

Prior Authorization – Approval in advance to get certain drugs that may or may not be on our formulary. Some drugs may be covered only if your doctor or other network provider gets "prior authorization" 1 us. Covered drugs that need prior authorization are marked in the formulary.

MATERIAL	us. Covered drugs that need pri	or authorization are marked in the formulary.							
MANUSCRIPT   MAN		Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
DEPOSE FOR ATTERIOR PROPERTY OF THE PROPERTY	5HT3 ANTI-NAUSEA G: AGENT BVD H		PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE						
DIRECTION OF THE PROPERTY OF T	ABATACEPT O	RENCIA			IDIOPATHIC ARTHRITIS: EXPERIENCED OR MAINTAINED 20% OR GREATER IMPROVEMENT IN		SUPERVISED BY A		(METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE) AND TRIAL OF HUMIRA OR CIMZIA. FOR JUVENILE IDIOPATHIC ARTHRITIS: TRIAL OF AT LEAST ONE OF THE FOLLOWING: TRIAL/FAILURE OF AT LEAST ONE DMARD (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE)
ADD-REASTIZIMANI KAIXYLA ALL MEMERIALY ACCEPTED PROBLEMENTS NOT OTHERWISE EXCLUDED PROM PART D.  SEMANSEN.  OTHERWISE EXCLUDED PROM PART D.  OTHERWISE EXCLUDED PROM	ABATACEPT SQ O	RENCIA			EXPERIENCED OR MAINTAINED 20% OR GREATER IMPROVEMENT IN TENDER AND SWOLLEN JOINT	18 YEARS OR OLDER.	SUPERVISED BY A		"INITIAL: RHEUMATOID ARTHRITIS: TRIALFAILURE OF AT LEAST ONE DMARD (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE) AND TRIAL OF HUMIRA OR CIMZIA."
ADD-TRASTIZUMAB RACYLA ALL MEDICALLY ACCEPTED DIODECTIONS NOT OTHERWISE EXCLUDED FROM PART D  ALL MEDICALLY ACCEPTED DIODECTIONS NOT DIERWISE EXCLUDED FROM PART D  ALL MEDICALLY ACCEPTED DIODECTIONS NOT DIERWISE EXCLUDED FROM PART D  ARRANGA  ANASTRO  ANA	ABIRATERONE Z	YTIGA						12 MONTHS	
ANARINGA ANA		ADCYLA	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	
ANALYSIA NEET OLD APPROVED INDICATIONS NOT OTHERWISE EXCLLEDE PROMPART D.  APREPITANT IND DETERMINATION DETERMINED AND SWOLLEN JOINT COUNT.  APREPITANT IND DETERMINATION PRICE OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST ANCES, INFORMATION MAY BEED TO BE SERVED BY ANALYSIN OF THE CIRCUMST AND ANALYSIN OF THE CIRCUMST ANALYSIN OF THE CIRCUMST AND ANALYSIN OF THE CI		ALTRAP	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	
ARREPTIANT BY D DETERMINATION	ANAKINRA K	INERET	ALL FDA APPROVED INDICATIONS NOT		EXPERIENCED OR MAINTAINED 20% OR GREATER IMPROVEMENT IN TENDER AND SWOLLEN JOINT		SUPERVISED BY A	RENEWAL: 12 MONTHS	. TRIAL/FAILURE OF AT LEAST ONE DMARD (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE)
INHIBITORS  OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT  OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT  OTHERWISE EXCLUDED FROM PART D.  12 MONTHS  TRIAL OF AT LEAST ONE:  FOR THE TREATHOR OF ITE (SUNITINIS), VOTRIENT (P.  (SUNITINIS), VOTRIENT (P.  (BEVACIZUMAB) IN COMB  INTERFERON."  AUTOANTIBODY POSITIVE LUPUS TEST.  12 MONTHS  TINTIAL: SELENA-SELDAT  OR EQUAL TO 6, REMEDIE  CONTROS FROM BASELINA:  ALL FDA APPROVED INDICATIONS NOT  OTHERWISE EXCLUDED FROM PART D.  AUTOANTIBODY POSITIVE LUPUS TEST.  12 MONTHS  TINTIAL: SELENA-SELDAT  OR EQUAL TO 6, REMEDIE  CONTROS FROM BASELINA:  MMUNOSUPPRISTS WE AGE  CONTROS FROM BASELINA:  MMUNOSUPRISTS SEVER AGE  FOR DIAGRAPHS, OR INT  CYCLOPHOSAMIDE:  BEVACIZUMAB  AVASTIN  ALL MEDICALLY ACCEPTED INDICATIONS NOT		MEND	PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE						
AXTINIB  INLYTA  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  INLYTA  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  INLYTA  ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  AUTOANTIBODY POSITIVE LUPUS TEST.  INLYTA  AUTOANTIBODY POSITIVE LUPUS TEST.  INLYT		NASTROZOLE   EXEMESTANE   LETROZOLE						12 MONTHS	
OTHERWISE EXCLUDED FROM PART D.  OR EQUAL TO 6. RENEWAI 4 POINT REDUCTION IN SEE FROM BASELINE. MEMBER CORTICOSTEROIDS, ANTIN IMMUNOSUPPRESSIVE AGI FOR DIAGNOSIS OF SEVER NETHERITS OR SEVERE CEI SYSTEM LUPUS OR CONCLUDIOGIC AGENTS, OR INTOCYCLOPHOSAMIDE."  BEVACIZUMAB AVASTIN ALL MEDICALLY ACCEPTED INDICATIONS NOT		NLYTA	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	"TRIAL OF AT LEAST ONE SYSTEMIC THERAPY FOR THE TREATMENT OF RCC SUCH AS NEXAVAR (SORAFENIB), TORISEL (TEMSIROLIMUS), SUTENT (SUNITINIB), VOTRIENT (PAZOPANIB), OR AVASTIN (BEVACIZUMAB) IN COMBINATION WITH INTERFERON."
