontrol.ie.

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