## **Formulary Adherence Checklist for NICE Technology Appraisals About Medicines**

Milton Keynes Health Economy



Developed for the NHS by: East and South East England Specialist Pharmacy Services

This spreadsheet is updated monthly and enables self-audit of a medicines formulary for adherence to current NICE Technology Appraisals. Version 20 November 2017 All guidelines refer to adults unless indicated. No copyright is asserted on this material if used for non-commercial purposes within the NHS.

Technology appraisal (TA)  Titles are hyperlinks to full guidance	Date of TA Release	Availability of medicine for NHS patients with this medical condition, as indicated by NICE	Adherence of local formulary to NICE					
			<b>Yes</b> (mark 'x' if applicable)	<b>N/A</b> (mark 'x' if applicable)	Date of local decision (DD/MM/YY)	Time to implement (days)	<b>Notes</b> (e.g. rationale, method of making available)	
2017/2018								
Venetoclax for treating chronic lymphocytic leukaemia (TA487)	01/10/2017	Venetoclax (Venclyxto®) is recommended for use within the Cancer Drugs Fund, within its marketing authorisation, as an option for treating chronic lymphocytic leukaemia	х		29/11/2017	59	NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements and the National Cancer Drugs Fund requirements	
Aflibercept for treating choroidal neovascularisation (TA486)	01/10/2017	Aflibercept (Eylea®) is recommended, within its marketing authorisation, as an option for treating visual impairment because of myopic choroidal neovascularisation in adults	х		29/11/2017	59	CCG commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements	
Sarilumab for moderate to severe rheumatoid arthritis (TA485)	01/10/2017	Sarilumab (Kevzara®) with methotrexate, is recommended as an option for treating active rheumatoid arthritis in adults whose disease has responded inadequately to intensive therapy with a combination of conventional diseasemodifying anti-rheumatic drugs (DMARDs)	х		29/11/2017	59	CCG commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements	
Nivolumab for previously treated non-squamous non-small-cell lung cancer (TA484)	01/10/2017	Nivolumab (Opdivo®) is recommended for use within the Cancer Drugs Fund as an option for treating locally advanced or metastatic non-squamous non-small-cell lung cancer in adults after chemotherapy	х		29/11/2017	59	NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements and the National Cancer Drugs Fund requirements	
Nivolumab for previously treated squamous non-small- cell lung cancer (TA483)	01/10/2017	Nivolumab (Opdivo®) is recommended for use within the Cancer Drugs Fund as an option for treating locally advanced or metastatic squamous non-small-cell lung cancer in adults after chemotherapy	х		29/11/2017	59	NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements and the National Cancer Drugs Fund requirements	
Immunosuppressive therapy for kidney transplant in children and young people (TA482)	01/10/2017	Immunosuppressive therapy for kidney transplant in children and young people	х		29/11/2017	59	NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements	
Immunosuppressive therapy for kidney transplant in adults (TA481)	01/10/2017	Immunosuppressive therapy for kidney transplant in adults	х		29/11/2017	59	NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements	