	BELIMUMAB BI	ENLYSTA			AUTOANTIBODY POSITIVE LUPUS TEST.			12 MONTHS	"INITIAL: SELENA-SELDAI SCORE GREATER THAN OR EQUAL TO 6. RENEWAL: MAINTAIN AT LEAST A 4 POINT REDUCTION IN SELENA-SELDAI SCORE FROM BASELINE. MEMBER IS CURRENTLY TAKING CORTICOSTEROIDS, ANTIMALARIALS, NSAIDS, OR IMMUNOSUPPRESSIVE AGENTS. NO APPROVAL FOR DIAGNOSIS OF SEVERE ACTIVE LUPUS NEPHRITIS OR SEVERE CENTRAL NERVOUS SYSTEM LUPUS OR CONCURRENT USE OF BIOLOGIC AGENTS, OR INTRAVENOUS CYCLOPHOSAMIDE."
OTHER WIDE EACEODED FROM FAIR D.	BEVACIZUMAB A	VASTIN						12 MONTHS	
BEXAROTENE TARGRETIN ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	BEXAROTENE TA	ARGRETIN	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	

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Prior Authorization	Drug Nama	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
Group Description	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
BOCEPREVIR	VICTRELIS	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	"TREATMENT WITH BOCEPREVIR WILL NOT BE APPROVED FOR A PATIENT WHO HAS FAILED SHORT TRIAL OR HAS CONTRAINDICATION TO TELAPREVIR (INCIVEK) OR HAS PREVIOUS FAILURE OF FULL COURSE OF TRIPLE THERAPY WITH TELAPREVIR (INCIVEK) OR BOCEPREVIR (VICTRELIS) OR CURRENTLY TAKING CARBAMAZEPINE, PHENOBARBITAL, PHENYTOIN, OR RIFAMPIN OR HAS A CO-INFECTION WITH HEPATITIS B. DETECTABLE HCV RNA LEVEL/VIRAL LOAD OR HCV RNA LEVEL/VIRAL LOAD GREATER THAN OR EQUAL TO 100 IU/ML AFTER TRIPLE THERAPY."	"CHRONIC HEPATITIS C, GENOTYPE I. NATIVE PATIENT: HCV RNA LEVEL/VIRAL LOAD AT TRIPLE THERAPY TREATMENT WEEK 4, 8, 12, AND 24 OF BOCEPREVIR THERAPY. PARTIAL RESPONDER, NULL RESPONDER, OR RELAPSER: HCV RNA LEVEL/VIRAL LOAD AT WEEK 8 AND 20 OF BOCEPREVIR THERAPY. RENEWAL HCV RNA LEVELS TO DETERMINE LENGTH OF TREATMENT."	PATIENT 18 YEARS OF AGE OR OLDER.	T, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE		CONCURRENT USE OF RIBAVIRIN AND PEGINTERFERON ALFA.
BORTEZOMIB	VELCADE	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	
BOSUTINIB	BOSULIF	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	CML: BCR-ABL MUTATIONAL ANALYSIS CONFIRMING THAT BOTH T3151 AND V299L MUTATIONS ARE NOT PRESENT.
	CINRYZE	ALL FDA APPROVED INDICATIONS NOT				"HEMATOLOGIST,	12 MONTHS	TRIAL OF OR INTOLERABLE SIDE EFFECTS TO
INHIBITOR CABOZANTINIB	COMETRIQ	OTHERWISE EXCLUDED FROM PART D. ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.				IMMUNOLOGIST"	12 MONTHS	DANAZOL.
CALCINEURIN INHIBITORS	ELIDEL   PROTOPIC	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	NOT TRIED/FAILED OR INTOLERABLE ADVERSE EFFECTS TO TOPICAL CORTICOSTEROIDS		ELIDEL 1% AND PROTOPIC 0.03%: 2 YEARS OR OLDER. PROTOPIC 0.1%: OVER 14 YEARS.		12 MONTHS	
CANAKINUMAB	ILARIS	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			CAPS: 4 YEARS AND OLDER. SJIA: 2 YEARS AND OLDER.	PRESCRIBED OR SUPERVISED BY RHEUMATOLOGIST	12 MONTHS	
CERTOLIZUMAB PEGOL	CIMZIA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		RENEWAL: RHEUMATOID ARTHRITIS: EXPERIENCED OR MAINTAINED 20% OR GREATER IMPROVEMENT IN TENDER AND SWOLLEN JOINT COUNT.		PRESCRIBED BY OR SUPERVISED BY A GASTROENTEROLOGIS T OR RHEUMATOLOGIST.	INITIAL: 3 MONTHS RENEWAL: 12 MONTHS	"FOR MODERATE TO SEVERE CROHN'S DISEASE: TRIAL/FAILURE OF ONE OR MORE CONVENTIONAL THERAPIES FOR CROHN'S DISEASE SUCH AS CORTICOSTEROIDS, AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. FOR MODERATE TO SEVERE RHEUMATOID ARTHRITIS: TRIAL/FAILURE OF AT LEAST ONE DMARD AGENT (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE)."
