

Clinical Policy: Step Therapy

Reference Number: MCPB.ST.00

Effective Date: 01.01.21 Last Review Date: 05.24

Line of Business: Medicare Part B

See <u>Important Reminder</u> at the end of this policy for important regulatory and legal information.

Description

This policy provides a list of drugs that require step therapy. Step therapy is when we require the trial of a preferred therapeutic alternative prior to coverage of a non-preferred drug for a specific indication.

FDA Approved Indication(s)

Various.

Policy/Criteria

This policy does not replace existing Medicare rules and regulations for the applicable agent(s).

I. Approval Criteria (NEW STARTS ONLY – member has not received the drug for the past 365 days)

A. Step Therapy:

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Drug Name	Part B Required Step-Through Agents* By Indication
	*May require prior authorization
Abatacept (Orencia®)	PART B STEP:
	• All indications: a tumor necrosis factor (TNF)
	inhibitor (e.g., infliximab)* (note credit may be given
	if another TNF inhibitor was tried)
Ado-trastuzumab	PART B STEP:
emtansine (Kadcyla®)	• Breast cancer : trastuzumab-based therapy* and a
	taxane* (note some IV chemo may not require prior
	authorization)
Aflibercept (Eylea®,	PART B STEP:
Eylea® HD)	Neovascular (wet) age-related macular
	degeneration (AMD), macular edema following
	retinal vein occlusion (RVO), diabetic macular
	edema (DME), or diabetic retinopathy (DR):
	intravitreal bevacizumab solution
Atezolizumab	PART B STEP:
(Tecentriq®)	• Urothelial carcinoma: member is ineligible for
	platinum-containing chemotherapy as first-line
	systemic therapy* (note some IV chemo may not
	require prior authorization)
	• Non-small cell lung cancer that is high-risk stage
	IIA with programmed death-ligand 1 (PD-L1)



Drug Name	Part B Required Step-Through Agents* By Indication
Axicabtagene ciloleucel (Yescarta®)	expression ≥ 1% OR is recurrent, advanced, or metastatic and anaplastic lymphoma kinase (ALK) or epidermal growth factor receptor (EGFR) mutation negative or unknown: prior platinum-containing chemotherapy (note some IV chemo may not require prior authorization), UNLESS one of the following is met: ○ Request is for use as a single agent, and disease is stage II to IIIA with previous resection ○ Request is for use as a single agent as first-line therapy for tumors that have high PD-L1 expression, defined as PD-L1 ≥ 50% (tumor cells [TC] ≥ 50%) or tumor-infiltrating immune cells (IC) covering ≥ 10% of the tumor area [IC ≥ 10%] ○ Disease is non-squamous, and Tecentriq is prescribed as combination therapy ○ No prior progression on a programmed death receptor-1 (PD-1) or PD-L1 inhibitor (e.g., Tecentriq, nivolumab, pembrolizumab, durvalumab), and Tecentriq is prescribed as single agent as subsequent therapy PART B STEP: • Large B-cell lymphoma: one of the following: ○ 2 lines of systemic therapy that includes rituximab* and one anthracycline-containing regimen (e.g., doxorubicin) ○ First-line chemoimmunotherapy that includes an anti-CD20 monoclonal antibody (e.g.,
	durvalumab), and Tecentriq is prescribed as single
Axicabtagene ciloleucel	PART B STEP:
(Yescarta®)	• Large B-cell lymphoma: one of the following:
	 2 lines of systemic therapy that includes
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	rituximab*) and anthracycline-containing regimen (e.g., doxorubicin), if disease was
	refractory (defined as no complete remission) to
	or relapsed (defined as complete remission
	followed by biopsy-proven disease relapse) no
	more than 12 months after chemoimmunotherapy
	• Relapsed or refractory follicular lymphoma: 2
	lines of systemic therapy that includes a combination
	of an anti-CD20 monoclonal antibody* (e.g., rituximab or Gazyva) and an alkylating agent (e.g.,
	bendamustine, cyclophosphamide, chlorambucil)
	Only for initial treatment dose; subsequent doses will not
	be covered



