Indication	Dostarlimab in combination with platinum-containing chemotherapy for the 1 st line treatment of mismatch repair deficient (dMMR) / microsatellite instability-high (MSI-H) primary advanced or recurrent endometrial cancer (EC) in patients who are not candidates for potentially curative surgery or radiotherapy or chemoradiotherapy but are eligible for systemic therapy. Patients should not have previously received any systemic chemotherapy for endometrial carcinoma unless it has been as neoadjuvant or adjuvant chemotherapy or chemoradiotherapy and the patient has progressed or recurred at least 6 months since the completion of treatment.			
Treatment	Palliative			
Intent				
Frequency	6 cycles of dostarlimab in combination with chemotherapy followed by dostarlimab monotherapy.			
and number				
of cycles	Cycles 1-6 combination therapy repeat every 21 days			
	Cycle 7 onwards repeat every 42 days until disease progression, unacceptable toxicity, patient			
	choice or to a maximum duration of 3 calendar years.			
	A formal medical review should take place after the first 6 weeks of treatment.			
Monitoring	Virology screening: All new patients referred for systemic anti-cancer treatment should be			
Parameters	screened for hepatitis B and C and the result reviewed prior to the start of treatment. Patients			
pre-treatment	not previously tested who are starting a new line of treatment, should also be screened for			
	hepatitis B and C. Further virology screening will be performed following individual risk			
	assessment and clinician discretion.			
	Thyroid function must be assessed at baseline then at least every 6 weeks throughout			
	treatment.			
	Cortisol monitoring should be undertaken in line with ESMO immunotherapy toxicity guidance			
	available on KMCC website (see link below). Cortisol level should not be taken within 24hours of			
	the last steroid dose.			
	Monitor for signs and symptoms of myocarditis. Carry out ECG as clinically indicated.			
	• Cycles 1-6			
	• EDTA/DTPA should be used to measure GFR prior to cycle 1. C+G may be used to estimate CrCl if there is a delay in obtaining EDTA result.			
	 Monitor U+Es, LFTs, FBC and glucose at each cycle. If CrCl falls by >25% repeat EDTA and d/w consultant. 			
	• If neuts <1.5 and/or PLT <100 defer treatment by one week. Consider dose reduction on			
	subsequent cycles.			
	Cycle 7 onwards			
	FBC, U&E, LFTs and glucose at each cycle.			
	Hepatic impairment:			
	o Dostarlimab - No recommended dose adjustment in mild hepatic impairment. Limited data			
	in moderate impairment and no available data in severe impairment.			
	Carboplatin - No dose adjustment required.			
	Paclitaxel - If bilirubin < 1.25 x ULN and transaminase < 10 x ULN, dose at full dose. Otherwise consider dose reduction, not recommended in source benefit impairment.			
	Otherwise consider dose reduction, not recommended in severe hepatic impairment.			
	 Renal impairment: Dostarlimab - No recommended dose adjustment in mild or moderate renal impairment. 			
	 Dostarlimab - No recommended dose adjustment in mild or moderate renal impairment. Limited data in severe impairment or end-stage renal disease undergoing dialysis. 			
	Carboplatin - stop if CrCl<30ml/min			
	Paclitaxel - no dose reduction necessary.			
l	1 <i>I</i>			

Protocol No	GYN-049 POST-EAMS	Kent and Medway SACT Protocol Disclaimer: No responsibility will be accepted for the accuracy of this information when used elsewhere.		
Version	V2	Written by	M.Archer	
Supersedes	V1	Checked by	C.Waters	
version			O. Adebayo	
Date	14.11.2023	Authorising consultant (usually NOG Chair)	L.Kivat	

 The use of systemic corticosteroids or immunosuppressants before starting dostarlimab should be avoided; dexamethasone is permitted as prescribed within this protocol. Systemic corticosteroids or other immunosuppressants can be used after starting dostarlimab to treat immune-related adverse reactions.

• Infusion-related reactions:

Dostarlimab

For severe infusion reactions (grade 3-4), infusion should be stopped and dostarlimab permanently discontinued.

