Medication	FDA Indications Note: Although every effort is made to keep this FDA indication list up to date, please consult the web link in the far right column for the most accurate information.	MFC-DC Specifications	Manufacturer's Prescribing Info (Hold CTRL and click on
Absons	Indicated for the treatment of adult reticute with relevand as refusetent monthly language.		link to open)
Abecma (idecabtagene	Indicated for the treatment of adult patients with relapsed or refractory multiple myeloma after four or more prior lines of therapy, including an immunomodulatory agent, a proteasome	***MFC-DC	ABECMA PI
vicleucel)	inhibitor, and an anti-CD38 monoclonal antibody.	Pharmacist	
		Review***	
	Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.	Review	
Actimmune	Indicated for:	***MFC-DC	ACTIMMUNE PI
(interferon gamma-1b)	 reducing the frequency and severity of serious infections associated with Chronic Granulomatous Disease. 	Pharmacist	
	2. delaying time to disease progression in patients with severe, malignant osteoporosis.	Review***	
	Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.		
Adcirca	Indicated for the treatment of pulmonary arterial hypertension (PAH) (WHO Group	Rx by Pulmonologist,	ADCIRCA PI
(tadalafil)	1) to improve exercise ability.	Cardiologist, or Rheumatologist	
Aimovig	Indicated for the preventive treatment of migraine in adults.	Rx by Neurologist	AIMOVIG PI
(erenumab-aooe)		Member must have tried and	
		failed at least 2 previous	
		migraine prophylaxis	
		medications. Examples of	
		migraine prophylaxis	
		medications include, but are not	
		limited to, divalproex,	
		metoprolol, propranolol, timolol,	
		topiramate, amitriptyline,	
		venlafaxine, atenolol.	
Alecensa	Indicated for the treatment of patients with anaplastic lymphoma kinase (ALK)-positive	Rx by Oncologist	ALECENSA PI
(alectinib)	metastatic non-small cell lung cancer (NSCLC) as detected by an FDA-approved test.		
Alunbrig (brigatinib)	Indicated for the treatment of patients with anaplastic lymphoma kinase (ALK)-positive metastatic non-small cell lung cancer (NSCLC) who have progressed on or are intolerant to	Rx by Oncologist	ALUNBRIG PI
(~64611110)	crizotinib.		

Amitiza (lubiprostone)	 Indicated for: chronic idiopathic constipation (CIC) in adults. opioid-induced constipation (OIC) in adult patients with chronic, non-cancer pain, including patients with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation. Limitations of Use: Effectiveness of Amitiza in the treatment of OIC in patients taking diphenylheptane opioids (e.g., methadone) has not been established. irritable bowel syndrome with constipation (IBS-C) in women ≥ 18 years old. 	Failure of at least 2 of the following: docusate, mineral oil, sennosides, psyllium, psyllium /aspartame, calcium polycarbophil, polyethylene glycol 3350, lactulose, methylcellulose	AMITIZA PI
Amondys 45 (casimersen)	Indicated for the treatment of Duchenne muscular dystrophy (DMD) in patients who have a confirmed mutation of the DMD gene that is amenable to exon 45 skipping.	***MFC-DC Pharmacist Review***	AMONDYS 45 PI
Ampyra (dalfampridine)	Indicated to help improve walking in adults with MS.	 Rx by Neurologist. Documentation of MS with ambulatory dysfunction but must be able to walk 25 feet within 8-45 seconds at baseline. Members must have a baseline gait assessment by PT within 90 days of beginning Ampyra. Members must have a repeat evaluation after 3 months on Ampyra. Improvement in walking speed must be documented in order to obtain further refills. Members must not have a history of seizure disorder or renal impairment. 	AMPYRA PI
Apretude (cabotegravir extended- release)	Indicated to reduce the risk of sexually acquired HIV-1 infection in at-risk adults and adolescents weighing at least 35 kg for PrEP (Pre-exposure prophylaxis). PA SUBMISSION REQUIREMENTS: 1. Attestation that patient is considered high-risk for HIV infection 2. Risk-reduction and medication adherence counseling documentation		APRETUDE PI
	 Risk-reduction and medication adherence counseling documentation Negative HIV-1 test prior to initiating therapy and before each injection 		

	 4. Patient will not receive concomitant therapy with any of the following medications due to contraindication and decreased levels of cabotegravir seen with coadministration: a. Anticonvulsants: Carbamazepine, oxcarbazepine, phenobarbital, phenytoin b. Antimycobacterials: Rifampin, rifapentine 		
Austedo (deutetrabenazine)	Indicated for the treatment of: 1. Chorea associated with Huntington's disease. 2. Tardive dyskinesia in adults.		AUSTEDO PI
Ayvakit (avapritinib)	Indicated for the treatment of adults with unresectable or metastatic gastrointestinal stromal tumor (GIST) harboring a platelet-derived growth factor receptor alpha (PDGFRA) exon 18 mutation, including PDGFRA D842V mutations.	Rx by Oncologist	AYVAKIT PI
Balversa (erdafitinib)	Indicated for the treatment of adult patients with locally advanced or metastatic urothelial carcinoma that has susceptible FGFR3 or FGFR2 genetic alterations and progressed during or following at least one line of prior platinum-containing chemotherapy including within 12 months of neoadjuvant or adjuvant platinum-containing chemotherapy.	Rx by Oncologist	BALVERSA PI
Benlysta (belimumab)	Indicated for the treatment of adult patients with active, autoantibody-positive, systemic lupus erythematosus who are receiving standard therapy.		BENLYSTA PI
Bethkis (tobramycin inh sol)	Indicated for management of cystic fibrosis patients with <i>Pseudomonas aeruginosa.</i> *** Safety and efficacy have not been demonstrated in patients under the age of six years, patients with a forced expiratory volume in one second (FEV1) less than 40% or greater than 80% predicted, or patients colonized with Burkholderia cepacia.	Rx by Pulmonologist	BETHKIS PI
Bosulif (bosutinib)	 Indicated for the treatment of adult patients with: Newly diagnosed chronic phase Ph+ chronic myelogenous leukemia (CML). This indication is approved under accelerated approval based on molecular and cytogenetic response rates. Continued approval for this indication may be contingent upon verification and confirmation of clinical benefit in an ongoing long-term follow up trial. Chronic, accelerated, or blast phase Ph+ CML with resistance or intolerance to prior therapy. 	Rx by Oncologist	BOSULIF PI
Botox (onabotulinumtoxin A)	 Indicated for: treatment of overactive bladder (OAB) with symptoms of urge urinary incontinence, urgency, and frequency, in adults who have an inadequate response to or are intolerant of an anticholinergic medication. treatment of urinary incontinence due to detrusor overactivity associated with a neurologic condition [e.g., spinal cord injury (SCI), multiple sclerosis (MS)] in adults who have an inadequate response to or are intolerant of an anticholinergic medication. prophylaxis of headaches in adult patients with chronic migraine (≥15 days per month with headache lasting 4 hours a day or longer). treatment of spasticity in adult patients. 	Rx by Neurologist, Urologist, Ophthalmologist Botox will NOT be approved for cosmetic purposes.	BOTOX PI

		1	1
	5. treatment of spasticity in pediatric patients 2 to 17 years of age, excluding spasticity		
	caused by cerebral palsy.		
	6. treatment of cervical dystonia in adult patients, to reduce the severity of abnormal head		
	position and neck pain.		
	7. treatment of severe axillary hyperhidrosis that is inadequately managed by topical agents		
	in adult patients.		
	8. treatment of blepharospasm associated with dystonia in patients ≥12 years of age.		
	9. treatment of strabismus in patients ≥12 years of age.		
Braftovi	Indicated, in combination with binimetinib, for the treatment of patients with unresectable or	Rx by Oncologist	BRAFTOVI PI
(encorafenib)	metastatic melanoma with a BRAF V600E or V600K mutation, as detected by an FDA-approved		
	test.		
Breyanzi	Indicated for:	***MFC-DC	BREYANZI PI
(lisocabtagene	1. treatment of adult patients with relapsed or refractory large B-cell lymphoma after two	Dhawaasist	
maraleucel)	or more lines of systemic therapy, including:	Pharmacist	
	a. Diffuse large B-cell lymphoma (DLBCL) not otherwise specified (including DLBCL	Review***	
	arising from indolent lymphoma)		
	b. High-grade B-cell lymphoma		
	c. Primary mediastinal large B-cell lymphoma		
	d. Follicular lymphoma grade 3B.		
	Continued approval for this indication may be continuent upon varification and description of		
	Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.		
Cabometyx	Indicated for the treatment of patients with advanced renal cell carcinoma.	Rx by Oncologist	CABOMETYX PI
(cabozantinib)	 Indicated for the treatment of patients with hepatocellular carcinoma (HCC) who have 	KX by Offcologist	CABOWETTA PI
(Cabozantinib)	been previously treated with sorafenib.		
Cialis	Indicated for the treatment of:	Rx by Pulmonologist or	CIALIS PI
(tadalafil)	1. erectile dysfunction (ED) – (<i>drug coverage not provided for this indication</i>)	Cardiologist	CIALIS FI
(tauaiaiii)	2. the signs and symptoms of benign prostatic hyperplasia (BPH)	Cardiologist	
	3. ED and the signs and symptoms of BPH (ED/BPH)		
Cinryze	Indicated for routine prophylaxis against angioedema attacks in adults, adolescents, and	******	CINRYZE PI
(C1 Esterase Inhibitor	pediatric patients (6 years of age and older) with Hereditary Angioedema.	***MFC-DC	SHWIZETT
[Human])		Pharmacist	
1,	Continued approval for this indication may be contingent upon verification and description of		
	clinical benefit in a confirmatory trial.	Review***	
Cometriq	Indicated for treatment of progressive, metastatic medullary thyroid cancer.	Rx by Oncologist	COMETRIQ PI
(cabozantinib)	,,	, ,	
Cosela	Indicated to decrease the incidence of chemotherapy-induced myelosuppression in adult	Rx by Oncologist	COSELA PI
(trilaciclib)	patients when administered prior to a platinum/etoposide-containing regimen or topotecan-		
,	containing regimen for extensive-stage small cell lung cancer.		
Cotellic	Indicated for the treatment of patients with unresectable or metastatic melanoma with a BRAF	Rx by Oncologist	COTELLIC PI
(cobimetinib)	V600E or V600K mutation, in combination with vemurafenib.		
,			

Cutaquig (Immune Globulin Subcutaneous (Human) - hipp)	Indicated for treatment of primary humoral immunodeficiency (PI) in adults.	Rx by Immunologist	CUTAQUIG PI
Darzalex Faspro (daratumumab and hyaluronidase-fihj, SQ admin)	 Indicated for the treatment of adult patients with multiple myeloma: in combination with bortezomib, melphalan and prednisone in newly diagnosed patients who are ineligible for autologous stem cell transplant. in combination with lenalidomide and dexamethasone in newly diagnosed patients who are ineligible for autologous stem cell transplant and in patients with relapsed or refractory multiple myeloma who have received at least one prior therapy. in combination with bortezomib and dexamethasone in patients who have received at least one prior therapy. as monotherapy, in patients who have received at least three prior lines of therapy including a proteasome inhibitor (PI) and an immunomodulatory agent or who are double-refractory to a PI and an immunomodulatory agent. 	Rx by Oncologist	DARZALEX FASPRO PI
DESMOPRESSIN NASAL SPRAY PRODUCTS: DDAVP spray- 0.01% Stimate spray- 1.5 mg/mL	DDAVP is indicated for: 1. antidiuretic replacement therapy in the management of central cranial diabetes insipidus. 2. treatment of transient polyuria and polydipsia post head trauma or surgery in the pituitary region. Stimate is indicated for: 1. hemophilia A with Factor VIII coagulant activity levels greater than 5% - will stop bleeding in patients with hemophilia A with episodes of spontaneous or trauma-induced injuries such as hemarthroses, intramuscular hematomas or mucosal bleeding. 2. mild to moderate classic von Willebrand's disease (Type I) with Factor VIII levels greater than 5% - will stop bleeding in patients with episodes of spontaneous or trauma-induced injuries such as hemarthroses, intramuscular hematomas, mucosal bleeding or menorrhagia.	1. STIMATE: Hemophilia A with factor VIII coagulant activity greater than 5%: ➤ *peri-operatively to prevent bleeding ➤ to treat spontaneous or trauma induced bleeding ***Note- Patients with factor VIII levels equal to or less than 5% or patients who have factor VIII antibodies are not candidates for the drug. It is contraindicated in patients under 3 months old. It is NOT indicated for Hemophilia B. 2. STIMATE: Patients with von Willebrand's Disease (type I) with factor VIII coagulant activity greater than 5%: ➤ used peri-operatively to prevent bleeding. ➤ to treat spontaneous or trauma induced bleeding. ***Note- The drug is NOT indicated for treatment of	STIMATE PI