CETUXIMAB	ERBITUX	ALL MEDICALLY ACCEPTED INDICATIONS NOT		METASTATIC COLORECTAL CANCER : WILD TYPE			12 MONTHS	
CHENODIOL	CHENODAL	OTHERWISE EXCLUDED FROM PART D. ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CEREBROTENDINOUS XANTHOMATOSIS.	RADIOLUCENT GALLSTONES: NO FAILED TREATMENT WITH URSODIOL	KRAS (WITHOUT MUTATION)			12 MONTHS	
	ARICEPT	ALL FDA APPROVED INDICATIONS NOT		MINI MENTAL STATE EXAM (MMSE) SCORE OF 26			12 MONTHS	
INHIBITORS CLOBAZAM	ONFI	OTHERWISE EXCLUDED FROM PART D. ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		OR LESS	2 YEARS OF AGE OR OLDER		12 MONTHS	TRIAL OF LAMOTRIGINE OR TOPIRAMATE.
DETERMINATION		OTHERWISE EACLUSED FROM PART THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.			JOLDIER .			
	H.P. ACTHAR	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	USED FOR DIAGNOSTIC PURPOSES. ACUTE EXACERBATION OF MULTIPLE SCLEROSIS: IV ACCESS OR IV ACCESS CAN BE OBTAINED.		INFANTILE SPASMS: LESS THAN 2 YEARS OF AGE.	,	INFANTILE SPASMS: 28 DAYS. MULTIPLE SCLEROSIS: 21 DAYS.	
CRIZOTINIB	XALKORI	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		LOCALLY ADVANCED OR METASTATIC NON SMALL CELL LUNG CANCER IS ANAPLASTIC LYMPHOMA KINASE POSITIVE.			12 MONTHS	

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Prior Authorization Group Description	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
	CYCLOPHOSPHAMIDE	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES, INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
CYCLOSPORINE OPHTHALMIC	RESTASIS	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		KERATOCONJUNCTIVITIS SICCA (KCS) OR DRY EYE DISEASE.		"PRESCRIBED BY OR SUPERVISED BY A OPHTHALMOLOGIST, OPTOMETRIST, OR RHEUMATOLOGIST."	12 MONTHS	
DABIGATRAN	PRADAXA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	TRIAL OF OR CONTRAINDICATION TO XARELTO.
DABRAFENIB MESYLATE	TAFINLAR	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	
DALFAMPRIDINE	AMPYRA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		WALKING DISABILITY SUCH AS MILD TO MODERATE BILATERAL LOWER EXTREMITY WEAKNESS OR UNILATERAL WEAKNESS PLUS LOWER EXTREMITY OR TRUNCAL ATAXIA.		NEUROLOGIST	INITIAL: 3 MONTHS. RENEWAL: 12 MONTHS	RENEWAL: PATIENT HAS EXPERIENCED OR MAINTAINED AT LEAST 15% IMPROVEMENT IN WALKING ABILITY.
DASATINIB	SPRYCEL	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	"PREVIOUSLY TREATED CML REQUIRES MUTATIONAL ANALYSIS NEGATIVE FOR THE FOLLOWING MUTATIONS FOLLOWING BCR-ABL MUTATIONAL ANALYSIS - T3151, V299L, T315A, F317L/V/I/C."
DENOSUMAB	PROLIA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		"A PATIENT WITH EITHER A HISTORY OF OSTEOPORTIC FRACTURE(S) OR GREATER THAN OR EQUAL TO TWO FACTORS FOR FRACTURE (E.G. HISTORY OF MULTIPLE RECENT LOW TRAUMA FRACTURES, BMD T-SCORE LESS THAN OR EQUAL TO -2.5, CORTICOSTEROID USE, OR USE OF GNRH ANALOGS), OR FAILED AN ADEQUATE TRIAL OF BISPHOSPHONATES, IS INTOLERANT, OR HAS A CONTRAINDICATION TO BISPHOSPHONATES."			12 MONTHS	
DENOSUMAB-XGEVA	XGEVA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	DIAGNOSIS OF MULTIPLE MYELOMA				12 MONTHS	
DIMETHYL FUMARATE	TECFIDERA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			18 YEARS AND OLDER		12 MONTHS	"TRIAL OF OR CONTRAINDICATION TO INTERFERON THERAPY (SUCH AS REBIF, AVONEX BETASERON, EXTAVIA) AND COPAXONE."
ENDOTHELIN RECEPTOR ANTAGONISTS	LETAIRIS   TRACLEER	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		DIAGNOSIS OF PULMONARY ARTIERAL HYPERTENTION GREATER OR EQUAL TO NYHA/WHO FUNCTIONAL CLASS II.		CARDIOLOGIST OR PULMONOLOGIST.	12 MONTHS	
ENZALUTAMIDE	XTANDI	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	TRIAL OF OR CONTRAINDICATION TO DOCETAXEL.
EPIDERMAL GROWTH FACTOR RECEPTOR INHIBITORS - ERLOTNIB		ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	
ERIBULIN	HALAVEN	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	"PREVIOUS TREATMENT WITH AN ANTHRACYCLINE (DAUNORUBICIN, DOXORUBICIN, IDARUBICIN, EPIRUBICIN, OR MITOXANTRONE) AND A TAXANE (DOCETAXEL OR PACLITAXEL)."