Grade 2 reaction, withhold dose. If reaction resolves within 1 hour of stopping, the infusion may be restarted at 50 % of the original infusion rate, or restart when symptoms resolve with pre-medication. If grade 2 recurs with adequate premedication, permanently discontinue.

Paclitaxel

Patients developing hypersensitivity reactions to paclitaxel may be rechallenged with full dose paclitaxel following prophylactic medication (e.g. famotidine 40mg po given 4 hours prior to treatment plus hydrocortisone 100mg iv and chlorphenamine 10mg iv 30 minutes prior to treatment, then give paclitaxel over 3-6 hours (i.e. starting at over 6 hours and gradually increase rate if possible).

Carboplatin

- Mild/moderate reactions (grade 1-2): If symptoms resolve after treatment with hydrocortisone and chlorphenamine, the infusion may be restarted at 50% rate for 30 mins, then, if no further reaction, increase to 100% rate.
- If symptoms do not resolve after treatment with hydrocortisone and chlorphenamine, do
 not restart the infusion. At consultant's discretion, patients may be rechallenged at a later
 date with additional prophylaxis. In the event of further reaction (grade 1-3), stop infusion
 and consider alternative treatment.
- Severe (grade 3): Do not restart infusion. Consider alternative treatment.
- Anaphylaxis (grade 4): Follow anaphylaxis protocol. Discontinue permanently and consider alternative treatment.

• Adverse reactions and dose modification:

- Immune-related adverse reactions may appear during or after treatment with dostarlimab. The most common immune-related reactions are: anaemia (including autoimmune haemolytic anaemia), pneumonitis, colitis, hyperthyroidism, hypothyroidism and arthralgia. The following additional, immune related adverse reactions have been reported in patients receiving dostarlimab: type 1 diabetes, nephritis, hepatitis, pancreatitis, severe skin reactions, encephalitis, Guillain-Barre syndrome, myocarditis, iridocyclitis, uveitis and diabetic ketoacidosis. See guidelines for management of immune-related adverse reactions following immunotherapy: https://www.kmcc.nhs.uk/medicines-and-prescribing-incorporating-sact-pathways/immunotherapy/
- Cases of Stevens-Johnson syndrome (SJS) and toxic epidermal necrolysis (TEN), have been
 reported. If SJS or TEN is confirmed, dostarlimab should be permanently discontinued. Caution
 should be used when considering the use of dostarlimab in a patient who has previously
 experienced a severe or life-threatening skin adverse reaction on prior treatment with other
 immune-stimulatory anticancer agents.

• Dose Modification:

O Dose reduction of either chemotherapy agents (paclitaxel or carboplatin) and not the other agent is appropriate if the toxicity is clearly related to one of the chemotherapy agents. If the toxicity is related to both agents, they should both be modified according to their recommended dose modification. If the toxicity is related to the combination of dostarlimab and both chemotherapy agents, the doses of the chemotherapy agents should be reduced, or the dose of all three drugs should be interrupted or discontinued. Patients may have

Protocol No	GYN-049	Kent and Medway SACT Protocol		
	POST-EAMS	Disclaimer: No responsibility will be accepted for the accuracy of this information when used elsewhere.		
Version	V2	Written by	M.Archer	
Supersedes	V1	Checked by	C.Waters	
version			O. Adebayo	
Date	14.11.2023	Authorising consultant (usually NOG Chair)	L.Kivat	