Dexcom G6 Continuous Glucose Monitoring (CGM) System	Indicated for the management of diabetes in persons age 2 years and older.	severe classic von Willebrand's Disease (type I) or when there is evidence of an abnormal molecular form of Factor VIII antigen Rx by Endocrinologist. Please click link below for CGM Policy: Continuous Glucose Monitoring Devices	DEXCOM G6 PI
Dificid (fidaxomicin)	Indicated for the treatment of <i>Clostridium difficile</i> -associated diarrhea in adults (≥18 years of age).	Pt must have documented failures with both metronidazole and vancomycin, or contraindication(s) to the use of these agents.	DIFICID PI
Doptelet (avatrombopag)	Indicated for the treatment of thrombocytopenia in adult patients with chronic liver disease who are scheduled to undergo a procedure.	Rx by Hematologist A recent (less than 1 month old) platelet count must be supplied with the clinical request, as well as information regarding the planned procedure.	DOPTELET PI
Dupixent (dupilumab)	 Indicated for: treatment of adult and pediatric patients aged 6 months and older with moderate-to-severe atopic dermatitis whose disease is not adequately controlled with topical prescription therapies or when those therapies are not advisable. DUPIXENT can be used with or without topical corticosteroids. add-on maintenance treatment of adult and pediatric patients aged 6 years and older with moderate-to-severe asthma with an eosinophilic phenotype or with oral corticosteroid dependent asthma. add-on maintenance treatment in adult patients with inadequately controlled chronic rhinosinusitis with nasal polyposis. treatment of adult and pediatric patients aged 12 years and older, weighing at least 40 kg, with eosinophilic esophagitis 	Rx by Allergist or Dermatologist	DUPIXENT PI
Egrifta SV (tesamorelin injection) Eligard	Indicated for the reduction of excess abdominal fat in HIV-infected adult patients with lipodystrophy. see Leuprolide		EGRIFTA SV PI
(leuprolide SQ)	See Leupi Oilue		
Elzonris (tagraxofusp-erzs)	Indicated for the treatment of blastic plasmacytoid dendritic cell neoplasm (BPDCN) in adults and in pediatric patients 2 years and older.	Rx by Oncologist	ELZONRIS PI
Emgality (galcanezumab-gnlm)	A calcitonin-gene related peptide antagonist indicated for the preventative treatment of migraine and treatment of episodic cluster headache.	Rx by Neurologist 1. Member must have tried and failed at least	EMGALITY PI

Empaveli	Indicated for the treatment of adult patients with paroxysmal nocturnal hemoglobinuria.	2 previous migraine prophylaxis medications. Examples of migraine prophylaxis medications include, but are not limited to, divalproex, metoprolol, propranolol, timolol, topiramate, amitriptyline, venlafaxine, atenolol. 2. Medication not to be used with another CGRP antagonist or inhibitor used for preventative treatment of migraines (Ajovy, Nurtec ODT, Vyepti (eptinezumab-jjmr))	EMPAVELI PI
(pegcetacoplan)		***MFC-DC Pharmacist	<u>EWIT / WEET T</u>
	Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.	Review***	
Enhertu	Indicated for the treatment of adult patients with unresectable or metastatic HER2-positive	Rx by Oncologist	ENHERTU PI
(fam-trastuzumab deruxtecan-nxk)	breast cancer who have received two or more prior anti-HER2-based regimens in the metastatic setting.	Tix by officiologist	<u>EMILITOTI</u>
Enspryng (satralizumab-mwge)	An interleukin-6 (IL-6) receptor antagonist indicated for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult patients who are anti-aquaporin-4 (AQP4) antibody positive.	Rx by Neurologist	ENSPRYNG PI
	PA SUBMISSION REQUIREMENTS: 1. Documentation of positive test for AQP4-IgG antibodies 2. Documentation of at least one relapse in the last 12 months OR 2 or more relapses that require rescue therapy in the last 24 months 3. Documentation of inadequate response, contraindication or intolerance to rituximab or Truxima (rituximab-abbs)		
Epclusa (sofosbuvir/ velpatasvir)	Indicated for the treatment of patients with chronic HCV genotype (GT) 1, 2, 3, 4, 5 or 6 infection without cirrhosis and with compensated cirrhosis (Child-Pugh A) and with decompensated cirrhosis for use in combination with ribavirin (Child-Pugh B and C). EPCLUSA	To get the latest copy of the hepatitis C Prior Authorization form, please click the link below:	EPCLUSA PI
SEE SPECIAL NOTE REGARDING PA REQUIREMENTS AND FORM	is also indicated for the treatment of adult and pediatric patients 6 years and older or weighing at least 17 kg with HCV genotypes 1 2, 3, 4, 5, or 6 infection, who previously have been treated with a regimen containing an HCV NS5A inhibitor or an NS5B protease inhibitor, but not both.	HEPATITIS C PA FORM	
*******	PA SUBMISSION REQUIREMENTS:		

	 A completed Hepatitis C Prior Authorization Form (see link on right). Medical records including: a. Most recent office visit note(s) which includes:		
Erwinaze (asparaginase Erwinia chrysanthemi)	BRAND Epclusa. Indicated as a component of a multi-agent chemotherapeutic regimen for the treatment of patients with ALL who have developed hypersensitivity to <i>E. coli</i> -derived asparaginase.	Rx by Oncologist	ERWINAZE PI
Esbriet (pirfenidone)	Indicated for the treatment of idiopathic pulmonary fibrosis IPF.	Rx by Pulmonologist or Cardiologist	ESBRIET PI
Evkeeza (evinacumab-dgnb)	Indicated as an adjunct to other low-density lipoprotein-cholesterol (LDL-C) lowering therapies for the treatment of adult and pediatric patients, aged 12 years and older, with homozygous familial hypercholesterolemia (HoFH).	Must submit genetic testing confirming homozygous familial hypercholesterolemia (HoFH).	EVKEEZA PI
Exkivity (mobocertinib)	Indicated for the treatment of adult patients with locally advanced or metastatic non-small cell lung cancer (NSCLC) with epidermal growth factor receptor (EGFR) exon 20 insertion mutations, as detected by an FDA-approved test, whose disease has progressed on or after platinum-based chemotherapy.		EXKIVITY PI
Fasenra (benralizumab)	Indicated for the add-on maintenance treatment of patients with severe asthma aged 12 years and older, and with an eosinophilic phenotype.	Rx by Pulmonologist or Allergist	FASENRA PI
Fentanyl SEE SPECIAL NOTE REGARDING PA REQUIREMENTS AND FORM ************	Indicated for the management of pain in opioid-tolerant patients, severe enough to require daily, around-the-clock, long-term opioid treatment and for which alternative treatment options are inadequate	All long-acting opioids require Prior Authorization (PA). The PA form can be accessed using the following link: Opioid PA Form	FENTANYL PI
Firazyr (icatibant)	Indicated for the treatment of acute attacks of hereditary angioedema (HAE) in adults ≥18 years of age (self-administered by the patient upon recognition of symptoms of an HAE attack after training under the guidance of a healthcare professional).	Rx by Allergist or ENT	FIRAZYR PI
Fotivda (tivozanib)	Indicated for the treatment of adult patients with relapsed or refractory advanced renal cell carcinoma (RCC) following two or more prior systemic therapies.	Rx by Hematology Oncology	FOTIVDA PI

Gavreto (pralsetinib)	Indicated for the treatment of adult patients with metastatic rearranged during transfection (RET) fusion- positive non-small cell lung cancer (NSCLC) as detected by an FDA approved test Rx by Oncologist	Rx by Oncologist	GAVRETO PI
Gralise (gabapentin)	Indicated for the management of Postherpetic Neuralgia (PHN). (GRALISE is not interchangeable with other gabapentin products because of differing 6 pharmacokinetic profiles that affect the frequency of administration)		GRALISE PI
Growth Hormone	See Norditropin; See Serostim		
Haegarda (C1 Esterase Inhibitor SubQ (Human))	Indicated for routine prophylaxis to prevent Hereditary Angioedema (HAE) attacks in adolescent and adult patients.	Rx by Allergist/Immunologist	HAEGARDA PI
Hycamtin caps (topotecan)	Indicated for treatment of patients with relapsed small cell lung cancer (SCLC) in patients with a prior complete or partial response and who are at least 45 days from the end of first-line chemotherapy.	Rx by Oncologist	HYCAMTIN PI
Ibrance (palbociclib)	 Indicated for the treatment of hormone receptor (HR)-positive, human epidermal growth factor receptor 2 (HER2)-negative advanced or metastatic breast cancer in combination with: 1. an aromatase inhibitor as initial endocrine based therapy in postmenopausal women or in men; 2. fulvestrant in patients with disease progression following endocrine therapy. 	Rx by Oncologist	IBRANCE PI
Iclusig (ponatinib)	 Indicated for: treatment of adult patients with chronic phase, accelerated phase, or blast phase chronic myeloid leukemia (CML) or Ph+ ALL for whom no other tyrosine kinase inhibitor (TKI) therapy is indicated. treatment of adult patients with T315I-positive CML (chronic phase, accelerated phase, or blast phase) or T315I-positive Philadelphia chromosome positive acute lymphoblastic leukemia (Ph+ ALL). 	Rx by Oncologist	ICLUSIG PI
Imbruvica (ibrutinib)	 Indicated for: Mantle cell lymphoma (MCL) who have received at least one prior therapy.	Rx by Oncologist	IMBRUVICA PI

Jakafi	Indicated for:	Rx by Hematologist/Oncologist	JAKAFI PI
(ruxolitinib)	1. treatment of patients with intermediate or high-risk myelofibrosis, including primary	, , , ,	
	myelofibrosis, post-polycythemia vera myelofibrosis, and post-essential thrombocythemia		
	myelofibrosis in adults.		
	2. treatment of adults with polycythemia vera who have had an inadequate response to or		
	are intolerant of hydroxyurea.		
	3. treatment of steroid-refractory acute graft-versus-host disease in adult and pediatric		
	patients 12 years and older.		
Jivi	Recombinant DNA-derived, Factor VIII concentrate indicated for use in previously treated	Rx by Hematologist	JIVI PI
(empagliflozin)	adults and adolescents (12 years of age and older) with hemophilia A (congenital Factor VIII		
	deficiency) for:		
	On-demand treatment and control of bleeding episodes.		
	Perioperative management of bleeding.		
to a description	3. Routine prophylaxis to reduce the frequency of bleeding episodes.	Bullion Conditations on	HIVTA DID DI
Juxtapid	Indicated as an adjunct to a low-fat diet and other lipid-lowering treatments, including LDL	Rx by Cardiology or	JUXTAPID PI
(lomitapide)	apheresis where available, to reduce low-density lipoprotein cholesterol (LDL-C), total	Endocrinologist	
	cholesterol (TC), apolipoprotein B (apo B), and non-high density lipoprotein cholesterol (non-		
Jynarque	HDL-C) in patients with homozygous familial hypercholesterolemia (HoFH). Indicated to slow kidney function decline in adults at risk of rapidly progressing autosomal	Rx by Nephrologist	JYNARQUE PI
(tolvaptan)	dominant polycystic kidney disease (ADPKD).	KX by Nephrologist	JYNARQUE PI
Kalbitor	Indicated for treatment of acute attacks of hereditary angioedema (HAE) in patients 12 years	Rx by Immunologist or Allergist	KALBITOR PI
(ecallantide)	of age and older.	ix by illimunologist of Allergist	KALBITOK FI
Kalydeco	Indicated for the treatment of cystic fibrosis (CF) in patients age 6 months and older who have	Rx by Pulmonologist	KALYDECO PI
(ivacaftor)	one mutation in the CFTR gene that is responsive to ivacaftor potentiation based on clinical	Tix by Full Tollogist	IVIET DECOTE
(,	and/or in vitro assay data. If the patient's genotype is unknown, an FDA-cleared CF mutation		
	test should be used to detect the presence of a CFTR mutation followed by verification with bi-		
	directional sequencing when recommended by the mutation test instructions for use.		
Kerendia	Indicated to reduce the risk of sustained eGFR decline, end stage kidney disease,		KERENDIA PI
(finerenone)	cardiovascular death, non-fatal myocardial infarction, and hospitalization for heart failure in		
	adult patients with chronic kidney disease (CKD) associated with type 2 diabetes (T2D).		
	PA SUBMISSION REQUIREMENTS:		
	1. Type II Diabetes Mellitus diagnosis		
	2. Laboratory results and clinical note		
	Serum creatinine ≤ 5.0 mEq/L		
	4. eGFR $\ge 25 \text{ mL/min/1.73 m}^2$		
	Urine albumin-to-creatinine ratio ≥ 30mg/g		
	6. Concomitant use with maximum tolerated doses of ACE-Inhibitor or ARB		
	7. Intolerance or contraindication to ACE-inhibitor or ARB if not taking		
	8. Intolerance, failed trial or contraindication to both Jardiance and Invokana		
Kisqali	Indicated in combination with:	Rx by Oncologist	KISQALI PI
(ribociclib)	1. an aromatase inhibitor for the treatment of pre/perimenopausal or postmenopausal		
	women with hormone receptor (HR)-positive, human epidermal growth factor receptor 2		