Drug Name	Part B Required Step-Through Agents* By Indication *May require prior authorization
Bevacizumab (Avastin®,	PART B STEP:
Alymsys [®] , Mvasi [®] ,	
Vegzelma [™] , Zirabev [™])	Oncology indications, if request is for Avastin, Alarmana or Vacatalman Musci and Zirahan
	Alymsys, or Vegzelma: Mvasi and Zirabev PART B STEP:
Brentuximab vedotin	
(Adcetris®)	Lymphomatoid papulosis, B-cell lymphomas other
	than monomorphic post-transplant
	lymphoproliferative disorder (T-cell type): prior
	systemic therapy* (note some IV chemo may not
D 1.	require prior authorization)
Brexucabtagene	PART B STEP:
autoleucel (Tecartus TM)	• Mantle cell lymphoma: 2 to 5 prior regimens that
	included all of the following: anthracycline (e.g.,
	doxorubicin*) or bendamustine*-containing
	chemotherapy; anti-CD20 monoclonal antibody
	therapy (e.g., rituximab*)
	B-cell precursor acute lymphoblastic leukemia: at
	least two prior systemic therapies*
	Only for initial treatment dose; subsequent doses will not
	be covered
Brolucizumab-dbll	PART B STEP:
(Beovu®)	Neovascular (wet) AMD, DME: intravitreal
	bevacizumab solution
Cemiplimab-rwlc	PART B STEP:
(Libtayo [®])	• Cutaneous squamous cell carcinoma: cisplatin*,
	unless curative radiation therapy or surgery is not
	feasible
Certolizumab (Cimzia®)	PART B STEP:
	• All indications: a different TNF inhibitor (e.g.,
	infliximab)* (note credit may be given if another TNF
	inhibitor was tried)
Ciltacabtagene autoleucel	PART B STEP:
(Carvykti [™])	• Lenalidomide-refractory multiple myeloma: at least
	1 prior line of therapy* that included both of the
	following: immunomodulatory agent (e.g.,
	lenalidomide, Pomalyst, Thalomid) and proteasome
	inhibitor (e.g., bortezomib, Kyprolis)
	Only for initial treatment dose; subsequent doses will not
Continuational distriction 1	be covered PART B STEP:
Corticosteroid intravitreal	
implants:	Macular edema following branch or central RVO
dexamethasone	(Ozurdex only): intravitreal bevacizumab solution and
(Ozurdex [®]), fluocinolone	intravitreal corticosteroid injection (e.g., Triesence)
acetonide (Iluvien®,	
Retisert [®] , Yutiq [™])	



Drug Name	Part B Required Step-Through Agents* By Indication
Drug Name	*May require prior authorization
	DME (Ozurdex or Iluvien): intravitreal bevacizumab
	solution and intravitreal corticosteroid injection (e.g.,
	Triesence)
	• Non-infectious uveitis (Ozurdex, Retisert, or Yutiq):
	intravitreal corticosteroid injection (e.g., Triesence)
Corticotropin	PART B STEP:
(H.P. Acthar®, Purified	• All indications, except infantile spasms, if request is
Cortrophin [™] Gel)	for H.P. Acthar: Purified Cortrophin Gel
	<u>IN ADDITION</u> :
	Multiple sclerosis: corticosteroid
Daratumumab	PART B STEP:
(Darzalex [®]),	• Multiple myeloma: 1 prior systemic therapy (e.g.,
daratumumab/	ixazomib*, bortezomib*, carfilzomib*) (note some IV
hyaluronidase-fihj	chemo may not require prior authorization) if
(Darzalex Faspro [™])	prescribed in combination with dexamethasone and
	either lenalidomide, bortezomib, or carfilzomib; OR 2
	prior systemic therapies (e.g., ixazomib*,
	bortezomib*, carfilzomib*) if prescribed as
	monotherapy or in combination with Pomalyst and
	dexamethasone; UNLESS Darzalex is prescribed as
	primary therapy in one of the following ways: o In combination with lenalidomide and
	dexamethasone or bortezomib, melphalan, and
	prednisone, and member is ineligible for
	autologous stem cell transplant (ASCT); or
	 In combination with bortezomib, thalidomide, and
	dexamethasone, and member is eligible for ASCT
	• Systemic light chain amyloidosis (<i>Darzalex only</i>): 1
	prior systemic therapy (e.g., bortezomib*) (note some
	IV chemo may not require prior authorization)
Darbepoetin alfa	PART B STEP:
(Aranesp [®])	All indications: Retacrit
	 If Retacrit is unavailable due to shortage: Epogen
Denosumab	PART B STEP:
(Xgeva [®])	Systemic mastocytosis, hypercalcemia of
	malignancy: zoledronic acid (Zometa)* or
	pamidronate*
Eflapegrastim-xnst	PART B STEP:
(Rolvedon [™])	• All indications: Zarxio, unless member requires ≥ 10
	doses of Zarxio, member is unable to self-administer
	Zarxio due to lack of caregiver or support system for
	assistance with administration and inadequate access
	to healthcare facility or home care interventions