	chemotherapy discontinued and continue dostarlimab as monotherapy, in the same way patients may discontinue dostarlimab and continue chemotherapy alone. Dostarlimab - Dose reduction is not recommended. Based on the severity of the adverse reaction, treatment with dostarlimab should be withheld or permanently discontinued and corticosteroids (1 to 2 mg/kg/day prednisone or equivalent) or other appropriate therapy administered (see table 1 and EAMs protocol). Upon improvement to Grade =1, corticosteroid taper should be initiated and continued for 1 month or longer. Treatment with dostarlimab should be permanently discontinued for any Grade 3 immune-related adverse reaction that recurs and for any Grade 4 immune-related adverse reaction toxicity, except for endocrinopathies that are controlled with replacement hormones and unless otherwise specified in table 1 / EAMs protocol. Paclitaxel - Dose reduce Paclitaxel by 20% in the event of /= grade 2 neuropathy and consider delay until recovery to = grade 1. Consider omitting paclitaxel in event of recurrent grade /= 3 neuropathy OR recurrent or persistent >/= grade 2 neuropathy following a dose reduction. Dose reduction of carboplatin and paclitaxel should be considered if any other grade 3 or 4 non-haematological toxicity or repeat appearance of grade 2 (except N&V and alopecia). Delay until resolution of toxicity to = grade 1. Common drug interactions (for comprehensive list refer to BNF/SPC): Dostarlimab - No interaction studies have been performed. Paclitaxel - Caution should be exercised when administering paclitaxel concomitantly with medicines known to inhibit either CYP2C8 or CYP3A4 (e.g. ketoconazole, erythromycin, fluoxetine, clopidogrel, cimetidine, ritonavir and nelfinavir); toxicity may be increased. CYP2C8 or CYP3A4 inducers (e.g. rifampicin, carbamazepine, phenytoin, efavirenz, nevirapine) may reduce efficacy. Carboplatin - Caution with other nephrotoxic drugs. Each patient should be given a copy of the Jemperli * patient alert card at</th
References	SPC accessed online 25.09.2023 KMCC protocol GYN-049 EAMS Blueteq form accessed online
	20.10.2023

 $\ensuremath{\mathsf{NB}}$ For funding information, refer to CDF and NICE Drugs Funding List

Protocol No	GYN-049 POST-EAMS	Kent and Medway SACT Protocol Disclaimer: No responsibility will be accepted for the accuracy of this information when used elsewhere.		
Version	V2	Written by	M.Archer	
Supersedes	V1	Checked by	C.Waters	
version			O. Adebayo	
Date	14.11.2023	Authorising consultant (usually NOG Chair)	L.Kivat	

Table 1: Dose modification for dostarlimab

Immune-related adverse reactions	Severity grade	Dose modification
C-list-	2 to 3	Withhold dose. Restart dosing when toxicity resolves to grade 0-1.
Colitis	4	Permanently discontinue.
Honotitis	Grade 2 with AST or ALT > 3 and up to 5 × ULN Or total bilirubin > 1.5 and up to 3 × ULN	Withhold dose. Restart dosing when toxicity resolves to grade 0 to 1.
Hepatitis	Grade >/=3 with AST or ALT > 5 × ULN Or total bilirubin > 3 × ULN	Permanently discontinue (see exception below*)
Type 1 diabetes mellitus (T1DM)	3 to 4 (hyperglycaemia)	Withhold dose. Restart dosing in appropriately managed, clinically and metabolically stable patients.
Hypophysitis or adrenal insufficiency	2, 3 or 4	Withhold dose. Restart dosing when toxicity resolves to grade 0 to 1. Permanently discontinue for recurrence or worsening while on adequate hormonal therapy.
Hypothyroidism or hyperthyroidism	3 to 4	Withhold dose. Restart dosing when toxicity resolves to grade 0 to 1.
Pneumonitis	2	Withhold dose. Restart dosing when toxicity resolves to grade 0-1. If grade 2 recurs, permanently discontinue.
	3 to 4	Permanently discontinue.
A1 1 11	2	Withhold dose. Restart dosing when toxicity resolves to grade 0-1.
Nephritis	3 to 4	Permanently discontinue.
Exfoliative dermatologic conditions (e.g., SJS, TEN,	Suspected	Withhold dose for any grade. Restart dosing if not confirmed and when toxicity resolves to grade 0-1.
DRESS)	Confirmed	Permanently discontinue.
Myocarditis	2,3, or 4	Permanently discontinue.
Severe neurological toxicities (myasthenic syndrome/myasthenia gravis, Guillain-Barré syndrome, encephalitis, transverse myelitis)	2,3 or 4	Permanently discontinue
Other immune-related adverse	3	Withhold dose. Restart dosing when toxicity resolves to grade 0-1.
reactions (including but not limited to myositis, sarcoidosis, autoimmune haemolytic anaemia, pancreatitis, iridocyclitis, uveitis, diabetic ketoacidosis, arthralgia, solid organ transplant rejection, graft-versus-host disease	4	Permanently discontinue.
Recurrence of immune-related adverse reactions after resolution to ≤ grade 1 (except for pneumonitis, see above)	3 to 4	Permanently discontinue.