Tofacitinib for moderate to severe rheumatoid arthritis (TA480)	01/10/2017	Tofacitinib (Xeljanz®) is recommended as an option for treating active rheumatoid arthritis in adults whose disease has responded inadequately to intensive therapy with a combination of conventional disease-modifying anti-rheumatic drugs (DMARDs)	x	29/11/2017	59 CCG commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Reslizumab for treating severe eosinophilic asthma (TA479)	01/10/2017	Reslizumab (Cinqaero®) is recommended as an option for the treatment of severe eosinophilic asthma that is inadequately controlled in adults despite maintenance therapy with high-dose inhaled corticosteroids plus another drug	x	29/11/2017	59 NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Brentuximab vedotin for treating relapsed or refractory systemic anaplastic large cell lymphoma (TA478)	01/10/2017	Brentuximab vedotin (Adcetris®) is recommended as an option for treating relapsed or refractory systemic anaplastic large cell lymphoma in adults	х	29/11/2017	59 NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements and the National Cancer Drugs Fund requirements
Autologous chondrocyte implantation for treating symptomatic articular cartilage defects of the knee (TA477)	01/10/2017	Autologous chondrocyte implantation (ACI) ChondroCelect® is recommended as an option for treating symptomatic articular cartilage defects of the knee	х	29/11/2017	59 NICE Appraisal N/A at MK
Paclitaxel as albumin-bound nanoparticles with gemcitabine for untreated metastatic pancreatic cancer (TA476)	06/09/2017	Paclitaxel as albumin-bound nanoparticles (nab-paclitaxel) with gemcitabine is recommended as an option for untreated metastatic adenocarcinoma of the pancreas in	х	27/09/2017	Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Dimethyl fumarate for treating moderate to severe plaque psoriasis (TA475)	06/09/2017	<b>Dimethyl fumarate</b> is recommended as an option for treating plaque psoriasis in adults	х	29/11/2017	84 Confirmed place in therapy with dermatologists. CCG commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements.
Sorafenib for treating advanced hepatocellular carcinoma (TA474)	06/09/2017	Sorafenib is recommended as an option for treating advanced hepatocellular carcinoma only for people with Child-Pugh grade A liver impairment	х	27/09/2017	Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Cetuximab for treating recurrent or metastatic squamous cell cancer of the head and neck (TA473)	31/08/2017	Cetuximab in combination with platinum-based chemotherapy is recommended as an option for treating recurrent or metastatic squamous cell cancer of the head and neck in adults	x	27/09/2017	27 CCG commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements.
Obinutuzumab with bendamustine for treating follicular lymphoma refractory to rituximab (TA472)	30/08/2017	Obinutuzumab in combination with bendamustine followed by obinutuzumab maintenance is recommended for use within the Cancer Drugs Fund as an option for treating adults with follicular lymphoma that did not respond or progressed during or up to 6 months after treatment with rituximab or a rituximab-containing regimen	x	27/09/2017	28 Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Eluxadoline for treating irritable bowel syndrome with diarrhoea (TA471)	30/08/2017		x	27/09/2017	28 Approved for addition to the formulary and to be used in line with NICE guidance with Secondary Care Initiation and stabilisation including one month review.