Drug Name	Part B Required Step-Through Agents* By Indication
	 *May require prior authorization If unable to use Zarxio for any of the reasons listed above: Udenyca[^] If unable to use Udenyca[^]: biosimilar pegfilgrastim product (e.g., Fulphila, Fylnetra, Nyvepria, Stimufend, Ziextenzo)
	^ Udenyca refers to all formulations (prefilled syringe, autoinjector, and on-body injector)
Elranatamab-bcmm (Elrexfio [™])	 PART B STEP: Multiple myeloma: 4 prior lines of therapy* that include all of the following: immunomodulatory agent (e.g., lenalidomide, Pomalyst, Thalomid), proteasome inhibitor (e.g., bortezomib, Kyprolis), and anti-CD38 antibody (e.g., Darzalex/Darzalex Faspro, Sarclisa)
Elotuzumab (Empliciti®)	PART B STEP: • Multiple myeloma: prior line of systemic therapy (e.g., bortezomib*) (note some IV chemo may not require prior authorization)
Emapalumab-lzsg	PART B STEP:
(Gamifant [™])	• Primary hemophagocytic lymphohistiocytosis (HLH): conventional HLH therapy* (note some IV chemo may not require prior authorization)
Epoetin alfa (Epogen®,	PART B STEP:
Procrit [®])	All indications: Retacrit
·	 If Retacrit is unavailable due to shortage: Epogen
Faricimab-svoa (Vabysmo	PART B STEP:
®)	• Neovascular (wet) AMD, DME, macular edema following RVO: bevacizumab intravitreal solution
Ferric carboxymaltose	PART B STEP:
(Injectafer®)	• Iron deficiency anemia (IDA) with chronic kidney
	 disease (CKD): Ferrlecit and Venofer If unable to use or failure of Ferrlecit and Venofer: generic Feraheme
	• IDA without CKD: two of the following: Ferrlecit, Infed, Venofer
	If unable to use or failure of Ferrlecit, Infed, and Venofer: generic Feraheme
Ferric derisomaltose	PART B STEP:
(Monoferric®)	 IDA with CKD: Ferrlecit and Venofer If unable to use or failure of Ferrlecit and Venofer: generic Feraheme
	• IDA without CKD: two of the following: Ferrlecit, Infed, Venofer