	(HER2)-negative advanced or metastatic breast cancer, as initial endocrine-based therapy;		
	or		
	2. fulvestrant for the treatment of postmenopausal women with HER2-positive, HER2-		
	negative advanced or metastatic breast cancer, as initial endocrine based therapy or		
	following disease progression on endocrine therapy.		
Kymriah	Indicated for:	Rx by Oncologist	KYMRIAH PI
(tisagenlecluecel)	Patients up to 25 years of age with B-cell precursor acute lymphoblastic leukemia (ALL)	in by encologist	<u> </u>
(wagamaaaa,	that is refractory or in second or later relapse.		
	2. Adult patients with relapsed or refractory (r/r) large B-cell lymphoma after two or more		
	lines of systemic therapy including diffuse large B-cell lymphoma (DLBCL) not otherwise		
	specified, high grade B-cell lymphoma and DLBCL arising from follicular lymphoma.		
	<u>Limitation of Use</u> : KYMRIAH is not indicated for treatment of patients with primary central		
	nervous system lymphoma.		
	MedStar Family Choice considers Kymriah medically necessary when all of the following		
	conditions are met:		
	1. Recipient has relapsed or refractory B-cell ALL, defined as		
	Second or greater bone marrow relapse; OR		
	 Any bone marrow relapse after allogeneic stem cell transplantation; OR 		
	 Primary refractory as defined by not achieving a complete remission after 2 		
	cycles of a standard chemotherapy regimen or chemorefractory as defined		
	by not achieving a complete remission after 1 cycle of standard		
	chemotherapy for relapsed leukemia; OR		
	 Patients with Philadelphia chromosome positive (Ph+) ALL are eligible if they 		
	are intolerant to or have failed 2 lines of tyrosine kinase inhibitor therapy		
	(TKI), or if TKI therapy is contraindicated; AND		
	2. Recipient is 25 years of age or younger; AND		
	3. Documentation of CD19 tumor expression; AND		
	4. Performance score on Karnofsky or Lansky Scale is greater than or equal to 50%; AND		
	5. Life expectancy > 12 weeks; AND		
	Adequate cardiac, pulmonary, and other organ function (as determined by protocol from treatment facility); AND		
	7. The treatment facility that dispenses and administers Kymriah is enrolled and		
	complies with the Risk Evaluation and Mitigation Strategy; AND		
	8. One-time, single administration with dosing in accordance with the FDA label.		
	Kymriah is considered investigational and not medically necessary when the above medically		
	necessary criteria are not met, and for all other indications, including but not limited to:		
	 Isolated extra-medullary disease relapse; or 		
	2. Patients with Burkitt's lymphoma/leukemia (i.e. patients with mature B-cell ALL,		
	leukemia with B-cell [slg positive and kappa or lambda restricted positivity] ALL, with		
	FAB L3 morphology and /or a MYC translocation); or		
	3. Prior malignancy, except carcinoma in situ of the skin or cervix treated with curative		
	intent and with no evidence of active disease; or		

	 Treatment with any other chimeric antigen receptor therapy or genetically modified T cell therapy; or Any active uncontrolled infection; or Hepatitis B or C (if viral load is detectable); or Human Immunodeficiency Virus (HIV); or Presence of grade 2 to 4 acute or extensive chronic graft-versus-host disease (GVHD); or Active CNS involvement by malignancy, defined by CNS-3 per NCCN guidelines. 		
Latuda	Indicated for:		LATUDA PI
(lurasidone	Schizophrenia in adults and adolescents (13 to 17 years).		
hydrochloride)	Depressive episode associated with Bipolar I Disorder (bipolar depression) in adults		
ilyar bemoriae)	and pediatric patients (10 to 17 years) as monotherapy.		
	3. Depressive episode associated with Bipolar I Disorder (bipolar depression) in adults as		
	adjunctive therapy with lithium or valproate.		
LEUPROLIDE	Indicated for:		ELIGARD PI
PRODUCTS:	1. palliative treatment for advanced prostate cancer (Eligard).		<u>EE.O/MOTT</u>
TROBUCIO.	2. treatment of pediatric patients with central precocious puberty (Lupron Depot- PED).		LUPRON 3.75 mg PI
Eligard	3. treatment of endometriosis (Lupron and Lupron Depot).		<u> </u>
(leuprolide SQ)	4. Management of endometriosis, including pain relief, recurrence symptoms and reduction		LUPRON DEPOT 11.25 MG
(leapronae 3Q)	of endometriotic lesions, (Lupron and Lupron Depot).		PI
	5. uterine leiomyomata (fibroids) along with concurrent iron therapy in preparation for		11
Lupron	surgery [duration of treatment should be for 6 months or less (Lupron and Lupron		
(leuprolide acetate)	Depot)].		LUPRON DEPOT-PED PI
(leuprollue acetate)	Depotyj.		LOFKON BLFOT-FLB FI
Lupron Depot			
(leuprolide acetate for			
depot suspension)			
depot suspension)			
Lupron Depot-PED			
(leuprolide acetate for			
depot suspension)			
depot suspension)			
Librium	Indicated for the management of anxiety disorders or for the short-term relief of symptoms of		LIBRIUM PI
(chlordiazepoxide)	anxiety, withdrawal symptoms of acute alcoholism, and preoperative apprehension and		<u>Libition 1 1</u>
(cinoralazepoxiae)	anxiety.		
Libtayo	Indicated for the treatment of patients with metastatic cutaneous squamous cell carcinoma	Rx by Oncologist	LIBTAYO PI
(cemiplimab-rwlc)	(CSCC) or locally advanced CSCC who are not candidates for curative surgery or curative	The Sy Officerogist	<u> </u>
(cernipiiiiab-i wie)	radiation.		
Livtencity	A cytomegalovirus (CMV) pUL97 kinase inhibitor indicated for the treatment of adults and		LIVTENCITY PI
(maribavir)	pediatric patients (12 years of age and older and weighing at least 35 kg) with post-transplant		LIVILINCITIFI
(manbavii)	CMV infection/disease that is refractory to treatment (with or without genotypic resistance)		
	with ganciclovir, valganciclovir, cidofovir or foscarnet.		
	with gandidovir, valgandidovir, duolovir or loscarnet.		

LO Loestrin Fe (norethindrone, ethinyl estradiol and ferrous fumarate)	See Oral Contraceptive	ANY OCP on prior authorization requires documentation demonstrating a compelling reason why formulary OCPs cannot be used [ex: intolerance, prior side effects, failures, etc. documented after a 3-month trial of formulary OCPs]	LO LOESTRIN PI
Lorbrena (lorlatinib)	Indicated for the treatment of patients with anaplastic lymphoma kinase (ALK)-positive metastatic non-small cell lung cancer (NSCLC) whose disease has progressed on: 1. crizotinib and at least one other ALK inhibitor for metastatic disease; or 2. alectinib as the first ALK inhibitor therapy for metastatic disease; or 3. ceritinib as the first ALK inhibitor therapy for metastatic disease.	Rx by Oncologist	LORBRENA PI
Lovaza (omega-3-acid ethyl esters)	Indicated as an adjunct to diet to reduce triglyceride (TG) levels in adult patients with severe (≥500 mg/dL) hypertriglyceridemia.		LOVAZA PI
Lumakras (sotorasib)	Indicated for the treatment of adult patients with KRAS G12C-mutated locally advanced or metastatic non-small cell lung cancer (NSCLC), as determined by an FDA-approved test, who have received at least one prior systemic therapy.	Rx by Oncologist	LUMAKRAS PI
Lumoxiti (moxetumomab pasudotox-tdfk)	Indicated for the treatment of adult patients with relapsed or refractory hairy cell leukemia (HCL) who received at least 2 prior systemic therapies, including treatment with a purine nucleoside analog.	Rx by Oncologist	LUMOXITI PI
Lupron and Lupron Depot	See Leuprolide		
Lupkynis (voclosporin)	Indicated in combination with a background immunosuppressive therapy regimen for the treatment of adult patients with active lupus nephritis.		LUPKYNIS PI
Lybalvi (olanzapine and samidorphan)	Indicated for: 1. Schizophrenia in adults 2. Bipolar I disorder in adults a. Acute treatment of manic or mixed episodes as monotherapy and as adjunct to lithium or valproate b. Maintenance monotherapy treatment		LYBALVI PI
	PA SUBMISSION REQUIREMENTS: 1. Laboratory findings and clinical notes 2. Cholesterol levels, fasting blood glucose, A1c, current weight and BMI, complete blood count 3. Urine drug screen 4. No known substance use disorder 5. Do not initiate within 14 days of opioid medication use 6. 4-week trial and failure of at least two formulary atypical antipsychotic agents		

	7. If clinically stable on olanzapine provide documentation of metabolic syndrome,		
	weight gain, or glucose intolerance		
Lynparza (olaparib)	 Indicated for: First-Line Maintenance Treatment of BRCA-mutated Advanced Ovarian Cancer - For the maintenance treatment of adult patients with deleterious or suspected deleterious germline or somatic BRCA-mutated (gBRCAm or sBRCAm) advanced epithelial ovarian, fallopian tube or primary peritoneal cancer who are in complete or partial response to first-line platinum-based chemotherapy. Select patients for therapy based on an FDA-approved companion diagnostic for Lynparza. Maintenance Treatment of Recurrent Ovarian Cancer- For the maintenance treatment of adult patients with recurrent epithelial ovarian, fallopian tube, or primary peritoneal cancer, who are in complete or partial response to platinum-based chemotherapy. Advanced gBRCA-mutated Ovarian Cancer After 3 or More Lines of Chemotherapy- For the treatment of adult patients with deleterious or suspected deleterious germline BRCA-mutated (gBRCAm) advanced ovarian cancer who have been treated with 3 or more prior lines of chemotherapy. Select patients for therapy based on an FDA-approved companion diagnostic for LYNPARZA. Germline BRCA-mutated HER2-negative Metastatic Breast Cancer — In patients with deleterious or suspected deleterious gBRCAm, human epidermal growth factor receptor 2 (HER2)-negative metastatic breast cancer who have been treated with chemotherapy in the neoadjuvant, adjuvant or metastatic setting. Patients with hormone receptor (HR)-positive breast cancer should have been treated with a prior endocrine therapy or be considered inappropriate for endocrine therapy. Select patients for therapy based on FDA-approved companion diagnostic for Lynparza. 	Rx by Oncologist	LYNPARZA PI
Macrilen (macimorelin)	Indicated for the diagnosis of adult growth hormone deficiency.	Rx by Endocrinologist	MACRILEN PI
Mavyret (glecaprevir and pibrentasvir) SEE SPECIAL NOTE REGARDING PA REQUIREMENTS AND FORM *************	Indicated for the treatment of patients with chronic HCV genotype (GT) 1, 2, 3, 4, 5 or 6 infection without cirrhosis and with compensated cirrhosis (Child-Pugh A). MAVYRET is also indicated for the treatment of adult and pediatric patients 12 years and older or weighing at least 45 kg with HCV genotype 1 infection, who previously have been treated with a regimen containing an HCV NS5A inhibitor or an NS3/4A protease inhibitor, but not both. PA SUBMISSION REQUIREMENTS: 1. A completed Hepatitis C Prior Authorization Form (see link on right). 2. Medical records including: a. Most recent office visit note(s) which includes: i. detail on all previous hepatitis C treatments; if none, the note must say "treatment naïve." ii. Child-Pugh score (if cirrhotic). iii. social history with detail provided on use of ETOH and/or illicit substances. 3. Laboratory studies including: a. a recent (less than 6 months old) baseline viral load. b. genotype c. (if applicable) HIV viral load and/or hepatitis B viral load.	To get the latest copy of the hepatitis C Prior Authorization form, please click the link below: HEPATITIS C PA FORM	MAVYRET PI

