SA2498 - Pembrolizumab

APPLICATION FOR SUBSIDY BY SPECIAL AUTHORITY

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APPLICANT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:	First Names:	First Names:
Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Pembrolizumab		
Initial application — stage III or IV resectable melanoma - neoadjuvant Applications only from a relevant specialist or any relevant practitioner on the recommendation of a relevant specialist. Approvals valid Prerequisites(tick boxes where appropriate) The individual is currently on treatment with pembrolizumab and met all remaining criteria prior to commencing treatment or The individual has resectable stage IIIB, IIIC, IIID or IV melanoma (excluding uveal) (see note) The individual has not received prior funded systemic treatment in the perioperative setting for their stage IIIB, IIII and Treatment must be prior to complete surgical resection and Pembrolizumab must be administered as monotherapy and Pembrolizumab to be administered at a fixed dose of 200 mg every 3 weeks (or equivalent)		ia prior to commencing treatment al) (see note) ve setting for their stage IIIB, IIIC, IIID or IV

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Name	:			Surname:	Surname:
Addre	ss:			DOB:	Address:
				Address:	
Fax N	umbe	er:			Fax Number:
Pem	broli	izumab	- continued		
Curre	ent ap	proval Nui	I or IV resectable melanon mber (if known): m a relevant specialist or an oxes where appropriate)	•	evant specialist. Approvals valid for 4 months.
	The individual has received neoadjuvant treatment with an immune checkpoint inhibitor and The individual meets initial application criteria for pembrolizumab for stage III or IV resected melanoma – adjuvant or				
		and		d neoadjuvant and adjuvant treatment with an immune val criteria for pembrolizumab for stage III or IV resect	·
	or The individual has received neoadjuvant and adjuvant treatment with an immune checkpoint inhibitor and			checkpoint inhibitor	
		and	The individual has metasta	tic or unresectable melanoma (excluding uveal) stage	III or IV
	2.		The individual meets initial	application criteria for pembrolizumab for unresectable	e or metastatic melanoma
	or	and and	The individual has received	d neoadjuvant and adjuvant treatment with an immune d treatment with an immune checkpoint inhibitor for un val criteria for pembrolizumab for unresectable or meta	resectable or metastatic melanoma
Note	:				
a) S	tage I	IIB, IIIC, II	ID or IV melanoma defined	as per American Joint Committee on Cancer (AJCC) 8	th Edition
				ete surgical resection means either 13 weeks after res	ection (primary or lymphadenectomy) or 13 weeks

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Name:	Surname:	Surname:			
Address:	DOB:	Address:			
	Address:				
Fax Number:		Fax Number:			
Pembrolizumab - continued					
Initial application — stage III or IV resected me Applications only from a relevant specialist or any Prerequisites(tick boxes where appropriate)	lanoma - adjuvant relevant practitioner on the recommendation of a rele	evant specialist. Approvals valid for 4 months.			
The individual is currently on treati	ment with pembrolizumab and met all remaining crite	ria prior to commencing treatment			
The individual has resected and	stage IIIB, IIIC, IIID or IV melanoma (excluding uveal) (see note a)			
Adjuvant treatment with per	nbrolizumab is required				
	ved prior funded systemic treatment in the adjuvant so	etting for stage IIIB, IIIC, IIID or IV melanoma			
	on to complete surgical resection				
	within 13 weeks of complete surgical resection, unles	ss delay is necessary due to post-surgery			
Pembrolizumab must be adr	ministered as monotherapy				
The individual has ECOG pe	erformance score 0-2				
	istered at a fixed dose of 200 mg every 3 weeks (or e	equivalent)			
Note:	Note:				
a) Stage IIIB, IIIC, IIID or IV melanoma defined as	s per American Joint Committee on Cancer (AJCC) 8	th Edition			
b) Initiating treatment within 13 weeks of complete surgical resection means 13 weeks after resection (primary or lymphadenectomy)					

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Name:				Surname:	Surname:
Address	:			DOB:	Address:
				Address:	
Fax Nun	nber:				Fax Number:
Pembr	oliz	umab	- continued		
Renew	al —	stage III	l or IV resected melanoma -	adjuvant	
Applica	uisit	only fror	No evidence of disease recu Pembrolizumab must be adr Pembrolizumab to be adminitreatment course, including a	relevant practitioner on the recommendation of a rele	or equivalent) for a maximum of 12 months total 2 months total treatment course (equivalent to
O	The individual has received adjuvant treatment with an immune checkpoint inhibitor and The individual has metastatic or unresectable melanoma (excluding uveal) stage III or IV and				
	The individual meets initial application criteria for pembrolizumab for unresectable or metastatic melanoma			e or metastatic melanoma	
o	r				
		and	The individual has received	adjuvant treatment with an immune checkpoint inhibit	or
		and	The individual has received	treatment with an immune checkpoint inhibitor for unr	esectable or metastatic melanoma
			The individual meets renewa	al criteria for pembrolizumab for unresectable or meta	static melanoma