Drug Name	Part B Required Step-Through Agents* By Indication
	*May require prior authorization O If unable to use or failure of Ferrlecit, Infed, and
	Venofer: generic Feraheme
Ferric pyrophosphate	PART B STEP:
(Triferic [®] , Triferic Avnu [®])	• Iron replacement therapy with hemodialysis-
(Timerie , Timerie Tivia)	dependent CKD: Ferrlecit and Venofer
Ferumoxytol (Feraheme®)	PART B STEP:
returned (returned)	• All indications, if request is for Feraheme: generic
	ferumoxytol
	IN ADDITION:
	• IDA with CKD: Ferrlecit and Venofer
	• IDA without CKD: two of the following: Ferrlecit,
	Infed, Venofer
Filgrastim (Neupogen®,	PART B STEP:
Zarxio [®] , Nivestym TM ,	• All indications, if request is for an agent other than
Granix [®] , Releuko [®])	Zarxio: Zarxio
	 If unable to use Zarxio: Nivestym
	 If unable to use Zarxio and Nivestym and
	request is for Neupogen: biosimilar filgrastim
	product (e.g., Nivestym, Granix, Releuko)
Golimumab (Simponi®,	PART B STEP:
Simponi Aria®)	• All indications: a different TNF inhibitor (e.g.,
	infliximab)* (note credit may be given if another TNF
	inhibitor was tried)
Hyaluronate derivatives:	PART B STEP:
sodium hyaluronate	Osteoarthritis of the knee: intra-articular
(Euflexxa [®] , Gelsyn-3 [™] ,	glucocorticoid injection*, and:
GenVisc [®] 850, Hyalgan [®] ,	 If request is for a product other than
Supartz FX TM , Synojoynt TM ,	Synvisc/Synvisc One or Euflexxa:
Triluron TM , TriVisc TM ,	Synvisc*/Synvisc One* or Euflexxa*
VISCO-3 [™]), hyaluronic	
acid (Durolane®), cross-	
linked hyaluronate (Gel-	
One [®]), hyaluronan (Hymovis [®] , Orthovisc [®] ,	
Monovisc®), hylan	
polymers A and B	
(Synvisc [®] , Synvisc One [®])	
Idecabtagene vicleucel	PART B STEP:
(Abecma TM)	• Multiple myeloma: 2 prior lines of therapy* that
	include all of the following: immunomodulatory agent
	(e.g., lenalidomide, Pomalyst, Thalomid), proteasome
	inhibitor (e.g., bortezomib, Kyprolis), and anti-CD38
	antibody (e.g., Darzalex/Darzalex Faspro, Sarclisa)



Drug Name	Part B Required Step-Through Agents* By Indication
	*May require prior authorization
	Only for initial treatment dose; subsequent doses will not
	be covered
Immune globulins	PART B STEP:
(Asceniv [™] , Bivigam [®] ,	All indications except viral prophylaxis for
Cutaquig [®] , Cuvitru [™] ,	hepatitis A, measles, varicella, or rubella viruses, if
Flebogamma® DIF,	request is for an agent other than Gammagard:
GamaSTAN®,	Gammagard*
GamaSTAN® S/D,	IN ADDITION:
Gammagard® liquid,	Chronic idiopathic demyelinating polyneuropathy:
Gammagard® S/D,	a systemic corticosteroid, unless the member has pure
Gammaked [™] ,	motor symptoms
Gammaplex®, Gamunex®-	Polymyositis, myasthenia gravis, bullous
C, Hizentra [®] , HyQvia [®] ,	pemphigoid, mucous membrane pemphigoid
Octagam®, Panzyga®,	(a.k.a. cicatricial pemphigoid), epidermolysis
Privigen®, Xembify®)	bullosa acquisita: a systemic corticosteroid
	Dermatomyositis: rituximab*
	Idiopathic thrombocytopenic purpura: a systemic
	corticosteroid or Rho(D) immune globulin*
	Pemphigus vulgaris, pemphigus foliaceus: one
	corticosteroid and rituximab*
	Adenosine deaminase (ADA)-severe combined COMPANIE
T 1 (1')	immunodeficiency disorders (SCID): Revcovi*
IncobotulinumtoxinA	PART B STEP:
(Xeomin®)	Upper and lower limb spasticity, cervical dystonia,
	blepharospasm, overactive bladder and urinary incontinence, chronic migraine, primary axillary
	hyperhidrosis: Botox and Dysport
Lisocabtagene maraleucel	PART B STEP:
(Breyanzi [®])	
(Bieyalizi)	• Large B-cell lymphoma: one of the following:
	o 2 lines of systemic therapy that includes an anti- CD20 therapy (e.g., rituximab)* and one
	anthracycline-containing regimen (e.g.,
	doxorubicin)
	First-line chemoimmunotherapy that includes an
	anti-CD20 monoclonal antibody (e.g., rituximab*)
	and anthracycline-containing regimen (e.g.,
	doxorubicin), if disease was refractory (defined as
	no complete remission) to or relapsed (defined as
	complete remission followed by biopsy-proven
	disease relapse) no more than 12 months after
	chemoimmunotherapy
	Relapsed or refractory follicular lymphoma: 2
	lines of systemic therapy that includes a combination
	of an anti-CD20 monoclonal antibody* (e.g.,