Mekinist	Indicated:	Rx by Oncologist	MEKINIST PI
(trametinib)	 BRAF V600E or V600K Mutation-Positive Unresectable or Metastatic Melanoma - as a single agent in BRAF-inhibitor treatment-naïve patients or in combination with dabrafenib, for the treatment of patients with unresectable or metastatic melanoma with BRAF V600E or V600K mutations, as detected by an FDA-approved test. Adjuvant Treatment of BRAF V600E or V600K Mutation-Positive Melanoma-indicated in combination with dabrafenib, for the adjuvant treatment of patients with melanoma with BRAF V600E or V600K mutations as detected by an FDA-approved test, and involvement of lymph node(s), following complete resection. BRAF V600EMutation-Positive Metastatic NSCLC- in combination with dabrafenib, for the treatment of patients with metastatic non-small cell lung cancer (NSCLC) with BRAF V600E mutation as detected by an FDA-approved test. BRAF V600EMutation-Positive Locally Advanced or Metastatic Anaplastic Thyroid Cancer - in combination with dabrafenib, for the treatment of patients with locally advanced or metastatic anaplastic thyroid cancer. 		
Mektovi (binimetinib)	Indicated, in combination with encorafenib, for the treatment of patients with unresectable or metastatic melanoma with a BRAF V600E or V600K mutation, as detected by an FDA-approved test.	Rx by Oncologist	MEKTOVI PI
methadone (for pain) SEE SPECIAL NOTE REGARDING PA REQUIREMENTS AND FORM **********	Indicated for management of pain severe enough to require daily, around-the-clock, long-term opioid treatment and for which alternative treatment options are inadequate.	All long-acting opioids require Prior Authorization (PA). The PA form can be accessed using the following link: Opioid PA Form	METHADONE PI
Minastrin 24 Fe (norethindrone, ethinyl estradiol and ferrous fumarate)	See Oral Contraceptive ***CHEWABLE***	ANY OCP on prior authorization requires documentation demonstrating a compelling reason why formulary OCPs cannot be used [ex: intolerance, prior side effects, failures, etc. documented after a 3-month trial of formulary OCPs]	MINASTRIN 24 Fe PI
Movantik (naloxegol)	Indicated for the treatment of opioid-induced constipation in adults with chronic non-cancer pain, including patients with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation.	Failure of at least 2 of the following: docusate, sennosides, polyethylene glycol 3350, lactulose, methylcellulose	MOVANTIK PI
MS Contin (morphine sulfate extended release) SEE SPECIAL NOTE REGARDING PA	Indicated for the management of moderate to severe pain when a continuous, around-the-clock opioid analgesic is needed for an extended period of time.	All long-acting opioids require Prior Authorization (PA). The PA form can be accessed using the following link: Opioid PA Form	MS CONTIN PI

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Nucala (mepolizumab)	 Indicated for: The add-on maintenance treatment of patients with severe asthma aged 6 years and older, and with an eosinophilic phenotype. The treatment of adult patients with eosinophilic granulomatosis with polyangiitis (EGPA). 	Rx by Allergist or Pulmonologist	NUCALA PI
Nulibry (fosdenopterin)	Indicated to reduce the risk of mortality in patients with molybdenum cofactor deficiency (MoCD) Type A.	***MFC-DC Pharmacist Review***	NULIBRY PI
Nuvigil (armodafinil)	Indicated to improve wakefulness in adult patients with excessive sleepiness associated with obstructive sleep apnea, narcolepsy, or shift work disorder.		NUVIGIL PI
Nuedexta (dextromethorphan hydrobromide and quinidine sulfate)	Indicated for the treatment of pseudobulbar affect.	Rx by Neurologist	NUEDEXTA PI
Ofev (nintedanib)	Indicated for: 1. The treatment of idiopathic pulmonary fibrosis. 2. To slow the rate of decline in pulmonary function in patients with systemic sclerosis associated interstitial lung disease (SSc-ILD).	Rx by Pulmonologist	OFEV PI
OmniPod-Insulin Management (EIM) Systems	Indicated for subcutaneous delivery of insulin at set and variable rates for the management of diabetes mellitus in persons requiring insulin and for the quantitative measurement of glucose in fresh whole capillary blood (in vitro) from the finger.	RX Endocrinology Must meet criteria found in Policy 1413.DC Please click link below for EIM Policy: MFC External Insulin Pumps Policy	OMNIPOD PI
Onpattro (patisiran)	Indicated for the treatment of the polyneuropathy of hereditary transthyretin-mediated amyloidosis in adults.	Rx by Rheumatology or Neurology	ONPATTRO PI
Onureg (azacitidine)	Indicated for continued treatment of adult patients with acute myeloid leukemia who achieved first complete remission or complete remission with incomplete blood count recovery following intensive induction chemotherapy and are not able to complete intensive curative therapy.	Rx by Oncologist	ONUREG PI
OPIOIDS PRIOR AUTHORIZATION	IMPORTANT INFORMATION ABOUT PRESCRIBING OPIOIDS FOR MEDSTAR FAMILY CHOICE MEMBERS	The PA form can be accessed using the following link:	
TERMS	"Early" Opioid Refills Will No Longer be Covered by MedStar Family Choice - Effective 1/1/2019 Beginning 1/1/2019, MedStar Family Choice will not authorize early refills of controlled medications. Specifically, MedStar Family Choice will not approve early refills, override Managed Drug Limitations (MDL), replace lost/stolen medications, or provide early	Opioid PA Form	

SEE SPECIAL NOTE
REGARDING PA
REQUIREMENTS AND
FORM

refills for travel for controlled medications. Exceptions may be granted if a member is receiving controlled medication(s) for cancer treatment, sickle cell disease, or is in hospice/receiving palliative care.

PRIOR AUTHORIZATION

Prior Authorization will be required for:

- Prescriptions > 50 MME/day or more than 7 day for an opioid naïve patient (no opioids taken in the previous 90 days or one ≤ 50 MME per day, ≤ 7 day prescription taken in the previous 90 days) as described in Section I below.
- opioid experienced patients as described in Section II below.

SECTION I. OPIOID NAÏVE PATIENTS (defined as no opioids in the previous 90 days or one fill of ≤ 50 MME per day for ≤ 7 days prescription taken in the previous 90 days)

A "new" prescription means that a patient has not had an opioid medication filled under MedStar Family Choice in the preceding 90 days or had one short-acting opioid at ≤ 50 morphine equivalents per day for 7 or fewer days in previous 90 days. New prescriptions for more than 7-days' supply or greater than 50 MME per day will require Prior Authorization. It is our hope that limiting opioid quantities to a 7-day supply will discourage abuse, both by our patients and by the community at large. This change is also consistent with Medicare policy (effective 2019) which limits opioid naïve patients to a 7-day supply.

According to the CDC 2016 Guidelines for Prescribing Opioids, "When opioids are used for acute pain, clinicians should prescribe the lowest effective dose of immediate-release opioids and should prescribe no greater quantity than needed for the expected duration of pain severe enough to require opioids. Three days or less will often be sufficient; more than seven days will rarely be needed."

Examples of a typical 3-day supply and a 7-day supply of frequently prescribed opioids are below:

	3-day supply	7-day supply quantity* (maximum
Medication	quantity*	allowable)
HYDROMORPHONE TAB 2MG	18 tablets	42 tablets
MORPHINE SULFATE TAB 15MG	18 tablets	42 tablets
OXYCODONE SOLUTION 5MG/5ML	180 mL	420 mL
OXYCODONE TAB 5MG	18 tablets	42 tablets
TRAMADOL HCL TAB 50MG	18 tablets	42 tablets

^{*}Quantities are based on starting dose recommendations in the respective FDA Package Inserts for each medication.

Please contact MedStar Family Choice at 888-798-4244 for Prior Authorization of new opioid prescriptions that exceed the limits. Should you have any questions or concerns about this new policy, please call Dr. Kazmi at 202-469-6727.

MedStar Family Choice strongly encourages you to prescribe the least amount of opioid at the lowest dose possible to achieve pain relief goals.

SECTION II. OPIOID EXPERIENCED PATIENTS

Prior Authorization is required for the following medications:

- Long-acting opioids
- Fentanyl products
- Methadone for pain
- Any opioid prescription (or combination of opioid prescriptions) that results in a
 patient exceeding 90 morphine milliequivalents (MME) per day. Instructions on
 calculating MME are available at the <u>CDC website</u>.

For the sake of illustration of what constitutes 90 MME, the following is a list of daily doses of commonly prescribed opioids that **equal 90 MME/day**:

Fentanyl 112.5 mcg/day Hydrocodone 90 mg/day Hydromorphone 22.5 mg/day Morphine 90 mg/day Oxycodone 60 mg/day Oxymorphone 30 mg/day

The following are examples of common prescriptions that **equal 90 MME/day**: oxycodone 20 mg tid methadone 20 mg qd hydrocodone 10/325, 3 tabs tid

Additionally, some smaller doses of immediate release medications will require prior authorization at **less than 90 MME**. The decision to limit these medications was made in an effort to decrease the number of pills available for diversion. These medications are as follows:

Medication	Max per 30 days	Unit
Codeine compounds (all)	1,000	mL
	180	tab/ cap

Hydrocodone compounds (all)	2,750	mL
	180	tab/ cap
Hydromorphone (1 mg/mL solution, 2 mg tablet, 3 mg	675	mL
suppository)	180	tab/ supp
Morphine (5 mg suppository, 10 mg/5mL solution, 10 mg	1,350	mL
suppository)	180	supp
Oxycodone compounds (2.5 mg, 5 mg, 7.5 mg of all formulations)	1,800	mL
	180	tab/ cap
Tramadol (100 mg, 200 mg)	180	tab/ cap

In order to receive prior authorization, prescribers **must** attest to the following:

- Prescriber has reviewed controlled substance prescriptions in a Prescription Drug Monitoring Program.
- Prescriber will utilize random Urine Drug Screens.
- Prescriber has provided or offered a prescription for naloxone to the patient or patient's household if the patient has:
 - o a history of substance use disorder
 - requires more than 50 MME (for example, more than Fentanyl 62.5 mcg/72 hours, hydrocodone 50 mg/day, hydromorphone 12.5 mg/day, morphine 50 mg/day, oxycodone 33 mg/ day, and oxymorphone 16 mg/day)
 - o is prescribed both opioids and benzodiazepines
 - o is prescribed other sedative hypnotics
 - o or for any other reason deemed clinically appropriate
- Prescriber and patient have signed a Pain Management/Opioid Treatment Agreement/Contract and it is stored in the patient's medical record.