Ofatumumab with chemotherapy	23/08/2017	Ofatumumab with chemotherapy - was unable		27/09/2017	35	Terminated Appraisal by NICE
for treating chronic lymphocytic		to make a recommendation about the use in the				
leukaemia (terminated appraisal)		NHS of ofatumumab with chemotherapy for				
(TA470)		treating chronic lymphocytic leukaemia because	х			
<u>(1711-0)</u>		no evidence submission was received from				
		Novartis Pharmaceuticals UK				
		Novartis Pharmaceuticais UK				
Idelalisib with ofatumumab for	23/08/2017	NICE is unable to make a recommendation about		27/09/2017	35	Terminated Appraisal by NICE
treating chronic lymphocytic		the use in the NHS of idelalisib with ofatumumab	x			
leukaemia (terminated appraisal)		for treating chronic lymphocytic leukaemia	X			
(TA469)		because no evidence submission was received				
Methylnaltrexone bromide for	23/08/2017	Methylnaltrexone bromide - was unable to make		27/09/2017	35	Terminated Appraisal by NICE
treating opioid-induced	-,,	a recommendation about the use in the NHS of		, ,		
constipation (terminated		methylnaltrexone bromide for treating opioid-				
appraisal) (TA468)		induced constipation because no evidence	х			
<u>appraisal) (1A400)</u>						
		submission was received from Swedish Orphan				
	101001	Biovitrum Ltd		0=1001		
Holoclar for treating limbal stem	16/08/2017	Holoclar (ex vivo expanded autologous human		27/09/2017	42	Approved for addition to the formulary and to be used in line with NICE guidance and
cell deficiency after eye burns		corneal epithelial cells containing stem cells) is				commissioning statements
<u>(TA467)</u>		recommended as an option in people with	x			
		moderate to severe limbal stem cell deficiency				
		after eve burns				
Baricitinib for moderate to severe	09/08/2017			27/09/2017	49	Approved for addition to the formulary and to be used in line with NICE guidance and
rheumatoid arthritis (TA466)		Baricitinib, with methotrexate is recommended				commissioning statements
		as an option for treating active rheumatoid				
		arthritis in adults whose disease has responded	x			
		inadequately to intensive therapy with a				
		combination of conventional disease-modifying				
		antirheumatic drugs (DMARDs),				
Olaratumab in combination with	09/08/2017	Olaratumab, in combination with doxorubicin,		27/09/2017	49	Approved for addition to the formulary and to be used in line with NICE guidance and
doxorubicin for treating advanced	03,00,201,	is recommended for use within the Cancer Drugs		27/03/2017	.5	commissioning statements
		Fund as an option for advanced soft tissue	X			commissioning statements
soft tissue sarcoma (TA465)		i ·				
Bisphosphonates for treating	09/08/2017	sarcoma in adults Oral bisphosphonates (alendronic acid,		27/09/2017	40	Approved for addition to the formulary and to be used in line with NICE guidance and
	09/08/2017	ibandronic acid and risedronate sodium) are		27/09/2017	49	,
osteoporosis (TA464)		,	x			commissioning statements
		recommended as options for treating				
	20/20/20:=	osteoporosis in adults		27/22/224		A 16 100 - 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Cabozantinib for previously	09/08/2017	Cabozantinib is recommended, within its		27/09/2017	49	Approved for addition to the formulary and to be used in line with NICE guidance and
treated advanced renal cell		marketing authorisation, as an option for				commissioning statements
carcinoma (TA463)		treating advanced renal cell carcinoma in adults	x			
		after vascular endothelial growth factor (VEGF)-				
		targeted therapy				
Nivolumab for treating relapsed	26/07/2017	Nivolumab is recommended, within its		27/09/2017	63	Approved for addition to the formulary and to be used in line with NICE guidance and
or refractory classical Hodgkin	20/07/2017	· ·		27/03/2017	03	
		marketing authorisation, as an option for				commissioning statements
<u>lymphoma (TA462)</u>		treating relapsed or refractory classical Hodgkin				
		lymphoma in adults after autologous stem cell				
		transplant and treatment with brentuximab	x			
		vedotin, when the company provides nivolumab				
		with the discount agreed in the patient access				
		scheme.				

Roflumilast for treating chronic	26/07/2017	-		27/09/2017	63 Approved for addition to the formulary and to be used in line with NICE guidance and
obstructive pulmonary disease	26/07/2017	Roflumilast, as an add-on to bronchodilator		27/09/2017	
		therapy, is recommended as an option for	x		commissioning statements
<u>(TA461)</u>		treating severe chronic obstructive pulmonary	^		
		disease in adults with chronic bronchitis			
Adalimumab and dexamethasone	26/07/2017			27/09/2017	63 Approved for addition to the formulary and to be used in line with NICE guidance and
for treating non-infectious uveitis	26/07/2017	•		27/09/2017	
		treating non-infectious uveitis in the posterior			commissioning statements
<u>(TA460)</u>		segment of the eye in adults with inadequate			
		response to corticosteroids.			
			x		
		Dexamethasone (Ozurdex®) intravitreal implant			
		is recommended as an option for treating non-			
		infectious uveitis in the posterior segment of the			
		eye in adults			
Collagenase clostridium	26/07/2017	Collagenase clostridium histolyticum for people		27/09/2017	63 Approved for addition to the formulary and to be used in line with NICE guidance and
	20/07/2017			27/09/2017	, , ,
histolyticum for treating		who meet the inclusion criteria for the ongoing			commissioning statements
<u>Dupuytren's contracture (TA459)</u>		clinical trial (HTA-15/102/04), comparing			
		collagenase clostridium histolyticum (CCH) with			
		limited fasciectomy, are encouraged to	х		
		participate in the study.			
		For people not taking part in the ongoing clinical			
		trial, CCH is recommended as an option for			
		treating Dupuytren's contracture with a palpable			
		cord in adults			
Trastuzumab emtansine for	19/07/2017	Trastuzumab emtansine is recommended,		27/09/2017	70 Approved for addition to the formulary and to be used in line with NICE guidance and
treating HER2-positive advanced		within its marketing authorisation, as an option			commissioning statements
breast cancer after trastuzumab		for treating human epidermal growth factor			
and a taxane (TA458)		receptor 2 (HER2)-positive, unresectable, locally			
<u> </u>		advanced or metastatic breast cancer in adults	x		
		who previously received trastuzumab and a			
		taxane, separately or in combination.			
		taxane, separately of in combination.			
Carfilzomib for previously treated	19/07/2017	Carfilzomib in combination with		27/09/2017	70 Approved for addition to the formulary and to be used in line with NICE guidance and
multiple myeloma (TA457)		dexamethasone is recommended as an option	x		commissioning statements
		for treating multiple myeloma in adults	^		
<u>Ustekinumab for moderately to</u>	12/07/2017	Ustekinumab is recommended, within its		27/09/2017	77 Approved for addition to the formulary and to be used in line with NICE guidance and
severely active Crohn's disease		marketing authorisation, as an option for			commissioning statements
after previous treatment (TA456)		treating moderately to severely active Crohn's			
		disease, that is, for adults who have had an			
		inadequate response with, lost response to, or	x		
		were intolerant to either conventional therapy			
		or a TNF-alpha inhibitor or have medical			
		contraindications to such therapies.			
		contractions to sacriffications.			
Adalimumab, etanercept and	12/07/2017	Adalimumab, Etanercept and Ustekinumab is		27/09/2017	77 Approved for addition to the formulary and to be used in line with NICE guidance and
ustekinumab for treating plaque		recommended as an option for treating plaque	X		commissioning statements
psoriasis in children and young		psoriasis in children and young people aged 4	X		
people (TA455)		vears or older			
<del></del>					