Drug Name	Part B Required Step-Through Agents* By Indication *May require prior authorization
	rituximab or Gazyva) and an alkylating agent (e.g., bendamustine, cyclophosphamide, chlorambucil) • Mantle cell lymphoma: 2 lines of systemic therapy that includes an anti-CD20 monoclonal antibody therapy (e.g., rituximab)* and an alkylating agent (e.g., bendamustine, cyclophosphamide, platinum [carboplatin, cisplatin, or oxaliplatin]) Only for initial treatment dose; subsequent doses will not be covered
Lurbinectedin	PART B STEP:
(Zepzelca [™])	• Small cell lung cancer : platinum-containing regimen (e.g., cisplatin, carboplatin)* (note some IV chemo may not require prior authorization)
Luspatercept-aamt (Reblozyl®)	PART B STEP: • Myelodysplastic syndrome with ring sideroblasts < 15% (or ring sideroblasts < 5% with SFB3B1 mutation): erythropoiesis-stimulating agent
Lutetium Lu 177 dotatate (Lutathera®)	PART B STEP: • Neuroendocrine tumor: somatostatin analog (e.g., octreotide, lanreotide), unless member has a well-differentiated grade 3 neuroendocrine tumor
Motixafortide (Aphexda®)	PART B STEP: • Multiple myeloma: plerixafor
Nadofaragene firadenovec-vncg (Adstiladrin®) Natalizumab (Tysabri®,	PART B STEP: • Non-muscle invasive bladder cancer: Bacillus Calmette-Guerin (BCG) treatment* PART B STEP:
Tyruko [®])	 Crohn's disease: a TNF inhibitor (e.g., infliximab*) (note credit may be given if another TNF inhibitor was tried)
Nivolumab (Opdivo®)	 PART B STEP: Non-small cell lung cancer: prior systemic therapy*, UNLESS one of the following is met: Tumor is positive for the tumor mutation burden (TMB) biomarker, or Prescribed in combination with Yervoy for disease with RET rearrangement or unknown/negative mutation status for EGFR, ALK, ROS1, BRAF, MET exon 14 skipping, and NTRK gene fusion, or Prescribed as neoadjuvant treatment Malignant pleural mesothelioma: prior therapy*, unless prescribed in combination with Yervoy



Drug Name	Part B Required Step-Through Agents* By Indication *May require prior authorization
Pegfilgrastim (Neulasta [®] , Fulphila [™] , Fylnetra [®] , Nyvepria [™] , Stimufend [®] , Udenyca [™] , Ziextenzo [™])	 Classical or pediatric Hodgkin lymphoma, anal carcinoma, vulvar cancer, extranodal NK/T-cell lymphoma - nasal type, small cell lung cancer, cervical cancer, pediatric primary mediastinal large B-cell lymphoma: prior therapy* Squamous cell carcinoma of the head and neck: platinum-containing regimen* Urothelial carcinoma: platinum-containing regimen*, unless prescribed as adjuvant treatment and member is at high risk of recurrence after undergoing resection, or member is at high risk of recurrence and did not previously receive a platinum-containing regimen Esophageal squamous cell carcinoma: fluoropyrimidine-based (e.g., 5- fluorouracil, capecitabine) and platinum-based chemotherapy* Gestational trophoblastic neoplasia: platinum/etoposide-containing regimen*, unless disease is methotrexate-resistant and high-risk (note some IV chemo may not require prior authorization) PART B STEP: All indications: Zarxio*, unless member requires ≥ 10 doses of Zarxio, member is unable to self-administer Zarxio due to lack of caregiver or support system for assistance with administration and inadequate access to healthcare facility or home care interventions If unable to use Zarxio for any of the reasons listed above and request is for an agent other than Udenyca²: Udenyca³* If unable to use Udenyca² and request is for Neulasta: biosimilar pegfilgrastim product (e.g., Fulphila, Fylnetra, Nyvepria, Stimufend, Ziextenzo)*
Dombaolissassas	^ Udenyca refers to all formulations (prefilled syringe, autoinjector, and on-body injector) PART B STEP:
Pembrolizumab (Keytruda®)	 Head and neck squamous cell carcinoma: platinum-containing chemotherapy*, unless prescribed as part of combination therapy or prescribed as a single agent for a tumor that expresses PD-L1 with a combined positive score (CPS) ≥ 1