Oral Contraceptives	While some oral contraceptives have additional indications (ex: Beyaz for acne, PMDD, folate replacement; Estrostep Fe for acne; Safyral for folate replacement; Natazia for heavy periods), most are simply indicated for the prevention of pregnancy.	ANY OCP on prior authorization requires documentation demonstrating a compelling reason why formulary OCPs cannot be used [ex: intolerance, prior side effects, failures, etc. documented after a 3-month trial of formulary OCPs]	
Orfadin (nitisinone)	Indicated for the treatment of adult and pediatric patients with hereditary tyrosinemia type 1 in combination with dietary restriction of tyrosine and phenylalanine. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.	***MFC-DC Pharmacist Review***	ORFADIN PI
Orkambi (lumacaftor/ivacaftor)	Indicated for the treatment of cystic fibrosis (CF) in patients age 2 years and older who are homozygous for the F508del mutation in the CFTR gene. If the patient's genotype is unknown, an FDA-cleared CF mutation test should be used to detect the presence of the F508del mutation on both alleles of the CFTR gene.	Rx by Pulmonologist	ORKAMBI PI
Oriahnn (elagolix, estradiol, and norethindrone acetate capsules)	Indicated for the management of heavy menstrual bleeding associated with uterine leiomyomas (fibroids) in premenopausal women.	Must have tried and failed at least two of the following: NSAIDs, hormonal options (OCP, progesterone, hormonal IUD), or have a contraindication to using these therapies.	ORIAHNN PI
Orilissa (elagolix)	Indicated for the management of moderate to severe pain associated with endometriosis.	Must have tried and failed at least two of the following: NSAIDs, hormonal options (OCP, progesterone, hormonal IUD), or have a contraindication to using these therapies.	ORILISSA PI
Orladeyo (berotralstat)	Indicated for prophylaxis to prevent attacks of hereditary angioedema (HAE) in adults and pediatric patients 12 years and older. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.	***MFC-DC Pharmacist Review***	ORLADEYO PI
Oxlumo (lumasiran)	Indicated for the treatment of primary hyperoxaluria type 1 (PH1) to lower urinary oxalate levels in pediatric and adult patients. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.	***MFC-DC Pharmacist Review***	OXLUMO PI
Oxymorphone ER SEE SPECIAL NOTE REGARDING PA REQUIREMENTS AND FORM **********	Indicated for the relief of moderate to severe pain in patients requiring continuous around-the-clock opioid treatment for an extended period of time.	All long-acting opioids require Prior Authorization (PA). The PA form can be accessed using the following link: Opioid PA Form	OPANA ER PI

Pemazyre (pemigatinib)	Indicated for the treatment of adults with previously treated, unresectable locally advanced or metastatic cholangiocarcinoma with a fibroblast growth factor receptor 2 (FGFR2) fusion or other rearrangement as detected by an FDA-approved test.	Rx by Oncologist	PEMAZYRE PI
Pretomanid	Indicated, as part of a combination regimen with bedaquiline and linezolid for the treatment of adults with pulmonary extensively drug resistant (XDR), treatment-intolerant or nonresponsive multidrug-resistant (MDR) tuberculosis (TB).	Rx by Pulmonologist	<u>PRETOMANID</u>
	 Pretomanid Tablets are not indicated for patients with: Drug-sensitive (DS) tuberculosis. Latent infection due to Mycobacterium tuberculosis Extra-pulmonary infection due to Mycobacterium tuberculosis. MDR-TB that is not treatment-intolerant or nonresponsive to standard therapy Safety and effectiveness of Pretomanid Tablets have not been established for its use in combination with drugs other than bedaquiline and linezolid as part of the recommended dosing regimen. 		
Piqray (alpelisib)	Indicated in combination with fulvestrant for the treatment of postmenopausal women, and men, with hormone receptor (HR)- positive, human epidermal growth factor receptor 2 (HER2)-negative, PIK3CA-mutated, advanced or metastatic breast cancer as detected by an FDA-approved test following progression on or after an endocrine-based regimen.	Rx by Oncologist	PIQRAY PI
Polivy (polatuzumab vedotin)	Indicated in combination with bendamustine and a rituximab product for the treatment of adult patients with relapsed or refractory diffuse large B-cell lymphoma, not otherwise specified, after at least two prior therapies.	Rx by Oncologist	POLIVY PI
Prolia (denosumab)	 Indicated for: treatment of postmenopausal women with osteoporosis at high risk for fracture. treatment to increase bone mass in men with osteoporosis at high risk for fracture. treatment of glucocorticoid-induced osteoporosis in men and women at high risk for fracture. treatment to increase bone mass in men at high risk for fracture receiving androgen deprivation therapy for non-metastatic prostate cancer. treatment to increase bone mass in women at high risk for fracture receiving adjuvant aromatase inhibitor therapy for breast cancer. 		PROLIA PI
Provigil (modafinil)	Indicated to improve wakefulness in adult patients with excessive sleepiness associated with narcolepsy, obstructive sleep apnea, or shift work disorder.		PROVIGIL PI
Pulmozyme (dornase alfa) Inhalation solution	Indicated in conjunction with standard therapies for the management of cystic fibrosis (CF) patients to improve pulmonary function.	Rx by Pulmonologist	PULMOZYME PI
Pyrukynd (mitapivat)	Indicated for the treatment of hemolytic anemia in adults with pyruvate kinase (PK) deficiency. PA SUBMISSION REQUIREMENTS 1. Laboratory evidence of reduced PK enzymatic activity in red blood cell 2. Confirmatory genetic testing of PKLR gene showing 2 mutant alleles with at least onemissense mutation 3. Concomitant use with folic acid	Rx by or in consultation with a Hematologist	<u>PYRUKYND</u>

Qbrexza (glycopyrronium)	Indicated for topical treatment of primary axillary hyperhidrosis in adults and pediatric patients 9 years of age and older	1. Must have tried and failed OTC Clinical Strength antiperspirants and at least one prescription strength antiperspirant (ex: Drysol). 2. Documentation that symptoms are persistent despite previous treatment attempts and that the degree of symptomatology impacts quality of life must be clearly indicated in a recent (<6 month old) clinical encounter note.	QBREXZA PI
Qelbree (viloxazine extended- release capsules)	Indicated for the treatment of Attention Deficit Hyperactivity Disorder (ADHD) in pediatric patients 6 to 17 years of age.	clinical encounter note.	QELBREE PI
Qulipta (atogepant)	Indicated for the preventive treatment of episodic migraine in adults	1.Trial and failure or intolerance to at least three of the following agents: beta blockers, topiramate, Aimovig, and Ubrelvy, in medical documentation submitted. 2. Patient must have at least 4 headache days per month on average.	QULIPTA PI
Rasuvo (methotrexate inj)	 Indicated for: Management of patients with severe, active rheumatoid arthritis (RA) and polyarticular juvenile idiopathic arthritis (pJIA), who are intolerant of or had an inadequate response to first-line therapy. Symptomatic control of severe, recalcitrant, disabling psoriasis in adults who are not adequately responsive to other forms of therapy. 	Rx by Rheumatology or Dermatology	RASUVO PI
Ravicti (glycerol phenylbutyrate)	Indicated for chronic management of patients with urea cycle disorders (UCDs) who cannot be managed by dietary protein restriction and/or amino acid supplementation alone. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.	***MFC-DC Pharmacist Review***	RAVICTI PI
Repatha (evolocumab)	 Indicated: to reduce the risk of myocardial infarction, stroke, and coronary revascularization in adults with established cardiovascular disease. as an adjunct to diet, alone or in combination with other lipid-lowering therapies (e.g., statins, ezetimibe), for treatment of adults with primary hyperlipidemia (including heterozygous familial hypercholesterolemia) to reduce low-density lipoprotein cholesterol. 	Rx by Cardiologist or by Lipid Specialist.	REPATHA PI

	3. as an adjunct to diet and other LDL-lowering therapies (e.g., statins, ezetimibe, LDL apheresis) in patients with homozygous familial hypercholesterolemia (HoFH) who require additional lowering of LDL-C.		
Restasis (cyclosporine ophthalmic emulsion)	Indicated to increase tear production in patients whose tear production is presumed to be suppressed due to ocular inflammation associated with keratoconjunctivitis sicca.	Must have tried and failed artificial tears.	RESTASIS PI
Retevmo (selpercatinib)	 Indicated for: Adult patients with metastatic RET (rearranged during transfection) fusion-positive non-small cell lung cancer (NSCLC). Adult and pediatric patients 12 years of age and older with advanced or metastatic RET-mutant medullary thyroid cancer (MTC) who require systemic therapy. Adult and pediatric patients 12 years of age and older with advanced or metastatic RET fusion-positive thyroid cancer who require systemic therapy and who are radioactive iodine-refractory (if radioactive iodine is appropriate. 	Rx by Oncologist	RETEVMO PI
Revatio (sildenafil)	Indicated for the treatment of pulmonary arterial hypertension (PAH) (WHO Group I) in adults to improve exercise ability and delay clinical worsening. Studies establishing effectiveness were short-term (12 to 16 weeks) and included predominately patients with NYHA Functional Class II–III symptoms. Etiologies were idiopathic (71%) or associated with connective tissue disease (25%).	Rx by Pulmonologist or Cardiologist	REVATIO PI
Revcovi (elapegademase-lvlr)	Indicated for the treatment of adenosine deaminase severe combined immune deficiency (ADA-SCID) in pediatric and adult patients. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial	***MFC-DC Pharmacist Review***	REVCOVI PI
Revlimid (lenalidomide)	Indicated for the treatment of adult patients with: 1. Multiple myeloma (MM), in combination with dexamethasone. 2. MM, as maintenance following autologous hematopoietic stem cell transplantation (auto-HSCT). 3. Transfusion-dependent anemia due to low- or intermediate-1-risk myelodysplastic syndromes. 4. (MDS) associated with a deletion 5q abnormality with or without additional cytogenetic abnormalities. 5. Mantle cell lymphoma (MCL) whose disease has relapsed or progressed after two prior therapies, one of which included bortezomib. 6. Previously treated follicular lymphoma (FL), in combination with a rituximab product. 7. Previously treated marginal zone lymphoma (MZL), in combination with a rituximab product		REVLIMID PI
Reyvow (lasmiditan)	Indicated for the acute treatment of migraine with or without aura in adults.	Member must have tried and failed NSAIDs and Triptans or have a contraindication to taking either of these medications.	REYVOW PI

Rezurock (belumosudil)	Indicated for the treatment of adult and pediatric patients 12 years and older with chronic graft-versus-host disease (chronic GVHD) after failure of at least two prior lines of systemic therapy.	Member must have tried and failed, have intolerance or medical contraindication to at least three of these medications: cyclosporine, methotrexate, mycophenolate, sirolimus, and glucocorticoids.	REZUROCK PI
Rituxan	Indicated for:		<u>RITUXAN PI</u>
(rituximab)	 Pediatric patients aged 6 months and older with mature B-cell NHL and mature B-cell acute leukemia (B-AL) a. Previously untreated, advanced stage, CD20-positive, diffuse large B-cell lymphoma (DLBCL), Burkitt lymphoma (BL), Burkitt-like lymphoma (BLL) or mature B-cell acute leukemia (B-AL) in combination with chemotherapy. Moderate to severe Pemphigus Vulgaris (PV) in adult patients Demonstrated failure or intolerance to Truxima for the following indications: Adult patients with Non-Hodgkin's Lymphoma (NHL) Relapsed or refractory, low grade or follicular, CD20-positive B-cell NHL as a single agent. Previously untreated follicular, CD20-positive, B-cell NHL in combination with first line chemotherapy and, in patients achieving a complete or partial response to a rituximab product in combination with chemotherapy, as single-agent maintenance therapy. Non-progressing (including stable disease), low-grade, CD20-positive, Bcell NHL as a single agent after first-line cyclophosphamide, vincristine, and prednisone (CVP) chemotherapy. Previously untreated diffuse large B-cell, CD20-positive NHL in combination with (cyclophosphamide, doxorubicin, vincristine, and prednisone) (CHOP) or other anthracycline-based chemotherapy regimens. Adult patients with Chronic Lymphocytic Leukemia (CLL)		
	considered and PA Criteria will not apply.		
Rituxan Hycela	Indicated for:	Rx by Oncologist	RITUXAN HYCELA PI
(rituximab and	1. Follicular Lymphoma (FL)		
hyaluronidase human)	Relapsed or refractory, follicular lymphoma as a single agent.		<u> </u>