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Prior Authorization	Drug Nama	Covered Uses	Exclusion Criteria	Required Medical Information	Aga Dastwistians	Proscriber Destrictions	Coverage Duration	Other Cuitouis
	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	rrescriber Restrictions	Coverage Duration	Otner Criteria
Prior Authorization Group Description ERYTHROPOIESIS STIMULATING AGENTS - EPOETIN ALFA	Drug Name  EPOGEN   PROCRIT	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ADDITIONAL OFF LABEL ANEMIA IN HEPATITIS C BEING TREATED IN COMBINATION WITH RIBAVIRIN AND AN INERFERON ALFA OR PEGINTERFERON ALFA.	Exclusion Criteria	"CHRONIC RENAL FAILURE HEMAGLOBIN LEVELS LESS THAN 10 G/DL IF NOT ON DIALYSIS AND LESS THAN 11 G/DL IF ON DIALYSIS OR HEMOGLOBIN HAS REACHED 11 G/DL IF ON DIALYSIS AND DOSE REDUCTION/INTERRUPITON IS REQUIRED TO REDUCE THE NEED FOR BLOOD TRANSFUSIONS OR HEMOGLOBIN HAS REACHED 10 G/DL IF NOT ON DIALYSIS AND DOSE REDUCTION/INTERRUPTION IS REQUIRED TO REDUCE THE NEED FOR BLOOD TRANSFUSIONS. ANEMIA DUE TO EFFECT OF CONCOMITANTLY ADMINISTERED CANCER CHEMOTHERAPY: HEMOGLOBIN LEVELS BETWEEN 10 AND 12 G/DL OR HEMOGLOBIN LEVEL LESS THAN 11 G/DL OR HEMOGLOBIN LEVEL DECREASED AT LEAST 2 G/DL BELOW THEIR BASELINE. ZIDOVUDINE THERAPY: HEMOGLOBIN LESS THAN 10 G/DL ELECTIVE, NONCARDIAC, NONVASCULAR SURGERY: HEMOGLOBIN LESS THAN 10 G/DL ELECTIVE, NONCARDIAC, NONVASCULAR SURGERY: HEMOGLOBIN LESS THAN 10 G/DL ESS DETWEEN 10 AND 12 G/DL OR SES BETWEEN 10 AND 12 G/DL OR SES BETWEEN 10 AND 12 G/DL TESS BETWEEN 10 AND 12 G/DL TESS BETWEEN 10 AND 12 G/DL TESS BETWEEN 10 AND 12 G/DL TO RESS BETWEEN 10 AND 12 G/DL TO RESS BETWEEN 10 AND 12 G/DL FOR PATIENTS CURRENTLY TAKING REQUESTED MEDICATION OR CONTRAINDICATION TO RIBAVIRIN DOSE REDUCTION AND HEMOGLOBIN LESS THAN 10 G/DL FOR NEW	Age Restrictions	Prescriber Restrictions	ANEMIA FROM MYBLOSUPPRESSIVE CHEMO/CKD W/O DIALYSIS/ZIDOVUDINE 12 MOS, SURGERY:1 MO. HEP C:6 MOS.	Other Criteria  ALL INDICATIONS: TRIAL OF PROCRIT. PART D MEMBER RECEIVING DIALYSIS OR IDENTIFIED AS A PART D END STAGE RENAL DISEASE MEMBER: PAYS UNDER PART B.
				STARTS."				
	BONIVA   CALCITRIOL   CUBICIN   HECTOROL   HEPARIN SODIUM   LEVOCARNITINE   LIDOCAINE   LIDOCAINE HCL   LIDOCAINE-PRILOCAINE   MIACALCIN   PAMIDRONATE DISODIUM   VANCOMYCIN HCL   ZEMPLAR	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
EVEROLIMUS	AFINITOR	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	ADVANCED RENAL CELL CARCINOMA (RCC): TRIAL OF OR CONTRAINDICATION TO SUTENT OR NEXAVAR.
FENTANYL NASAL SPRAY	LAZANDA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					6 MONTHS	"CANCER: CURRENTLY ON A MAINTENANCE DOSE OF CONTROLLED-RELEASE OPIOID PAIN MEDICATION (SUCH AS MORPHINE SULFATE SR, OXYCODONE SR, OR FENTANYL). EITHER A TRIAL OR CONTRAINDICATION TO AT LEAST ONE (1) IMMEDIATE-RELEASE ORAL OPIOID PAIN AGENT (SUCH AS MORPHINE SULFATE IR, OXYCODONE/ASPIRIN, OXYCODONE/ASPIRIN, OXYCODONE/ACETAMINOPHEN, CODEINE/ACETAMINOPHEN, HYDROMORPHONE, OR MEPERIDINE) OR MEMBER HAS DIFFICULTY SWALLOWING TABLETS/CAPSULES AND TRIAL OR CONTRAINDICATION TO GENERIC FENTANYL CITRATE LOZENGE."
FENTANYL TRANSDERMAL PATCH	FENTANYL	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	TRIAL OF OR CONTRAINDICATION TO AT LEAST ONE SUSTAINED-RELEASE MORPHINE PRODUCT. EVERY 48 HOUR DOSING CONSIDERED FOR PATIENTS WHO FAIL EVERY 72 HOUR DOSING. NO APPROVAL WHEN PRESCRIBED FOR AS NEEDED DOSAGE FREQEUNCY.
FENTANYL TRANSMUCOSAL AGENTS - FENTANYL CITRATE	FENTANYL CITRATE	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					6 MONTHS	"CANCER: CURRENTLY ON A MAINTENANCE DOSE OF CONTROLLED-RELEASE OPIOID PAIN MEDICATION (SUCH AS MORPHINE SULFATE SR, OXYCODONE SR, OR FENTANYL). EITHER A TRIAL OR CONTRAINDICATION TO AT LEAST ONE (1) IMMEDIATE-RELEASE ORAL OPIOID PAIN AGENT (SUCH AS MORPHINE SULFATE IR, OXYCODONE/ASPIRIN, OXYCODONE/ASPIRIN, OXYCODONE/ACETAMINOPHEN, HYDROMORPHONE, OR MEPERIDINE) OR MEMBER HAS DIFFICULTY SWALLOWING TABLETS/CAPSULES."