Drug Name	Part B Required Step-Through Agents* By Indication
	*May require prior authorization • Classical Hodgkin lymphoma, primary mediastinal
	large B-cell lymphoma, esophageal squamous cell carcinoma, anal carcinoma, gestational
	trophoblastic neoplasia, extranodal NK/T-cell
	lymphoma, vulvar carcinoma, anaplastic large cell
	lymphoma, small cell lung cancer: at least 1 prior therapy*
	• Endometrial carcinoma: at least 1 prior therapy*, unless prescribed in combination with carboplatin and paclitaxel
	• Tumor mutational burden-high cancer: at least 1
	prior therapy*, unless member has ampullary
	adenocarcinoma or pancreatic adenocarcinoma
	• Cervical cancer: at least 1 prior therapy*, unless
	prescribed in combination with chemotherapy (e.g., paclitaxel/cisplatin, paclitaxel/carboplatin) or chemoradiotherapy
	Microsatellite instability-high/mismatch repair
	deficient cancer: at least 1 prior therapy*, unless
	member has colorectal cancer, ampullary
	adenocarcinoma, gallbladder cancer, gastric cancer,
	esophagogastric junction cancer, intrahepatic/
	extrahepatic cholangiocarcinoma, non-nasopharyngeal head and neck cancer, occult primary tumor,
	pancreatic adenocarcinoma, or small bowel
	adenocarcinoma
	Urothelial carcinoma: platinum-containing
	chemotherapy*, unless ineligible
	 Non-muscle invasive bladder cancer: Bacillus Calmette-Guerin*
	• Squamous cell esophagogastric junction cancer: at
	least 1 prior therapy (if $CPS \ge 10$), unless prescribed
	in combination with fluoropyrimidine- and platinum-
	containing chemotherapy
	• Stage IB, II, or IIIA non-small cell lung cancer:
	platinum-containing chemotherapy*, unless one of the
	following is met: O Disease is PD-L1 positive (tumor proportion)
	score [TPS] $\geq 1\%$)
	 Prescribed as part of combination therapy
	 Prescribed as part of combination therapy as
	neoadjuvant treatment, followed by single-agent
	adjuvant treatment after surgery for patients with