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Name:	Surname:	Surname:	
Address:	DOB:	Address:	
	Address:		
Fax Number:		Fax Number:	
Pembrolizumab - continued			
The individual has metastatic or unand Baseline measurement of overall to and The individual has ECOG performs and The individual has not received to and and The individual has not received to and The individual has not received to and The individual has received to an another the individual has received to an		/ ally	
or	The individual did not receive treatment in the perioperative setting with a PD-1/PD-L1 inhibitor		
The individual received	ed treatment in the perioperative setting with a PD-1/PD-L1 inhibitor		
	experience disease recurrence while on treatment with that PD-1/PD-L1 inhibitor		
	The individual did not experience disease recurrence within six months of completing perioperative treatment with a		

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Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Pembrolizumab - continued		
Renewal — unresectable or metastatic mel	anoma, less than 24 months on treatment	
Applications only from a relevant specialist or Prerequisites(tick boxes where appropriate) The individual's dior	any relevant practitioner on the recommendation of a release has had a complete response to treatment isease has had a partial response to treatment	evant specialist. Approvals valid for 4 months.
treatment period	in target lesions has been determined by comparable rac	diologic assessment following the most recent
The individual has previ	ously discontinued treatment with pembrolizumab for rea	asons other than severe toxicity or disease
The individual has signs of disease progression		
	ssed during previous treatment with pembrolizumab	
Current approval Number (if known):	any relevant practitioner on the recommendation of a relevant practitioner on the recommendation of a relevant practitioner on the recommendation of a relevant isease has had a complete response to treatment isease has had a partial response to treatment is stable disease in target lesions has been determined by comparable ractions of disease progression	diologic assessment following the most recent

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Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:		Fax Number:
Pembrolizumab - continued		
The individual has been on treatment and The individual has been on treatment and The individual's or The individual's or The individual has and Response to treatment the most recent treatment and The individual has predisease progression and The individual has sign and The individual has sign and	relevant practitioner on the recommendation of a relevant for more than 24 months disease has had a complete response to treatment disease has had a partial response to treatment as stable disease	le radiologic or clinical assessment following or reasons other than severe toxicity or

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APPLICANT (stamp or sticker acceptable)	PATIENT NHI:	REFERRER Reg No:
Reg No:	First Names:	First Names:
Name:	Surname:	Surname:
Address:	DOB:	Address:
	Address:	
Fax Number:Pembrolizumab - continued		Fax Number:
Initial application — non-small cell lung cancer Applications only from a medical oncologist or any Prerequisites(tick boxes where appropriate)	first-line monotherapy relevant practitioner on the recommendation of a monotherapy	edical oncologist. Approvals valid for 4 months.
and Patient has not had chemotherapy and Patient has not received prior funde and For patients with non-squamous his EGFR or ALK tyrosine kinase unles and Pembrolizumab to be used as mon and	otherapy irming the disease expresses PD-L1 at a level great	sease does not express activating mutations of
a validated test unless	n confirming the disease expresses PD-L1 at a level not possible to ascertain	
and Patient has an ECOG 0-2 and Pembrolizumab to be used at a ma	ximum dose of 200 mg every three weeks (or equiva	alent) for a maximum of 16 weeks
Baseline measurement of overall to	umour burden is documented clinically and radiologic	cally

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Addre	ess:	DOB:	Address:		
		Address:			
Fax N	lumber:		Fax Number:		
Peml	brolizumab - continued				
Rene	ewal — non-small cell lung cancer fi	rst line monotherapy			
Appli	ent approval Number (if known):ications only from a medical oncologist equisites (tick boxes where appropriate	or any relevant practitioner on the recor	nmendation of a medical oncologist. Approvals valid for 4 months.		
	or Patient's disease has	had a complete response to treatment			
	Patient has stable dis	eease			
period			mparable radiologic assessment following the most recent treatment		
	and No evidence of disease progression and				
The treatment remains clinically appropriate and patient is benefitting from treatment			ng from treatment		
	Pembrolizumab to be used at a maximum dose of 200 mg every three weeks (or equivalent) and				
	Treatment with pembrolizur 3 weeks)	nab to cease after a total duration of 24	months from commencement (or equivalent of 35 cycles dosed every		
Appl	Initial application — non-small cell lung cancer first-line combination therapy Applications only from a medical oncologist or any relevant practitioner on the recommendation of a medical oncologist. Approvals valid for 4 months. Prerequisites(tick boxes where appropriate)				
	Patient has locally advance	d or metastatic, unresectable, non-smal	cell lung cancer		
	The patient has not had che	emotherapy for their disease in the pallia	tive setting		
		or funded treatment with an immune che	eckpoint inhibitor for NSCLC		
	For patients with non-squar	nous histology there is documentation c se unless not possible to ascertain	onfirming that the disease does not express activating mutations of		
		in combination with platinum-based che	motherapy		
	Patient has an ECOG 0-2				
		at a maximum dose of 200 mg every thr	ee weeks (or equivalent) for a maximum of 16 weeks		
		verall tumour burden is documented clir	ically and radiologically		