	 Previously untreated follicular lymphoma in combination with first line chemotherapy and, in patients achieving a complete or partial response to rituximab in combination with chemotherapy, as single agent maintenance therapy. Non-progressing (including stable disease), follicular lymphoma as a single agent after first-line cyclophosphamide, vincristine, and prednisone (CVP) chemotherapy. Diffuse Large B-cell Lymphoma (DLBCL) previously untreated diffuse large B-cell lymphoma in combination with cyclophosphamide, doxorubicin, vincristine, prednisone (CHOP) or other anthracycline-based chemotherapy regimens. Chronic Lymphocytic Leukemia (CLL) previously untreated and previously treated CLL in combination with fludarabine and cyclophosphamide (FC). 		
Rozlytrek	Indicated for the treatment of:	Rx by Oncologist	ROZLYTREK
(entrectinib)	 Adult patients with metastatic non-small cell lung cancer (NSCLC) whose tumors are ROS1-positive. Adult and pediatric patients 12 years of age and older with solid tumors that have a neurotrophic tyrosine receptor kinase (NTRK) gene fusion without a known acquired resistance mutation, are metastatic or where surgical resection is likely to result in severe morbidity and have either progressed following treatment or have no satisfactory alternative therapy. This indication is approved under accelerated approval based on tumor response rate and durability of response. 		
Rubraca	Indicated for:	Rx by Oncologist	RUBRACA PI
(rucaparib)	 maintenance treatment of adult patients with recurrent epithelial ovarian, fallopian tube, or primary peritoneal cancer who are in a complete or partial response to platinum-based chemotherapy. treatment of adult patients with deleterious BRCA mutation (germline and/or somatic)-associated epithelial ovarian, fallopian tube, or primary peritoneal cancer who have been treated with two or more chemotherapies. 		
Rybrevant (amivantamab-vmjw)	Indicated for the treatment of adult patients with locally advanced or metastatic non-small cell lung cancer (NSCLC) with epidermal growth factor receptor (EGFR) exon 20 insertion mutations, as detected by an FDA-approved test, whose disease has progressed on or after platinum-based chemotherapy.	Rx by Oncologist	RYBREVANT PI
Sabril (vigabatrin)	 Indicated for: the treatment of Refractory Complex Partial Seizures as adjunctive therapy in patients 2 years of age and older who have responded inadequately to several alternative treatments; SABRIL is not indicated as a first line agent. Infantile Spasms - monotherapy in infants 1 month to 2 years of age for whom the potential benefits outweigh the potential risk of vision loss. 	Rx by Neurologist	SABRIL PI
Santyl Ointment Collagenase	Indicated for debriding chronic dermal ulcers and severely burned areas.	Rx by Dermatologist or Wound Care Specialist	SANTYL PI
Saphnelo	Indicated for the treatment of adult patients with moderate to severe systemic lupus		SAPHNELO PI
(anifrolumab-fnia)	erythematosus (SLE), who are receiving standard therapy.		
	PA SUBMISSION REQUIREMENTS:		
	Confirmed diagnosis of SLE		

		1	
	2. Lab report showing autoantibodies (e.g., ANA, anti-ds, anti-Sm)		
	3. Current therapy for SLE alone or in combination with:		
	a. Glucocorticoid (e.g. prednisone, methylprednisone, dexamethasone)		
	b. Antimalarials (e.g. hydroxychloroquine)		
	c. Immunosuppressants (e.g. azathioprine, methotrexate, mycophenolate,		
	cyclosporine, cyclophosphamide)		
	4. Excluded use with Benlysta		
	5. Excluded to use with active lupus nephritis or central nervous system lupus		
	6. Trial and failure or intolerance to Benlysta		
Scemblix	A kinase inhibitor indicated for the treatment of adult patients with:	Rx by Oncologist or	SCEMBLIX PI
(asciminib)	1. Philadelphia chromosome-positive chronic myeloid leukemia (Ph+ CML) in chronic	Hematologist	
	phase (CP), previously treated with two or more tyrosine kinase inhibitors (TKIs)This		
	indication is approved under accelerated approval based on major molecular		
	response (MMR). Continued approval for this indication may be contingent upon		
	verification and description of clinical benefit in a confirmatory trial(s).		
	2. Ph+ CML in CP with the T315I mutation.		
Serostim	Indicated for the treatment of HIV patients with wasting or cachexia to increase lean body	Rx by ID or HIV Specialist	SEROSTIM PI
(somatropin (rDNA	mass and body weight and improve physical endurance.		
origin))			
Seysara	Indicated for the treatment of inflammatory lesions of non-nodular moderate to severe acne	Rx by Dermatologist.	SEYSARA PI
(seracycline)	vulgaris in patients 9 years of age and older.		
		Failure of at least one other oral	
		tetracycline antibiotic.	
Signifor LAR	Indicated for the treatment of patients with acromegaly who have had an inadequate	Rx by Endocrinologist	SIGNIFOR LAR PI
(pasireotide)	response to surgery and/or for whom surgery is not an option.		
Sirturo	Indicated as part of combination therapy in adult and pediatric patients (12 to less than	Rx by ID	SITURO PI
(bedaquiline)	18 years of age and weighing at least 30 kg) with pulmonary multi-drug resistant tuberculosis		
	(MDR-TB). [Reserved for use when an effective treatment regimen cannot otherwise be		
	provided; not indicated for the treatment of latent, extra pulmonary or drug-sensitive		
	tuberculosis; should be administered by directly observed therapy. Safety and efficacy of		
	SIRTURO in HIV-infected patients with MDR-TB have not been established, as clinical data are		
	limited.		SOLUDIO DI
Soliris	Indicated for:	***MFC-DC	SOLIRIS PI
(eculizumab)	1. treatment of patients with paroxysmal nocturnal hemoglobinuria (PNH) to reduce		
	hemolysis.	Pharmacist	
	2. treatment of patients with atypical hemolytic uremic syndrome (aHUS) to inhibit	Review***	
	complement-mediated thrombotic microangiopathy. The effectiveness of Soliris in aHUS is		
	based on the effects on thrombotic microangiopathy (TMA) and renal function.		
	Continued approval for this indication may be contingent upon verification and description of		
	clinical benefit in a confirmatory trial		

Spinraza	Indicated for the treatment of spinal muscular atrophy (SMA) in pediatric and adult patients.	***MFC-DC	SPINRAZA PI
(nusinersen)	Continued approval for this indication may be contingent upon verification and description of	Pharmacist	
	clinical benefit in a confirmatory trial.		
	Clinical benefit in a committatory trial.	Review***	
Stimate nasal spray	See Desmopressin Products		
(desmopressin)			
Stivarga	Indicated for:	Rx by Oncologist	STIVARGA PI
(regorafenib)	1. treatment of metastatic colorectal cancer previously treated with ALL the following		
	therapies:		
	a. fluoropyrimidine-based chemotherapy		
	b. oxaliplatin-based chemotherapy		
	c. irinotecan-based chemotherapy		
	d. an anti-vascular endothelial growth factor (VEGF) therapy		
	e. if Kirsten RNA Associated Rat Sarcoma 2 Virus Gene (KRAS) wild type, an anti-		
	epidermal growth factor receptor (EGFR) therapy		
	2. treatment of locally advanced, unresectable or metastatic gastrointestinal stromal tumor		
	(GIST), previously treated with imatinib mesylate and sunitinib malate.		
Chuamaatal	hepatocellular carcinoma (HCC) who have been previously treated with sorafenib.	At this time a subsection to the	CTDOMACCTOL DI
Stromectol	Indicated for the treatment of the following infections:	At this time, outpatient use for	STROMECTOL PI
(ivermectin)	1. Strongulaidiasis of the intestinal treet (i.e., nandiscominated) strongulaidiasis due to the	COVID-19 treatment is	
	1. Strongyloidiasis of the intestinal tract (i.e., nondisseminated) strongyloidiasis due to the nematode parasite <i>Strongyloides stercoralis</i> .	prohibited.	
	2. Onchocerciasis due to the nematode parasite <i>Onchocerca volvulus</i> .		
	2. Offchocerciasis due to the hematode parasite offchocercu volvulus.		
Synagis	Indicated for prevention of serious lower respiratory tract disease caused by RSV in pediatric	Please submit:	SYNAGIS PI
(palivizumab)	Patients:	A COMPLETED PRIOR	
SEE SPECIAL NOTE		AUTHORIZATION FORM (see link	
REGARDING PA	1. with a history of premature birth (≤35 weeks gestational age) and who are 6 months	below)	
REQUIREMENTS AND	of age or younger at the beginning of RSV season		
FORM	2. with bronchopulmonary dysplasia (BPD) that required medical treatment within the	SYNAGIS PRIOR AUTHORIZATION	
*******	previous 6 months and who are 24 months of age or younger at the beginning of RSV	AND PRESCRIPTION FORM	
	season		
	3. with hemodynamically significant congenital heart disease (CHD) and who are 24	To view the most up to date AAP	
	months of age or younger at the beginning of RSV season.	Synagis Guidelines, follow the	
		link below:	
	MedStar Family Choice uses the newest recommendations of the American Academy of	AAP SYNAGIS GUIDELINES	
	Pediatrics (AAP).		
	Recommendations were last updated in the journal Pediatrics (7/28/2014 issue):		
	Updated Guidance for Palivizumab Prophylaxis Among Infants and Young Children at		
	Increased Risk of Hospitalization for Respiratory Syncytial Virus Infection.		
Synribo	Indicated to treat adults with chronic phase (CP) or accelerated phase (AP) CML with	Rx by Oncologist	SYNRIBO PI
(omacetaxine)	resistance and/or intolerance to two or more TKIs.		

Syprine (trientine hydrochloride)	Indicated in the treatment of patients with Wilson's disease who are intolerant of penicillamine.		SYPRINE PI
Tabrecta (capmatinib)	Indicated for treatment of adults with metastatic NSCLC whose tumors have a mutation that leads to mesenchymal-epithelial transition (MET) exon 14 skipping as detected by an approved test.		TABRECTA PI
Tafinlar (dabrafenib)	Indicated for the treatment of patients with unresectable or metastatic melanoma with BRAF V600E mutation as detected by an FDA-approved test. TAFINLAR in combination with trametinib is indicated for the treatment of patients with unresectable or metastatic melanoma with BRAF V600E or V600K mutations as detected by an FDA-approved test. The use in combination is based on the demonstration of durable response rate. Improvement in disease-related symptoms or overall survival has not been demonstrated for TAFINLAR in combination with trametinib	Rx by Oncologist	TAFINLAR PI
Tagrisso (osimertinib)	 Indicated: for the treatment of first-line treatment of patients with metastatic NSCLC whose tumors have epidermal growth factor receptor (EGFR) Exon 19 deletions or exon 21 L858R mutations, as detected by an FDA approved test. For the treatment of patients with metastatic EGFR T790M mutation-positive NSCLC, as detected by an FDA approved test, whose disease has progressed on or after EGFR TKI therapy. 	Rx by Oncologist	TAGRISSO PI
Talzenna (talazoparib)	Indicated for the treatment of adult patients with deleterious or suspected deleterious germline BRCA-mutated (<i>gBRCAm</i>) HER2-negative locally advanced or metastatic breast cancer.	Rx by Oncologist	TALZENNA PI
Tarceva (erlotinib)	 Indicated for: treatment of patients with metastatic non-small cell lung cancer (NSCLC) whose tumors have epidermal growth factor receptor (EGFR) exon 19 deletions or exon 21 (L858R) substitution mutations as detected by an FDA-approved test receiving first-line, maintenance, or second or greater line treatment after progression following at least one prior chemotherapy regimen. first-line treatment of patients with locally advanced, unresectable or metastatic pancreatic cancer, in combination with gemcitabine. 	Rx by Oncologist	TARCEVA PI
Tarpeyo (budesonide delayed- release capsules)	 Indicated to reduce proteinuria in adults with primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression, generally a urine protein-to-creatinine ratio (UPCR) ≥ 1.5 g/g. PA SUBMISSION REQUIREMENTS Confirmed diagnosis of primary immunoglobulin A nephropathy History of failure, contraindication or intolerance to a glucocorticoid Patient does not have severe hepatic impairment (Child-Pugh Class C) Estimated glomerular filtration rate (eGFR) ≥ 35 mL/min/1.73 m² Proteinuria ≥ 1g/day Patient is on a stable and maximally tolerated dose of a renin-angiotensin system (RAS) inhibitor (angiotensin converting enzyme [ACE] inhibitor or angiotensin receptor blocker [ARB]), for at least 3 months, unless contraindicated 	Rx by Nephrologist or Immunologist	TARPEYO PI