Drug Name	Part B Required Step-Through Agents* By Indication *May require prior authorization
	resectable (tumors ≥ 4 cm or node positive) disease Prescribed as single-agent continuation maintenance therapy if previously given first line as part of a chemotherapy regimen
Polatuzumab vedotin-piiq (Polivy [™])	 (note some IV chemo may not require prior authorization) PART B STEP: Diffuse large B-cell lymphoma, high-grade B-cell lymphoma: 1 prior therapy*, unless all of the
	 following are met: ○ Prescribed for previously untreated disease ○ Prescribed in combination with a rituximab product, cyclophosphamide, doxorubicin, and prednisone ○ Member has an International Prognostic Index score ≥ 2
	• Monomorphic post-transplant lymphoproliferative disorder (B-cell type), HIV-related B-cell lymphoma, follicular lymphoma: 1 prior therapy* (note some IV chemo may not require prior authorization)
Ramucirumab (Cyramza®)	 PART B STEP: Esophageal, esophagogastric junction, and gastric cancer: prior lines of systemic therapy* (note some IV chemo may not require prior authorization)
Ranibizumab (Lucentis [®] , Byooviz [®] , Cimerli [™] , Susvimo [™])	PART B STEP: • Neovascular (wet) AMD, macular edema following RVO, DME, DR, or myopic choroidal neovascularization (mCNV): intravitreal bevacizumab solution
RimabotulinumtoxinB (Myobloc®)	PART B STEP: • Cervical dystonia: Botox and Dysport • Chronic sialorrhea: Xeomin
Rituximab (Rituxan [®] , Riabni [™] , Ruxience [™] , Truxima [®]), rituximab/ hyaluronidase (Rituxan	PART B STEP: • All indications, if request is for Rituxan: Ruxience, Truxima, and Riabni† • All indications, if request is for Riabni: Ruxience*
Hycela [™])	 and Truxima* All oncology indications, if request is for Rituxan Hycela: member has received at least one full dose of Rituxan, Riabni, Ruxience, or Truxima IN ADDITION:
	Rheumatoid arthritis, if request is for Rituxan or Riabni: infliximab*, unless member has had a history of failure of two TNF inhibitors



Drug Name	Part B Required Step-Through Agents* By Indication *May require prior authorization
Romiplostim (Nplate®)	 PART B STEP: Immune thrombocytopenia: systemic corticosteroid (if intolerant or contraindicated to systemic corticosteroids, then immune globulin*) Myelodysplastic syndrome: hypomethylating agent (e.g., azacitadine*, decitabine*) or immunosuppressive therapy (e.g., Atgam*) Chemotherapy-induced thrombocytopenia: prior chemotherapy* (note some IV chemo may not require prior authorization)
Romosozumab-aqqg (Evenity [™])	PART B STEP: • Postmenopausal osteoporosis: bisphosphonate, unless member is very high risk for fracture (recent osteoporotic fracture within the past 12 months, BMD T-score at hip or spine ≤ -3.0, OR BMD T-score at hip or spine ≤ -2.5 and major osteoporotic fracture [i.e., hip, spine, forearm, wrist, humerus])
Sargramostim (Leukine®)	PART B STEP: • All indications: Zarxio
Sipuleucel-T (Provenge®)	PART B STEP: • Prostate cancer: androgen deprivation therapy* (e.g., Zoladex, Vantas, leuprolide, Trelstar, Firmagon)
Talquetamab-tgvs (Talvey [™])	 PART B STEP: Multiple myeloma: 4 prior lines of therapy* that include all of the following: immunomodulatory agent (e.g., lenalidomide, Pomalyst, Thalomid), proteasome inhibitor (e.g., bortezomib, Kyprolis), and anti-CD38 antibody (e.g., Darzalex/Darzalex Faspro, Sarclisa)
Teclistamab-cqyv (Tecvayli®)	 PART B STEP: Multiple myeloma: 4 prior lines of therapy* that include all of the following: immunomodulatory agent (e.g., lenalidomide, Pomalyst, Thalomid), proteasome inhibitor (e.g., bortezomib, Kyprolis, Ninlaro), and anti-CD38 antibody (e.g., Darzalex/Darzalex Faspro, Sarclisa)
Teprotumumab-trbw (Tepezza [™])	PART B STEP: Thyroid eye disease: a systemic corticosteroid
Tisagenlecleucel (Kymriah®)	PART B STEP: • B-cell precursor acute lymphoblastic leukemia: at least two prior systemic therapies* Only for initial treatment dose; subsequent doses will not be covered