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FINGOLIMOD	GILENYA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	"TRIAL OR CONTRAINDICATION TO INTERFERON THERAPY (AVONEX, BETASERON, EXTAVIA, OR REBIF) AND COPAXONE, OR RAPIDLY PROGRESSING DISEASE WHILE ON INTERFERON THERAPY OR COPAXONE."
GLP-1 ANALOGS	VICTOZA 3-PAK	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	"FAILURE TO REACH TREATMENT GOALS WITH METFORMIN, METFORMIN ER, GLYBURIDE/METFORMIN, GLIPIZIDE/METFORMIN, A FORMULARY SULFONYLUREA (GLYBURIDE, GLIPIZIDE), PIOGLITAZONE (ACTOS), PIOGLITAZONE/METFORMIN (ACTOSPLUS MET), OR PIOGLITAZONE/GLIMEPIRIDE (DUETACT) AND EXENATIDE EXTENDED RELEASE (BYDUREON)."
GLYCEROL	RAVICTI	ALL FDA APPROVED INDICATIONS NOT					12 MONTHS	TRIAL OF OR CONTRAINDICATION TO SODIUM
PHENYLBUTYRATE GOLIMUMAB	SIMPONI	OTHERWISE EXCLUDED FROM PART D. ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		RENEWAL: ACTIVE RHEUMATOID ARTHRITIS/PSORIATIC ARTHRITIS: MAINTAINED OR EXPERIENCED GREATER THAN 20% IMPROVEMENT IN TENDER JOINT COUNT AND SWOLLEN JOINT COUNT. ANKYLOSING SPONDYLITIS: MAINTAINED OR EXPERIENCED GREATER THAN 20% IMPROVEMENT IN ANKYLOSING SPONDYLITIS (ASAS20) CRITERIA.	18 YEARS OR OLDER	"PRESCRIBED BY OR SUPERVISED BY A RHEUMATOLOGIST, DERMATOLOGIST, OR GASTROENTEROLOGIS T."	INITIAL: 3 MONTHS RENEWAL: 12 MONTHS UC: 12 MONTHS.	PHENYLBUTYRATE (BUPHENYL).  "ACTIVE RHEUMATOID ARTHRITIS INITIAL: TRIAL OF HUMIRA OR CIMZIA AND TRIAL/FAILURE OF AT LEAST ONE DMARD AGENT (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE). PSORIATIC ARTHRITIS: TRIAL OF HUMIRA AND TRIAL/FAILURE OF AT LEAST ONE DMARD (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE) AND HUMIRA. ANKYLOSING SPONDYLITIS: TRIAL OF HUMIRA. ULCERATIVE COLITIS: TRIAL OF OR CONTRAINDICATION TO SULFASALAZINE, CORTICOSTEROIDS, METHOTREXATE, AZATHIOPRINE, OLSALAZINE, MESALAMINE, CYCLOSPORINE, OR MERCAPTOPURINE."
HEPATITIS A VACCINE (INACTIVATED) BVD DETERMINATION	HAVRIX   VAQTA	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
HEPATITIS B VACCINE BVD DETERMINATION	ENGERIX-B   RECOMBIVAX HB	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
HIGH RISK DRUGS IN THE ELDERLY - ANTI- INFECTIVE	NITROFURANTOIN	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	MEMBERS 65 YEARS OR OLDER WILL BE EVALUATED FOR LIMITED PRESCRIPTION USE TO NO MORE THAN 90 DAYS (TOTAL) OF CUMULATIVE USE. REQUESTS FOR GREATER THAN 90 DAYS OF CUMULATIVE USE WILL REQUIRE TRIAL OF OR CONTRAINDICATION TO SULFAMETHOXAZOLE/TRIMETHOPRIM (TMP-SMX) OR TRIMETHOPRIM.
HIGH RISK DRUGS IN THE ELDERLY - ANTICHOLINERGICS - BENZTROPINE TRIHEXYPHENIDYL	BENZTROPINE MESYLATE   TRIHEXYPHENIDYL HCL	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS DRUG IS LABELED AS HIGH RISK MEDICATION IN THE ELDERLY FOR PATIENTS 65 YEARS AND OLDER.
HIGH RISK DRUGS IN THE ELDERLY - ANTICHOLINERGICS - HYDROXYZINE	HYDROXYZINE HCL   HYDROXYZINE PAMOATE	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	"PRURITUS/URTICARIA/SEASONAL/PERENNIAL ALLERGY: TRIAL OR CONTRAINDICATION TO A NON-SEDATING ANTIHISTAMINE SUCH AS LEVOCETIRIZINE. ANXIETY: TRIAL OR CONTRAINDICATION TO TWO (2) OF THE FOLLOWING - BUSPIRONE, PAROXETINE, DULOXETINE, OR VENLAFAXINE."