Daratumumab with lenalidomide and dexamethasone for treating relapsed or refractory multiple myeloma (terminated appraisal)  (TA454)	05/07/2017	Daratumumab - was unable to make a recommendation about the use in the NHS of daratumumab, with lenalidomide and dexamethasone, for treating relapsed or refractory multiple myeloma because no evidence submission was received from Janssen-Cilag	х	26/07/2017	21	Terminated Appraisal by NICE
Bortezomib for treating multiple myeloma after second or subsequent relapse (terminated appraisal) (TA453)	05/07/2017	Bortezomib - was unable to make a recommendation about the use in the NHS of bortezomib for treating multiple myeloma after second or subsequent relapse because no evidence submission was received from Janssen-Cilag	х	26/07/2017	21	Terminated Appraisal by NICE
Ibrutinib for untreated chronic lymphocytic leukaemia without a 17p deletion or TP53 mutation (terminated appraisal) (TA452)	05/07/2017	Ibrutinib - was unable to make a recommendation about the use in the NHS of ibrutinib for untreated chronic lymphocytic leukaemia without a 17p deletion or TP53 mutation because no evidence submission was received from Janssen–Cilag	х	26/07/2017	21	Terminated Appraisal by NICE
Ponatinib for treating chronic myeloid leukaemia and acute lymphoblastic leukaemia (TA451)	28/06/2017	Ponatinib (Iclusig) is recommended, within its marketing authorisation, as an option for treating chronic-, accelerated- or blast-phase chronic myeloid leukaemia in adults	х	26/07/2017		NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Blinatumomab for previously treated Philadelphia-chromosome- negative acute lymphoblastic leukaemia (TA450)	28/06/2017	Blinatumomab (Blincyto) is recommended within its marketing authorisation as an option for treating Philadelphia-chromosome-negative relapsed or refractory precursor B-cell acute lymphoblastic leukaemia in adults, only if the company provides it with the discount agreed in the patient access scheme.	x	26/07/2017		NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Everolimus and sunitinib for treating unresectable or metastatic neuroendocrine tumours in people with progressive disease (TA449)	28/06/2017	Everolimus (Afinitor) and Sunitinib (Sutent) are recommended, within their marketing authorisations, as options for treating well- or moderately differentiated unresectable or metastatic neuroendocrine tumours (NETs) of pancreatic origin in adults with progressive disease.	х	26/07/2017		NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Etelcalcetide for treating secondary hyperparathyroidism (TA448)	28/06/2017	Etelcalcetide (Parsabiv) is recommended as an option for treating secondary hyperparathyroidism in adults with chronic kidney disease on haemodialysis	х	26/07/2017		NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Pembrolizumab for untreated PD- L1-positive metastatic non-small- cell lung cancer (TA447)	28/06/2017	Pembrolizumab (Keytruda) is recommended for use within the Cancer Drugs Fund as an option for untreated PD-L1-positive metastatic nonsmall-cell lung cancer in adults	х	26/07/2017	II.	NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements
Brentuximab vedotin for treating CD30-positive Hodgkin lymphoma (TA446)	28/06/2017	Brentuximab vedotin (Adcetris) is recommended as an option for treating CD30-positive Hodgkin lymphoma in adults	х	26/07/2017		NHSE commissioned. Approved for addition to the formulary and to be used in line with NICE guidance and commissioning statements