Drug Name	Part B Required Step-Through Agents* By Indication
	*May require prior authorization
	• Large B-cell lymphoma: 2 lines of systemic therapy
	that includes rituximab* and one anthracycline-
	containing regimen (e.g., doxorubicin*)
	Only for initial treatment dose; subsequent doses will not
	be covered
	• Relapsed or refractory follicular lymphoma: 2
	lines of systemic therapy that includes a combination
	of an anti-CD20 monoclonal antibody (e.g., rituximab
	or Gazyva)* and an alkylating agent (e.g.,
	bendamustine, cyclophosphamide, chlorambucil)
	Only for initial treatment dose; subsequent doses will not
	be covered
Tocilizumab (Actemra®,	PART B STEP:
Tofidence TM)	Polyarticular juvenile idiopathic arthritis,
Torractice)	rheumatoid arthritis: a TNF inhibitor (e.g.,
	infliximab)* (note credit may be given if another TNF
	inhibitor was tried)
Trastuzumab (Herceptin®,	PART B STEP:
Ontruzant [®] , Herzuma [®] ,	
Ogivri TM , Trazimera TM ,	• All indications, if request is for an agent other than Trazimera: Trazimera*
Kanjinti [™]),	If unable to use Trazimera and request is for
trastuzumab/hyaluronidase	Herceptin or Herceptin Hylecta: biosimilar
(Herceptin Hylecta TM)	trastuzumab product (e.g., Ogivri, Kanjinti) PART B STEP:
Triamcinolone ER	
injection (Zilretta®)	Osteoarthritis of the knee: intra-articular immediate-
	release glucocorticoid injection
Triamcinolone acetonide	PART B STEP:
suprachoroidal injection	• All indications: Triesence (triamcinolone) intravitreal
(Xipere ^{IM})	injection
Vedolizumab (Entyvio®)	PART B STEP:
	• All indications: a TNF inhibitor (e.g., infliximab)*
	(note credit may be given if another TNF inhibitor
	was tried)
Verteporfin (Visudyne®)	PART B STEP:
	Classic subfoveal CNV due to AMD, pathologic
	myopia, or presumed ocular histoplasmosis:
	intravitreal bevacizumab solution

For questions, please reach out to your provider relations.

Important Reminder

This clinical policy has been developed by appropriately experienced and licensed health care professionals based on FDA recommendation(s), peer-reviewed medical literature and evidence-based clinical practice guidelines.

CLINICAL POLICY Step Therapy



The purpose of this clinical policy is to provide a guide to medical necessity, which is a component of the guidelines used to assist in making coverage decisions and administering benefits. It does not constitute a contract or guarantee regarding payment or results. Coverage decisions and the administration of benefits are subject to all terms, conditions, exclusions, and limitations of the coverage documents (e.g., evidence of coverage, certificate of coverage, policy, contract of insurance, etc.), as well as to state and federal requirements and applicable Health Plan-level administrative policies and procedures.

This clinical policy is effective as of the date determined by the Health Plan or responsible business unit. The date of posting may not be the effective date of this clinical policy. This clinical policy may be subject to applicable legal and regulatory requirements relating to provider notification. If there is a discrepancy between the effective date of this clinical policy and any applicable legal or regulatory requirement, the requirements of law and regulation shall govern. The Health Plan retains the right to change, amend or withdraw this clinical policy, and additional clinical policies may be developed and adopted as needed, at any time.

This clinical policy does not constitute medical advice, medical treatment, or medical care. It is not intended to dictate to providers how to practice medicine. Providers are expected to exercise professional medical judgment in providing the most appropriate care, and are solely responsible for the medical advice and treatment of members. This clinical policy is not intended to recommend treatment for members. Members should consult with their treating physician in connection with diagnosis and treatment decisions.

Providers referred to in this clinical policy are independent contractors who exercise independent judgment and over whom the Health Plan has no control or right of control. Providers are not agents or employees of the Health Plan.

This clinical policy is the property of the Health Plan. Unauthorized copying, use, and distribution of this clinical policy or any information contained herein are strictly prohibited. Providers, members, and their representatives are bound to the terms and conditions expressed herein through the terms of their contracts. Where no such contract exists, providers, members and their representatives agree to be bound by such terms and conditions by providing services to members and/or submitting claims for payment for such services.

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