^{*}For patients with liver metastases who begin treatment with grade 2 increase of AST or ALT, if AST or ALT increases by >/= 50 % relative to baseline and lasts for at least 1 week, then treatment should be discontinued.

Protocol No	GYN-049 POST-EAMS	Kent and Medway SACT Protocol Disclaimer: No responsibility will be accepted for the accuracy of this information when used elsewhere.		
Version	V2	Written by	M.Archer	
Supersedes	V1	Checked by	C.Waters	
version			O. Adebayo	
Date	14.11.2023	Authorising consultant (usually NOG Chair)	L.Kivat	

Cycle 1 to 6: Repeat every 21 days

Day	Drug	Dose	Route	Infusion Duration	Administration
1	DOSTARLIMAB	500mg	IV	30mins	In 100ml Sodium Chloride 0.9% Administered via a 0.22 micron in-line filter
		Give pre-med	ls 30 minu	tes prior to	paclitaxel
	Dexamethasone	16mg	IV	Bolus	
	Chlorphenamine	10mg	IV	Slow bolus	Through the side of a fast running Sodium Chloride 0.9% intravenous infusion.
	Ondansetron	<75yrs 16mg <u>></u> 75yrs 8mg	IV	15 min	Sodium chloride 0.9% 50ml
	PACLITAXEL	175mg/m²	IV	3 hrs	In 500ml Sodium Chloride 0.9% (if dose <150mg in 250ml Sodium Chloride 0.9%) Use non-PVC bag and non-PVC administration set via in-line 0.22 microns filter. Flush with sodium chloride 0.9%
	CARBOPLATIN Dose = (GFR + 25) x AUC	AUC 5 Maximum dose 700mg	IV	30 mins	Glucose 5% 500ml
TTO	Drug	Dose	Route	Directions	
Day 1	Dexamethasone	6mg	РО	OM for 3 days starting the day after paclitaxel dos	
	Metoclopramide	10mg	РО	(Maximum of 30mg per day). Do not take for more than 5 days continuously. Take 4mg (2 capsules) initially, then 2mg (1 capsule) after each loose stool when required.	
	Loperamide	2-4mg	РО		

Protocol No	GYN-049	Kent and Medway SACT Protocol		
	POST-EAMS	Disclaimer: No responsibility will be accepted for the accuracy of this information when used elsewhere.		
Version	V2	Written by	M.Archer	
Supersedes	V1	Checked by	C.Waters	
version			O. Adebayo	
Date	14.11.2023	Authorising consultant (usually NOG Chair)	L.Kivat	

Cycle 7 onwards repeat every 42 days (cycle 7 to be started 21 days after cycle 6)

Day	Drug	Dose	Route	Infusion	Administration
				Duration	
1	DOSTARLIMAB	1000mg	IV	30mins	In 100ml Sodium Chloride 0.9% Administered via a 0.22 micron in-line filter
TTO	Drug	Dose	Route	Directions	
Day 1	Metoclopramide	10mg	РО	10mg TDS PRN. Do not take for more than 5 days continuously.	

Protocol No	GYN-049	Kent and Medway SACT Protocol		
	POST-EAMS	Disclaimer: No responsibility will be accepted for the accuracy of this information when used elsewhere.		
Version	V2	Written by	M.Archer	
Supersedes	V1	Checked by	C.Waters	
version			O. Adebayo	
Date	14.11.2023	Authorising consultant (usually NOG Chair)	L.Kivat	