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Name:	Surname:	Surname:	
Address:	DOB:	Address:	
	Address:		
Fax Number:		Fax Number:	
Pembrolizumab - continued			
Renewal — non-small cell lung cancer first line	combination therapy		
Current approval Number (if known):			
Applications only from a medical oncologist or any relevant practitioner on the recommendation of a medical oncologist. Approvals valid for 4 months. Prerequisites(tick boxes where appropriate)			
Patient's disease has had a	complete response to treatment		
or			
Patient's disease has had a	partial response to treatment		
Patient has stable disease			
Response to treatment in target lesions has been determined by comparable radiologic assessment following the most reception period		assessment following the most recent treatment	
No evidence of disease progressio	n		
	propriate and patient is benefitting from treatment		
	uximum dose of 200 mg every three weeks (or equiva	alent)	
Treatment with pembrolizumab to c 3 weeks)	cease after a total duration of 24 months from comme	encement (or equivalent of 35 cycles dosed every	

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	Address:		
Fax Number:		Fax Number:	
Pembrolizumab - continued			
Prerequisites(tick boxes where appropriate)	relevant practitioner on the recommendation of a rele		
or express ER, PR or HE	or de novo unresectable, inoperable locally advanced R2 IHC3+ or ISH+ [including FISH or other technology or de novo metastatic triple-negative breast cancer (the H or other technology])	gy])	
and			
Patient is treated with palliati			
Patient's cancer has confirmed and	Patient's cancer has confirmed PD-L1 Combined Positive Score (CPS) is greater than or equal to 10 and		
Patient has received no prior	Patient has received no prior systemic therapy in the palliative setting and		
Patient has an ECOG score of 0–2			
	Pembrolizumab is to be used in combination with chemotherapy		
Baseline measurement of ov	Baseline measurement of overall tumour burden is documented clinically and radiologically		
Pembrolizumab is to be used	d at a maximum dose of 200 mg every three weeks (or equivalent) for a maximum of 16 weeks	

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Address:		DOB:	Address:				
		Address:					
Fax Number:			Fax Number:				
Pembrol	izumab - continued						
Renewal -	— breast cancer, advanced						
Current ap	oproval Number (if known):						
	ns from any relevant practitioner. Approsites(tick boxes where appropriate)	vals valid for 6 months.					
Frerequis	sites(tick boxes where appropriate)						
	Patient's disease has had a complete response to treatment						
	or Patient's disease has had a	partial response to treatment					
	or Patient has stable disease						
and							
and	No evidence of disease progressi	on					
		esions has been determined by a comparable radiolog	ic assessment following the most recent				
and	·	maniferance does of 000 mm and the same lie (or and	in a land				
and		maximum dose of 200 mg every three weeks (or equi	·				
	Treatment with pembrolizumab is every 3 weeks)	to cease after a total duration of 24 months from com	mencement (or equivalent of 35 cycles dosed				
	Dication — head and neck squamous ons only from a relevant specialist or any	cell carcinoma relevant practitioner on the recommendation of a rele	evant specialist. Approvals valid for 4 months.				
Prerequis	sites(tick boxes where appropriate)						
or	Patient is currently on treatment v	vith pembrolizumab and met all remaining criteria prior	r to commencing treatment				
		tastatic head and neck squamous cell carcinoma of m	nucosal origin (excluding nasopharyngeal				
	carcinoma) that is incurable	by local therapies					
	Patient has not received pri	or systemic therapy in the recurrent or metastatic setti	ng				
	Patient has a positive PD-L	1 combined positive score (CPS) of greater than or eq	jual to 1				
	Patient has an ECOG perfo	rmance score of 0-2					
	and Pembrolizumab to be	used in combination with platinum-based chemothera	apy dan				
	or	•					
	and	used as monotherapy					
		ed at a maximum dose of 200 mg every three weeks (o	or equivalent) for a maximum of 16 weeks				