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								Your Care Comes First
Prior Authorization Group Description	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
HIGH RISK DRUGS IN THE ELDERLY - CARDIOVASCULAR	GUANFACINE HCL	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER, PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	"HYPERTENSION: TRIAL OR CONTRAINDICATION TO TWO (2) OF THE FOLLOWING - BENAZEPRIL, BENAZEPRIL/HYDROCHLOROTHIAZIDE, CAPTOPRIL, CAPTOPRIL/HYDROCHLOROTHIAZIDE, ENALAPRIL, ENALAPRIL/HYDROCHLOROTHIAZIDE, ENALAPRIL, ENALAPRIL/HYDROCHLOROTHIAZIDE, FOSINOPRIL, LISINOPRIL/HYDROCHLOROTHIAZIDE, LISINOPRIL, HYDROCHLOROTHIAZIDE, RAMIPRIL, MOEXIPRIL, MO
HIGH RISK DRUGS IN THE ELDERLY - CENTRAL NERVOUS SYSTEM - THIORIDAZINE	THIORIDAZINE HCL	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	SOTALOL, TIMOLOL MALEATE." 65 YEARS AND OLDER: SCHIZOPHRENIA - PRESCRIBER ACKNOWLEDGEMENT/AWARENESS DRUG IS LABELED AS HIGH RISK MEDICATION IN THE ELDERLY FOR PATIENTS 65 YEARS AND OLDER.
	DIGOXIN	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		DIGOXIN LEVEL	APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	APPROVAL FOR MEMBERS STABLE ON 250 MCG WITH DOCUMENTED THERAPEUTIC DIGOXIN LEVEL TAKEN WITHIN THE PAST YEAR.
HIGH RISK DRUGS IN THE ELDERLY - ENDOCRINE - GLYBURIDE	GLYBURIDE   GLYBURIDE MICRONIZED   GLYBURIDE-METFORMIN HCL	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS DRUG IS LABELED AS HIGH RISK MEDICATION IN THE ELDERLY FOR PATIENTS 65 YEARS AND OLDER.
HIGH RISK DRUGS IN THE ELDERLY - NON- BENZODIAZEPINE	ZALEPLON   ZOLPIDEM TARTRATE   ZOLPIDEM TARTRATE ER	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER, PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	MEMBERS 65 YEARS OR OLDER WILL BE EVALUATED FOR LIMITED PRESCRIPTION USE TO NO MORE THAN 90 DAYS (TOTAL) OF CUMULATIVE USE WITHIN THE CURRENT PLAN YEAR. REQUESTS GREATER THAN 90 DAYS OF CUMULATIVE USE REQUIRES PRESCRIBER ACKNOWLEDGEMENT/AWARENESS DRUG IS LABELED AS HIGH RISK MEDICATION IN THE ELDERLY FOR PATIENTS 65 YEARS AND OLDER.
HIGH RISK DRUGS IN THE ELDERLY - SKELETAL MUSCLE RELAXANTS	CARISOPRODOL   CHLORZOXAZONE   CYCLOBENZAPRINE HCL   METHOCARBAMOL	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.			APPROPRIATE HIGH RISK MEDICATION USE: 65 YEARS OF AGE OR OLDER. PA NOT REQUIRED FOR AGE 0- 64 YEARS.		12 MONTHS	PRESCRIBER ACKNOWLEDGEMENT/AWARENESS DRUG IS LABELED AS HIGH RISK MEDICATION IN THE ELDERLY FOR PATIENTS 65 YEARS AND OLDER.
IMATINIB MESYLATE	GLEEVEC	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					ALL DIAGNOSIS: 12 MONTHS. GIST (TWICE DAILY DOSE): 36 MONTHS.	"GASTROINTESTINAL STROMAL TUMOR (GIST) KIT (CDI17) POSITIVE USE FOR GLEEVEC 400MG TWICE DAILY: TRIAL OF GLEEVEC 400MG ONCE DAILY OR GIST TUMOR EXPRESSING A KIT EXON 9 MUTATION. PREVIOUSLY TREATED CML REQUIRES MUTATIONAL ANALYSIS NEGATIVE FOR THE FOLLOWING MUTATIONS FOLLOWING BCR-ABL MUTATIONAL ANALYSIS - T315I, V299L, F317L/V/I/C, Y253H, E255K/V, F359V/C/I."

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								Your Care Comes First
Prior Authorization Group Description	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
IMIQUIMOD - ALDARA	IMIQUIMOD	"ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ADDITIONAL OFF LABEL COVERAGE FOR ACTINIC KERATOSIS NOT LIMITED TO THE FACE AND SCALP IN NON-IMMUNOCOMPETENT PATIENTS, MOLLUSCUM CONTAGIOSUM, AND LETIGO MALIGNA."			EXTERNAL GENITAL OR PERIANAL WARTS: GREATER THAN OR EQUAL TO 12 YEARS OF AGE. ACTINIC KERATOSIS: GREATER THAN OR EQUAL TO 18 YEARS OF AGE.	ACTINIC KERATOSIS: DERMATOLOGIST ONLY. SUPERFICIAL BASAL CELL CARCINOMA/LETIGO MALIGNA: DERMATOLOGIST OR ONCOLOGIST ONLY.	4 MONTHS	CRITERIA APPLIES TO NEW STARTS ONLY. ACTINIC KERATOSIS: TRIAL OF TOPICAL 5- FLUOROURACIL. ACTINIC KERATOSIS BRAND DRUG REQUEST: TRIAL/FAILURE OF GENERIC IMIQUIMOD 5%. SUPERFICIAL BASAL CELL CARCINOMA: LESS THAN 2CM IN SIZE AND NOT ON THE FACE. MOLLUSCUM CONTAGISOUM LIMITED TO THE FACE.