Cortalizumah nagal and	24/05/2017	Certolizumab pegol (Cimzia) alone, or in			26/07/2017	62	CCG commissioned. Approved for addition to the formulary and to be used in line with
Certolizumab pegol and	24/05/2017				26/07/2017	03	
secukinumab for treating active		combination with methotrexate is					NICE guidance and commissioning statements.
psoriatic arthritis after inadequate		recommended as an option for treating active	х				
response to DMARDs (TA445)		psoriatic arthritis in adults					
Afatinib for treating advanced	22/03/2017	Tenofovir alafenamide - was unable to make a			24/05/2017	62	Terminated Appraisal by NICE
squamous non-small-cell lung	22/03/2017	recommendation about the use in the NHS of			24/03/2017	05	Terminated Appraisar by Nice
cancer after platinum-based		afatinib for treating locally advanced or					
chemotherapy (terminated		metastatic squamous non-small-cell lung cancer					
		after platinum-based chemotherapy because no	Х				
appraisal) (TA444)		evidence submission was received from					
		Boehringer Ingelheim.					
Obeticholic acid for treating	26/04/2017	Obeticholic acid (Ocaliva) is recommended,			24/05/2017	20	NHSE commissioned. Approved for addition to the formulary and to be used in line with
primary biliary cholangitis (TA443)					24/05/2017	28	
primary biliary cholangitis (1A443)		within its marketing authorisation, as an option					NICE guidance and commissioning statements
		for treating primary biliary cholangitis in					
		combination with ursodeoxycholic acid for people whose disease has responded					
		inadequately to ursodeoxycholic acid or as	x				
		monotherapy for people who cannot tolerate	^				
		ursodeoxycholic acid. Obeticholic acid is					
		recommended only if the company provides it					
		with the discount agreed in the patient access					
		scheme.					
Ixekizumab for treating moderate	26/04/2017	Ixekizumab (Taltz) is recommended as an option			24/05/2017	28	CCG commissioned. Approved for addition to the formulary and to be used in line with
to severe plaque psoriasis (TA442)		for treating plaque psoriasis in adults	х				NICE guidance and commissioning statements.
Daclizumab for treating	26/04/2017	Daclizumab (Zinbryta) is recommended as an			24/05/2017	28	NHSE commissioned. Approved for addition to the formulary and to be used in line with
relapsing-remitting multiple		option for treating multiple sclerosis in adults	х				NICE guidance and commissioning statements
sclerosis (TA441)							
Pegylated liposomal irinotecan for	26/04/2017	Pegylated liposomal irinotecan (Onivyde), in			24/05/2017	28	Not recommended
treating pancreatic cancer after		combination with 5-fluorouracil and leucovorin,			_ ,, _ , _ ,		
gemcitabine (TA440)		is not recommended, within its marketing					
		authorisation, for treating metastatic	x				
		adenocarcinoma of the pancreas in adults whose					
		disease has progressed after gemcitabine-based					
		therapy.					
			49	0			
						Average	
			% "Yes"	% "N/A"	_	implement	
				•	_	time (days)	
Adherence statistics for 2017-18			100%	0%		46	