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Address:	DOB:	Address:				
	Address:					
Fax Number:		Fax Number:				
Pembrolizumab - continued						
Renewal — head and neck squamous cell card	inoma					
Current approval Number (if known):						
Applications from any relevant practitioner. Approx Prerequisites (tick boxes where appropriate)	vals valid for 4 months.					
relequisites (lick boxes where appropriate)						
Patient's disease has had a complete response to treatment						
Patient's disease has had a	partial response to treatment					
or Patient has stable disease						
and						
No evidence of disease progression	on					
Pembrolizumab is to be used at a	maximum dose of 200 mg every three weeks (or equi	ivalent)				
	to cease after a total duration of 24 months from com	mencement (or equivalent of 35 cycles dosed				
Initial application — MSI-H/dMMR advanced colorectal cancer Applications only from a relevant specialist or any relevant practitioner on the recommendation of a relevant specialist. Approvals valid for 4 months. Prerequisites(tick boxes where appropriate)						
Individual is currently on treatment	with pembrolizumab and met all remaining criteria p	rior to commencing treatment				
Individual has deficier	nt mismatch repair (dMMR) or microsatellite instability	-high (MSI-H) metastatic colorectal cancer				
or Individual has deficier	nt mismatch repair (dMMR) or microsatellite instability	-high (MSI-H) unresectable colorectal cancer				
and						
Individual is treated with pal						
Individual has not previously and	received funded treatment with pembrolizumab for N	MSI-H/dMMR advanced colorectal cancer				
Individual has an ECOG per	formance score of 0-2					
	verall tumour burden is documented clinically and rad	iologically				
	at a maximum dose of 200 mg every three weeks (or	equivalent) for a maximum of 16 weeks				

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Address:	DOB:	Address:					
	Address:						
Fax Number:		Fax Number:					
Pembrolizumab - continued							
Renewal — MSI-H/dMMR advanced colorectal o	eancer						
Current approval Number (if known):							
Applications from any relevant practitioner. Approv Prerequisites (tick boxes where appropriate)	als valid for 4 months.						
No evidence of disease progression	n						
	aximum dose of 200 mg every three weeks (or equiva	alent)					
Treatment with pembrolizumab is to every 3 weeks)	o cease after a total duration of 24 months from com	mencement (or equivalent of 35 cycles dosed					
Initial application — Urothelial carcinoma Applications only from a relevant specialist or any relevant practitioner on the recommendation of a relevant specialist. Approvals valid for 4 months. Prerequisites(tick boxes where appropriate)							
Patient is currently on treatment with pembrolizumab and met all remaining criteria prior to commencing treatment or							
Patient has inoperable locall	y advanced (T4) or metastatic urothelial carcinoma						
Patient has an ECOG perfor	mance score of 0-2						
Patient has documented dise	ease progression following treatment with chemother	ару					
Pembrolizumab to be used a 16 weeks	s monotherapy at a maximum dose of 200 mg every	three weeks (or equivalent) for a maximum of					
Renewal — Urothelial carcinoma							
nenewai — Oromenai caremonia							
Current approval Number (if known): Applications only from a relevant specialist or any in Prerequisites (tick boxes where appropriate)	relevant practitioner on the recommendation of a rele	vant specialist. Approvals valid for 4 months.					
or Patient's disease has had a	complete response to treatment						
Patient's disease has had a or	partial response to treatment						
Patient has stable disease							
and No evidence of disease progressio	n						
and Pembrolizumab to be used at a ma	aximum dose of 200 mg every three weeks (or equiva	alent)					
and Treatment with pembrolizumab is to every 3 weeks)	o cease after a total duration of 24 months from com	mencement (or equivalent of 35 cycles dosed					

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Address:	DOB:	Address:				
	Address:					
Fax Number:		Fax Number:				
Pembrolizumab - continued						
Prerequisites(tick boxes where appropriate) Individual is currently on treatment or Individual has reand Individual has relapse and Individual has not previously and	relevant practitioner on the recommendation of a relevant practitioner on the recommendation of a relevant practition of a relevant period of the provided provided and met all remaining criteria provided provid	ore lines of chemotherapy Indergone an autologous stem cell transplant Dory Hodgkin lymphoma				
Renewal — relapsed/refractory Hodgkin lymphoma						
Current approval Number (if known):						
Applications only from a relevant specialist or any relevant practitioner on the recommendation of a relevant specialist. Approvals valid for 6 months. Prerequisites(tick boxes where appropriate)						
Patient has received a partial or complete response to pembrolizumab						
	o cease after a total duration of 24 months from com	mencement (or equivalent of 35 cycles dosed				