IMMUNE GLOBULIN BVD DETERMINATION	CARIMUNE NF NANOFILTERED   GAMASTAN S-D   GAMMAGARD LIQUID   GAMMAPLEX   GAMUNEX- C   PRIVIGEN	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
IMMUNOSUPPRESSANT BVD DETERMINATION	AZATHIOPRINE   AZATHIOPRINE SODIUM   CELLCEPT   CYCLOSPORINE   CYCLOSPORINE MODIFIED   GENGRAF   MYCOPHENOLATE MOFETIL   MYFORTIC   NULOJIX   PROGRAF   RAPAMUNE   SIMULECT   TACROLIMUS   ZORTRESS	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
INFLIXIMAB	REMICADE	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		"INITIAL: PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 10 PERCENT BODY SURFACE AREA OR PSORIATIC LESIONS AFFECT THE HANDS, FEET, OR GENITAL AREA. RENEWAL: RIEUMATOID/PSORIATIC ARTHRITIS: MAINTAINED OR EXPERIENCED GREATER THAN 20% IN TENDER JOINT COUNT AND SWOLLEN JOINT COUNT. PLAQUE PSORIASIS: MAINTAINED OR EXPERIENCED PASI OF GREATER THAN 50% OR SIGNIFCANT IMPROVEMENT IN QUALITY OF LIFE OBSERVED BY PHYSICIAN AND PATIENT. ANKYLOSING SPONDYLITIS: MAINTAINED OR EXPERIENCED IMPROVEMENT OF AT LEAST 50%, OR 2 UNITS (SCALE OF 1-10), IN THE BATH ANKYLOSING SPONDYLITIS DISEASE ACTIVITY INDEX (BASDAI) OR IMPROVEMENT OF AT LEAST 20% IN THE ASSESSMENT IN ANKYLOSING SPONDYLITIS (SPONDYLITIS DISEASE ACTIVITY INDEX (BASDAI) OR IMPROVEMENT OF AT LEAST 20% IN THE ASSESSMENT IN ANKYLOSING SPONDYLITIS (ASAS20) CRITERIA."			UC: 12 MO. OTHER INDICATIONS INITIAL: 4 MO RENEWAL: 12 MO	"INITIAL: MODERATE TO SEVERE CROHN'S DISEASE/ULCERATIVE COLITIS/ACUTE ENTEROCUTANEOUS FISTULA: TRIALFAILURE OF ONE OR MORE OF THE FOLLOWING PREFERRED THERAPY AGENTS SUCH AS SULFASALAZINE, CORTICOSTEROIDS, AZATHIOPRINE, METHOTREXATE, OLSALAZINE, MESALAMINE, CYCLOSPORINE, OR MERCAPTOPURINE. FOR MODERATE TO SEVERE RHEUMATOID ARTHRITIS: TRIAL OF HUMIRA OR CIMZIA AND TRIAL/FAILURE TO AT LEAST ONE DMARD AGENT (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE). FOR PSORIATIC ARTHRITIS: TRIAL OF HUMIRA AND TRIAL/FAILURE TO AT LEAST ONE DMARD AGENT (METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE.) FOR SEVERE PLAQUE PSORIASIS COVERING 10% BSA: TRIAL/FAILURE/INTOLERABLE SIDE AFFECTS TO AT LEAST ONE PREFERRED THERAPY (PUVA, UVB, ACITRETIN, METHOTREXATE OR CYCLOSPORINE), RENEWAL: FOR RHEUMATOID ARTHRITIS: CONCOMITANT METHOTREXATE USE."
INFUSIBLE DRUG BVD DETERMINATION	ABELCET   ACYCLOVIR SODIUM   ADRIAMYCIN   AMPHOTERICIN B   BLEOMYCIN SULFATE   CLADRIBINE   CYTARABINE   FLUOROURACIL   FOSCARNET SODIUM   GANCICLOVIR SODIUM   IFOSFAMIDE   METHOTREXATE   MITOMYCIN   REMODULIN   TORISEL   VINBLASTINE SULFATE   VINCRISTINE SULFATE	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
INTERFERON AGENTS - INTERFERON ALFA-2B	INTRON A	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		HEP C: PRETREATMENT HCV RNA LEVEL GREATER THAN OR EQUAL TO 50 IU/ML.	HEP C: 3 YEARS OR OLDER.	"HEP C: GASTROENTEROLOGIS T, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (E.G. HEPATOLOGIST)."	MOS. RENEWAL HEP C	HEP C: DRUG MUST BE USED IN COMBINATION WITH RIBAVIRIN UNLESS CONTRAINDICATED.
INTERFERON AGENTS - PEG-INTERFERON ALFA 2A	PEGASYS   PEGASYS PROCLICK	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		"INITIAL: HEP C: PRETREATMENT HCV RNA LEVEL GREATER THAN OR EQUAL TO 50 IU/ML. HEP C WITH HIV: CD4 COUNT GREATER THAN 100 CELLS/MM3, HCV RNA LEVELS/VIRAL LOAD GREATER THAN OR EQUAL TO 50 IU/ML. RENEWAL: HCV RNA LEVELS TO DETERMINE LENGTH OF TREATMENT."	5 YEARS OR OLDER.		HEP C 2 TO 6 MOS.	HEP C: TRIAL OR CONTRAINDICATION TO PEGINTRON, DRUG MUST BE USED IN COMBINATION WITH RIBAVIRIN UNLESS CONTRAINDICATED, RENEWAL: HEP C: USED IN COMBINATION OR CONTRAINDICATION WITH RIBAVIRIN, GENOTYPE 2 OR 3: NO RENEWAL.