Tasigna	Indicated for:	Rx by Oncologist	TASIGNA PI
(nilotinib)	 adult and pediatric patients greater than or equal to 1 year of age with newly diagnosed Philadelphia chromosome positive chronic myeloid leukemia (Ph+ CML) in chronic phase. adult patients with chronic phase (CP) and accelerated phase (AP) Ph+ CML resistant to or intolerant to prior therapy that included imatinib. pediatric patients greater than or equal to 1 year of age with Ph+ CML-CP resistant or intolerant to prior tyrosine-kinase inhibitor (TKI) therapy. 	nx by Officologist	TAJUNATI
Tavalisse	Indicated for the treatment of thrombocytopenia in adult patients with chronic immune	Rx by Hematologist	TAVALISSE PI
(fostamatinib disodium hexahydrate)	thrombocytopenia (ITP) who have had an insufficient response to a previous treatment.		
Tavneos	Indicated for:	Rx by Rheumatologist	TAVNEOS PI
(avacopan)	 adjunctive treatment of adult patients with severe active anti-neutrophil cytoplasmic autoantibody (ANCA)-associated vasculitis (granulomatosis with polyangiitis [GPA] and microscopic polyangiitis [MPA]) in combination with standard therapy including glucocorticoids. TAVNEOS does not eliminate glucocorticoid use. PA SUBMISSION REQUIREMENTS Patient must be ≥ 18 years of age. Patient must have a diagnosis of active anti-neutrophil cytoplasmic autoantibody (ANCA)-associated vasculitis. Laboratory tests before initiating Tavneos (avacopan) therapy: Liver test panel Screening the patient for hepatitis B infection by measuring HBsAg and anti-HBc Documentation of baseline Birmingham vasculitis activity score (BVAS), with either one of the following: At least one major item At least 2 renal items, proteinuria and hematuria are present Documentation that patient will continue standard therapy including glucocorticoids. 		
Taytulla	See Oral Contraceptive	ANY OCP on prior authorization	TAYTULLA PI
(norethindrone/ethinyl estradiol capsules and ferrous fumarate)		requires documentation demonstrating a compelling reason why formulary OCPs cannot be used [ex: intolerance, prior side effects, failures, etc. documented after a 3-month trial of formulary OCPs]	
Tazverik	Indicated for the treatment of adults and pediatric patients aged 16 years and older with	Rx by Oncologist	TAZVERIK PI
(tazemetostat)	metastatic or locally advanced epithelioid sarcoma not eligible for complete resection.		
Tibsovo (ivosidenib)	 Indicated for the treatment of adult patients with relapsed or refractory acute myeloid leukemia (AML) with a susceptible IDH1 mutation as detected by an FDA-approved test in: Adult patients with newly -diagnosed AML who are ≥ 75 years old or who have comorbidities that preclude use of intensive induction chemotherapy. Adult patients with relapsed or refractory AML. 	Rx by Oncologist	TIBSOVO PI

Trikafta (elexacaftor, ivacaftor, and tezacaftor)	Indicated for the treatment of cystic fibrosis (CF) in patients aged 12 years and older who have at least one F508del mutation in the CFTR gene. If the patient's genotype is unknown, an FDA-cleared CF mutation test should be used to confirm the presence of at least one F508del mutation.	Rx by Pulmonologist	TRIKAFTA PI
Trodelvy (sacituzumab govitecan-hziy)	Indicated for the treatment of adult patients with metastatic triple-negative breast cancer (mTNBC) who have received at least two prior therapies for metastatic disease.	Rx by Oncologist	TRODELVY PI
Truseltiq (infigratinib)	Indicated for the treatment of previously treated, unresectable locally advanced or metastatic cholangiocarcinoma (CCA) with a fibroblast growth factor receptor II (FGFR2) fusion in adult patients.	Rx by Oncologist	TRUSELTIQ PI
Tukysa (tucatinib)	Indicated in combination with trastuzumab and capecitabine for treatment of adult patients with advanced unresectable or metastatic HER2-positive breast cancer, including patients with brain metastases, who have received one or more prior anti-HER2-based regimens in the metastatic setting.	Rx by Oncologist	TUKYSA PI
Turalio (pexidartinib)	Indicated for the treatment of adult patients with symptomatic tenosynovial giant cell tumor (TGCT) associated with severe morbidity or functional limitations and not amenable to improvement with surgery.	Rx by Oncologist	TURALIO PI
Tykerb (lapatinib)	Indicated in combination with capecitabine for the treatment of patients with advanced or metastatic breast cancer whose tumors overexpress human epidermal growth factor receptor 2 (HER2) and who have received prior therapy including an anthracycline, a taxane, and trastuzumab. Patients should have disease progression on trastuzumab prior to initiation of treatment with	Rx by Oncologist	TYKERB PI
	TYKERB in combination with capecitabine		
Ubrelvy (ubrogepant)	Indicated for the acute treatment of migraine with or without aura in adults.	Member must have tried and failed NSAIDs and Triptans or have a contraindication to taking either of these medications.	UBRELVY PI
Uplizna (inebilizumab-cdon)	Indicated for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult patients who are anti-aquaporin-4 (AQP4) antibody positive		UPLIZNA PI
Vabysmo (faricimab-svoa)	A vascular endothelial growth factor (VEGF) and angiopoietin-2 (Ang-2) inhibitor for intravitreal indicated for the treatment of patients with: 1. Neovascular (Wet) Age-Related Macular Degeneration (nAMD) 2. Diabetic Macular Edema (DME)	Rx by an Ophthalmologist	VABYSMO PI
	PA SUBMISSION REQUIREMENTS		
	 Diagnosis of nAMD or DME History of prior use, intolerance or contraindication to bevacizumab Patient does not have ocular or peri-ocular infections No active intraocular inflammation No concomitant use with other ophthalmic VEGF inhibitors 		

	6. Best corrected visual activity measured at baseline and periodically during treatment		
Vazalore (aspirin)	OTC Product Indicated for: 1. Pain relief 2. Reduce fever 3. Anti-inflammatory 4. Cardiovascular event prevention	Documentation of significant side effects (gastrointestinal distress, GERD, PUD, persistent nausea and vomiting, abdominal pain, etc.) with standard enteric coated aspirin tablet formulation.	VAZALORE PI
Venclexta (venetoclax)	 Indicated for: for the treatment of adult patients with chronic lymphocytic leukemia (CLL) or small lymphocytic lymphoma (SLL), with or without 17p deletion, who have received at least one prior therapy. In combination with azacitidine or decitabine or low dose cytarabine for the treatment of newly diagnosed acute myeloid leukemia (AML) in adults who are age 75 years or older, or who have comorbidities that preclude use of intensive induction chemotherapy. his indication is approved under accelerated approval based on response rates. Continued approval for this indication may be contingent upon verification and description of clinical benefit in confirmatory trials. 	Rx by Oncologist	VENCLEXTA PI
V-Go	Wearable insulin device indicated for use in adult patients requiring insulin.	Rx by Endocrinologist Please click link below for External Insulin Pump Policy: External Insulin Pump Policy	V-GO WEBSITE
Viltepso (viltolarsen)	Indicated for the treatment of Duchenne muscular dystrophy (DMD) in patients who have a confirmed mutation of the DMD gene that is amenable to exon 53 skipping. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.		VILTEPSO PI
Vitrakvi (larotrectinib)	Indicated for the treatment of adult and pediatric patients with solid tumors that: 1. have a neurotrophic receptor tyrosine kinase (NTRK) gene fusion without a known acquired resistance mutation, 2. are metastatic or where surgical resection is likely to result in severe morbidity, and 3. have no satisfactory alternative treatments or that have progressed following treatment.	Rx by Oncologist	<u>VITRAKVI PI</u>
Vizimpro (dacomitinib)	Indicated for the first-line treatment of patients with metastatic non-small cell lung cancer (NSCLC) with epidermal growth factor receptor (EGFR) exon 19 deletion or exon 21 L858R substitution mutations as detected by an FDA-approved test.	Rx by Oncologist	VIZIMPRO PI
Vimizim (elosulfase alfa)	Indicated for patients with Mucopolysaccharidosis type IVA (MPS IVA; Morquio A syndrome).		<u>VIMIZIM PI</u>

	Continued approval for this indication may be contingent upon verification and description of		
	clinical benefit in a confirmatory trial.		
Vocabria (cabotegravir)	Indicated to reduce the risk of sexually acquired HIV-1 infection in at-risk adults and adolescents weighing at least 35 kg for short-term PrEP (Pre-exposure prophylaxis). Used as oral lead in for Apretude (cabotegravir extended-release injectable suspension) to assess tolerability and oral therapy for patients who miss a planned injection of Apretude. Note: HIV-treatment is covered under Fee-for-service (FFS) for DC Healthy Families enrollees		VOCABRIA PI
	and AIDS Drug Assistance Program (ADAP) for DC Alliance enrollees.		
	PA SUBMISSION REQUIREMENTS: 1. Attestation that patient is considered high-risk for HIV infection 2. Risk-reduction and medication adherence counseling documentation 3. Negative HIV-1 test prior to initiating therapy and before subsequent use 4. Patient will not receive concomitant therapy with any of the following medications due to contraindication and decreased levels of cabotegravir seen with coadministration: a. Anticonvulsants: Carbamazepine, oxcarbazepine, phenobarbital, phenytoin b. Antimycobacterials: Rifampin, rifapentine		
Vonjo (pacritinib)	A kinase inhibitor indicated for the treatment of adults with intermediate or high-risk primary or secondary (post-polycythemia vera or post-essential thrombocythemia) myelofibrosis with a platelet count below 50×10^9 /L.	Rx by Oncologist or Hematologist	VONJO PI
	PA SUBMISSION REQUIREMENTS: 1. Laboratory studies including: serum potassium level, CBC w/differential, CMP, LFTs, INR, PT, coagulation studies 2. Platelet count below 50 × 10 ⁹ /L 3. Electrocardiogram (ECG) showing baseline QTc below 480 msec 4. Avoid concomitant use with CYP3A4 inhibitors or inducers 5. Patient does not have hepatic impairment (Child-Pugh B and Child-Pugh C) 6. No active bleeding 7. eGFR > 30 mL/min/1.72m ²		
Voxzogo (vosoritide)	Indicated to increase linear growth in pediatric patients with achondroplasia who are 5 years of age and older with open epiphyses. This indication is approved under accelerated approval based on an improvement in annualized growth velocity. Continued approval for this indication may be contingent upon verification and description of clinical benefit in confirmatory trial(s).	Rx by or in consultation with a Pediatric Endocrinologist	VOXZOGO PI
	 PA SUBMISSION REQUIREMENTS: 1. Patient's age 5 – 18 years' old 2. Diagnosis of achondroplasia 3. Genetic testing confirming a mutation in the fibroblast growth factor receptor 3 (FGFR3) gene 4. Documentation of radiographic evidence indicating open epiphyses (growth plates) 		

	 Documentation of baseline annualized growth velocity, calculated based on standing height measured over the course of 6 months prior to request Documentation of member's current weight (in kg) for appropriate dosing Voxzogo is not prescribed concurrently with any human growth hormone products (e.g., Genotropin®, Humatrope®, Norditropin®, Nutropin AQ®, Omnitrope®, Saizen®, Zomacton®) 		
Vyepti (eptinezumab-jjmr)	Indicated for the preventive treatment of migraine in adults	1.Trial and failure or intolerance to at least three of the following agents: beta blockers, topiramate, Aimovig, Emgality, and Ubrelvy, in medical documentation submitted. 2. Patient must have at least 4 headache days per month on average.	VYEPTI PI
Vyvgart (efgartigimod alfa-fcab)	Indicated for the treatment of generalized myasthenia gravis (gMG) in adult patients who are anti-acetylcholine receptor (AChR) antibody positive	Rx by or in consultation with a Neurologist	VYVGART PI
	PA SUBMISSION REQUIREMENTS: 1. Diagnosis of gMG 2. Patient age at least 18 years 3. Myasthenia Gravis-Activities of Daily Living (MG-ADL) score ≥ 5 at baseline 4. Greater than 50% of the baseline MG-ADL score is due to non-ocular symptoms 5. Myasthenia Gravis Foundation of America (MGFA) clinical classification of Class II to IV 6. Documentation of positive serologic test for anti-AChR antibodies 7. Trial and failure of or documented intolerance or contraindication to a cholinesterase inhibitor 8. Trial and failure of or documented intolerance or contraindication to a corticosteroid 9. Trial and failure of or documented intolerance to at least two of the following immunosuppressive therapies or contraindication to all of the therapies below: a. Rituximab or biosimiliar Truxima b. Cyclophosphamide c. Azathioprine d. Mycophenolate mofetil 10. Vyvgart is not prescribed concurrently with Soliris® 11. Documentation of member's current weight (in kg) for appropriate dosing 12. Dose does not exceed 10 mg/kg (1,200 mg per infusion for members weighing 120 kg		
Xadago	or more) once weekly for the first 4 weeks of every 8-week cycle. Indicated as adjunctive treatment to levodopa/carbidopa in patients with Parkinson's disease	Rx by Neurologist	XADAGO PI
(safinamide)	(PD) experiencing "off" episodes.	. 0	

Xalkori (crizotinib)	Indicated for the treatment of patients with metastatic non-small cell lung cancer (NSCLC) whose tumors are anaplastic lymphoma kinase (ALK) or ROS1-positive as detected by an FDA-approved test.	Rx by Oncologist	XALKORI PI
Xenazine (tetrabenazine)	Indicated for the treatment of chorea associated with Huntington's disease.	Rx by Neurologist	XENAZINE PI
Xgeva (denosumab)	 Indicated for: Prevention of skeletal-related events in patients with multiple myeloma and in patients with bone metastases from solid tumors. Treatment of adults and skeletally mature adolescents with giant cell tumor of bone that is unresectable or where surgical resection is likely to result in severe morbidity. Treatment of hypercalcemia of malignancy refractory to bisphosphonate therapy. 	Rx by Oncologist	XGEVA PI
Xiidra (lifitegrast ophthal)	Indicated for the treatment of the signs and symptoms of dry eye disease.	Must have tried and failed artificial tears.	XIIDRA PI
Xolair (omalizumab)	 Indicated for: moderate to severe persistent asthma in patients 6 years of age and older with a positive skin test or in vitro reactivity to a perennial aeroallergen and symptoms that are inadequately controlled with inhaled corticosteroids. chronic idiopathic urticaria in adults and adolescents (12 years of age and above) who remain symptomatic despite H1 antihistamine treatment. 	Rx by Allergist or Pulmonologist Regarding ASTHMA indication only: 1. moderate to severe persistent ALLERGIC asthma (confirmed by a positive skin test or RAST for ≥ 1 perennial aeroallergen) 2. IgE level obtained prior to initiation of therapy 3. currently using an inhaled corticosteroid at maximum dose; compliance must be confirmed in the patient's Caremark profile 4. currently using a long-acting inhaled beta ₂ -agonist OR a leukotriene modifier; compliance must be confirmed in the patient's Caremark profile NOT approved for monotherapy	XOLAIR PI
Xospata (gilteritinib)	Indicated for the treatment of adult patients who have relapsed or refractory acute myeloid leukemia (AML) with a FLT3 mutation as detected by an FDA-approved test.	Rx by Oncologist	XOSPATA PI
Xpovio (selinexor)	Indicated for in combination with dexamethasone for the treatment of adult patients with relapsed or refractory multiple myeloma (RRMM) who have received at least four prior therapies and whose disease is refractory to at least two proteasome inhibitors, at least two immunomodulatory agents, and an anti-CD38 monoclonal antibody.	Rx by Oncologist	XPOVIO

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	This indication is approved under accelerated approval based on response rate. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial		
Xyrem (sodium oxybate)	Indicated for: 1. for the treatment of cataplexy or excessive daytime sleepiness (EDS) in patients 7 years of age and older with narcolepsy. ***Xyrem may only be dispensed to patients enrolled in the Xyrem Success Program	Rx by Neurologist 1. patient > 16 years old 2. alternative diagnoses must have been excluded 3. for cataplexy, must have failed tricyclic or SSRIs 4. for excessive daytime sleepiness, must have failed at least one formulary stimulant treatment (ex: methylphenidate or dextroamphetamine) 5. initial approval for maximum of 1-month supply with subsequent renewals for maximum approval period of 3 months at a time (Patients are to be re-evaluated by physician no less frequently than every 3 months)	XYREM PI
Yescarta (axicabtagene ciloleucel)	Indicated for the treatment of adult patients with relapsed or refractory large B-cell lymphoma after two or more lines of systemic therapy, including diffuse large B-cell lymphoma (DLBCL) not otherwise specified, primary mediastinal large B-cell lymphoma, high grade B-cell lymphoma, and DLBCL arising from follicular lymphoma. MedStar Family Choice considers Yescarta (Axicabtagene Ciloleucel) medically necessary when ALL of the following criteria are met: 1. Recipient is 18 years of age or older; AND 2. Histologically confirmed diagnosis of one of the following types of aggressive non-Hodgkin's lymphoma a. Diffuse large B-cell lymphoma (DLBCL), not otherwise specified; or b. High-grade B-cell lymphoma; or c. Primary mediastinal large B-cell lymphoma; or d. Transformed follicular lymphoma; AND 3. Relapsed or refractory disease, when a. Recipient has previously received two or more lines of systemic therapy; and b. Disease is refractory to the most recent therapy or relapsed within 1 year after autologous hematopoietic stem cell transplantation (HSCT); AND	Rx by Oncologist	YESCARTA PI

Zejula (niraparib) Indicated for: 1. for the maintenance treatment of adult patients with recurrent epithelial ovarian, fallopian tube, or primary peritoneal cancer who are in a complete or partial response to platinum-based chemotherapy. 2. for the treatment of adult patients with advanced ovarian, fallopian tube, or primary peritoneal cancer who have been treated with three more prior chemotherapy regimens and whose cancer is associated with homologous recombination deficiency (HRD) positive status defined by either: • a deleterious or suspected deleterious BRCA mutation, or • genomic instability and who have progressed more than six months after response to the last platinum-based chemotherapy. Zelboraf (vemurafenib) Indicated for: 1. the treatment of patients with unresectable or metastatic melanoma with BRAF V600E mutation as detected by an FDA-approved test.		 4. Must have received adequate prior therapy including, at a minimum, all of the following: a. An anthracycline-containing chemotherapy regimen; and b. For CD20+ disease, anti-CD20 monoclonal antibody; and c. For subjects with transformed follicular lymphoma, prior chemotherapy for follicular lymphoma with chemotherapy refractory disease after transformation to DLBCL; AND 5. Documentation of all of the following clinical findings: a. Eastern Cooperative Oncology Group (ECOG) performance status of 0-1; and b. Adequate cardiac, pulmonary, and other organ function (as determined by protocol from treatment facility); AND 6. The treatment facility that dispenses and administers Yescarta is enrolled and complies with the Risk Evaluation and Mitigation Strategy; AND 7. One-time, single administration with dosing in accordance with the FDA label Yescarta (Axicabtagene ciloleucel) is considered investigational and not medically necessary when the above medically necessary criteria are not met, and for all other indications, including but not limited to: History of malignancy other than nonmelanoma skin cancer or carcinoma in situ (e.g. cervix, bladder, breast) or follicular lymphoma unless disease free for at least 3 years; or Any central nervous system (CNS) disease, for example, detectable CSF malignant cells, brain metastases, CNS lymphoma, or a history or presence of CNS disorders such as seizure disorder, cerebrovascular ischemia/hemorrhage, dementia, cerebellar disease, or autoimmune disease with CNS involvement; or History of allogeneic stem cell transplant, chimeric antigen receptor therapy or other genetically modified T-cell therapy; or Active, uncontrolled infection; or Human immunodeficiency virus (HIV); or Hepatitis B or C (if viral load is detectable). 		
fallopian tube, or primary peritoneal cancer who are in a complete or partial response to platinum-based chemotherapy. 2. for the treatment of adult patients with advanced ovarian, fallopian tube, or primary peritoneal cancer who have been treated with three more prior chemotherapy regimens and whose cancer is associated with homologous recombination deficiency (HRD) positive status defined by either: • a deleterious or suspected deleterious BRCA mutation, or • genomic instability and who have progressed more than six months after response to the last platinum-based chemotherapy. Zelboraf (vemurafenib) Indicated for: 1. the treatment of patients with unresectable or metastatic melanoma with BRAF V600E mutation as detected by an FDA-approved test.	Zejula (niraparib)	Indicated for:	Rx by Oncologist	ZEJULA PI
1. the treatment of patients with unresectable or metastatic melanoma with BRAF V600E Dermatologist mutation as detected by an FDA-approved test.		fallopian tube, or primary peritoneal cancer who are in a complete or partial response to platinum-based chemotherapy. 2. for the treatment of adult patients with advanced ovarian, fallopian tube, or primary peritoneal cancer who have been treated with three more prior chemotherapy regimens and whose cancer is associated with homologous recombination deficiency (HRD) positive status defined by either: • a deleterious or suspected deleterious BRCA mutation, or • genomic instability and who have progressed more than six months after response to the last platinum-based chemotherapy.		
	Zelboraf (vemurafenib)	1. the treatment of patients with unresectable or metastatic melanoma with BRAF V600E		ZELBORAF PI

	2. the treatment of patients with Erdheim Chester Disease with BRAF V600 mutation.		
Zepzelca	Indicated for the treatment of adult patients with metastatic small cell lung cancer (SCLC) with		ZEPZULCA PI
(lurbinectedin)	disease progression on or after platinum -based chemotherapy.		
Zoladex	Indicated for:	Rx by Oncologist	ZOLADEX 3.6 mg PI
(goserelin)	1. palliative treatment of advanced carcinoma of the prostate. (3.6 and 10.8 mg)		
	2. use in combination with flutamide for the management of locally confined stage T2b-T4		
	(Stage B2-C) carcinoma of the prostate. (3.6 and 10.8 mg)		ZOLADEX 10.8 mg PI
	3. management of endometriosis. (3.6 mg)		
	4. palliative treatment of advanced breast cancer in pre- and peri-menopausal women. (3.6 mg)		
	5. use as an agent to cause endometrial thinning agent prior to endometrial ablation for dysfunctional uterine bleeding. (3.6 mg)		
	6. for the management of endometriosis, including pain relief and reduction of		
	endometriotic lesions for the duration of therapy. Experience with ZOLADEX for the		
	management of endometriosis has been limited to women 18 years of age and older		
	treated for 6 months.		
Zolgensma	Indicated for the treatment of pediatric patients less than 2 years of age with spinal muscular	***MFC-DC	ZOLGENSMA PI
(onasemnogene	atrophy (SMA) with bi-allelic mutations in the survival motor neuron 1 gene.		
abeparvovec-xioi)		Pharmacist	
	Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.	Review***	
Zontivity	Indicated for the reduction of thrombotic cardiovascular events in patients with a history of	Rx by Cardiology, Neurology or	ZONTIVITY PI
(vorapaxar)	myocardial infarction (MI) or with peripheral arterial disease (PAD).	Vascular Surgery	
Zurampic	Indicated in combination with a xanthine oxidase inhibitor for the treatment of hyperuricemia	Rx by Rheumatologist	ZURAMPIC PI
(lesinurad)	associated with gout in patients who have not achieved target serum uric acid levels with a		
	xanthine oxidase inhibitor alone.		
Zydelig	Indicated for:	Rx by Oncologist	ZYDELIG PI
(idelalisib)	1. treatment of patients with relapsed chronic lymphocytic leukemia (CLL), in combination		
	with rituximab, in patients for whom rituximab alone would be considered appropriate		
	therapy due to other co-morbidities.		
	2. treatment of patients with relapsed follicular B-cell non-Hodgkin lymphoma (FL) in		
	patients who have received at least two prior systemic therapies.		
	3. treatment of patients with relapsed small lymphocytic lymphoma (SLL) in patients who have received at least two prior systemic therapies.		
Zykadia	Indicated for the treatment of adults with metastatic non-small cell lung cancer (NSLC) whose	Rx by Oncologist	ZYKADIA PI
(ceritinib)	tumors are anaplastic lymphoma kinase-positive as detected by an FDA-approved test.	Total Sy Street Spice	2110101111
Zynlonta	Indicated for the treatment of adult patients with relapsed or refractory large B-cell lymphoma	Rx by Oncologist	ZYNLONTA PI
(loncastuximab tesirine-	after two or more lines of systemic therapy, including diffuse large B-cell lymphoma (DLBCL)	, 3	
lpyl)	not otherwise specified, DLBCL arising from low-grade lymphoma, and high-grade B-cell		
,	lymphoma.		