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Prior Authorization Group Description	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
INTERFERON AGENTS - PEG-INTERFERON ALFA 2B		ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		INITIAL: HEP C: PRETREATMENT HCV RNA LEVEL GREATER THAN OR EQUAL TO 50 IU/ML. RENEWAL: HCV RNA LEVELS TO DETERMINE LENGTH OF TREATMENT.	3 YEARS OR OLDER.	"GASTROENTEROLOGIS T, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (E.G. HEPATOLOGIST)."		HEP C: DRUG MUST BE USED IN COMBINATION WITH RIBAVIRIN UNLESS CONTRAINDICATED. RENEWAL HEP C: USED IN COMBINATION OR CONTRAINDICATION WITH RIBAVIRIN. GENOTYPE 2 OR 3: NO RENEWAL.
IPILIMUMAB	YERVOY	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					3 MONTHS	
IVACAFTOR	KALYDECO	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		G551D MUTATION	6 YEARS OF AGE OR OLDER.		12 MONTHS	
LENALIDOMIDE	REVLIMID	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	
METHOTREXATE BVD DETERMINATION	METHOTREXATE   TREXALL	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES, INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
METHYLNALTREXONE	RELISTOR	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		CONSTIPATION DUE TO OPIOIDS			UP TO 6 MONTHS	PATIENT IS RECEIVING PALLIATIVE CARE.
MIFEPRISTONE	KORLYM	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	
MIPOMERSEN	KYNAMRO	OTHERWISE EXCLUDED FROM PART D. ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	PATIENT IS CONCURRENTLY RECEIVING LDL APHERESIS.				12 MONTHS	"USE IN COMBINATION WITH A STATIN (EXAMPLE: SIMVASTATIN, ATORVASTATIN), BILE ACID SEQUESTRANT FENOFIBRATE OR NIACIN."
MODAFINIL AND ARMODAFINIL - PROVIGIL	MODAFINIL	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ADDITIONAL COVERAGE CONSIDERATION FOR CHRONIC FATIGUE SYNDROME RELATED TO MULTIPLE SCLEROSIS.					12 MONTHS	"NARCOLEPSY: TRIAL OF OR CONTRAINDICATION TO AMPHETAMINE, DEXTROAMPHETAMINE, OR METHYLPHENDATE."
NATALIZUMAB	TYSABRI	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					MULTIPLE SCLEROSIS:12 MONTHS CROHN'S DISEASE: 6 MONTHS. RENEWAL: CROHN'S: 12 MONTHS.	"MULTIPLE SCLEROSIS: TRIAL OF AN INTERFERON OR COPAXONE. CROHN'S DISEASE: TRIAL OF A TNI ALPHA INHIBITOR. RENEWAL: CROHN'S: PATIENT IS NOT ON CONCOMITANT CORTICOSTEROID TREATMENT AFTER 6 MONTHS ON NATALIZUMAB OR HAS NOT RECEIVED MORE THAN 3 MONTHS OI A CORTICOSTEROID WITHIN THE PAST 12 MONTHS."
NEBULIZER BVD DETERMINATION	ACETYLCYSTEINE   ALBUTEROL SULFATE   CROMOLYN SODIUM   NEBUPENT   PULMOZYME   TOBI   VENTAVIS	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
OFATUMUMAB	ARZERRA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	CHRONIC LYMPHOCYTIC LEUKEMIA: NO FAILED TREATMENT WITH FLUDARABINE AND ALEMTUZUMAB				6 MONTHS	
OMACETAXINE	SYNRIBO	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					INDUCTION: 3 MONTHS POST INDUCTION/RENEWAL: 3 TO 12 MONTHS	. "CML INDUCTION THERAPY: TRIAL OF OR CONTRAINDICATION TO AT LEAST TWO OF THE FOLLOWING GLEEVEC, SPRYCEL, TASIGNA, BOSULIF, OR ICLUSIG. DETERMINATION FOR THERAPY LENGTH OF APPROVAL THAT IS NOT INDUCTION THERAPY WILL DEPEND ON THE PATIENTS HEMATOLOGIC RESPONSE (DEFINED AS ABSOLUTE NEUTROPHIL COUNT (ANC) GREATER THAN OR EQUAL TO 1.5 X 10°9/L AND PLATELETS GREATER THAN OR EQUAL TO 100 X 10°9/L AND NO BLOOD BLASTS OR BONE MARROW BLASTS LESS THAN 5%). IF MEETS HEMATOLOGIC RESPONSE CRITERIA APPROVAL WILL BE 12 MONTHS. IF HEMATOLOGIC RESPONSE CRITERIA IS NOT MET APPROVAL WILL BE FOR 3 MONTHS."

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Drug Name BUPRENORPHINE-NALOXONE   XOLAIR	Covered Uses  ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	Exclusion Criteria	Required Medical Information "INITIAL. PATIENT MEETS THE CRITERIA OF	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
BUPRENORPHINE-NALOXONE   XOLAIR			"INITIAL. PATIENT MEETS THE CRITERIA OF	PATIENT 12 YEARS OF	SPECIALIST IN	12 MONTHS	
			MODERATE TO SEVERE ASTHMA, POSITIVE SKIN PRICK OR RAST TEST, FEVI LESS THAN 80%, DEMONSTRATED INADEQUATELY CONTROLLED	AGE OR OLDER	ALLERGY OR PULMONARY MEDICINE ONLY	12 MONTHO	
			SYMPTOMS ON INHALED CORTICOSTEROIDS AND SECOND ASTHMA CONTROLLER, BASELINE IGE SERUM LEVEL GREATER THAN OR EQUAL TO 30 IU/ML. RENEWAL: PATIENT REDUCED EXACERBATIONS BY AT LEAST 25% FROM BASELINE, REDUCTION IN ORAL OR INHALED CORTICOSTEROID USE FROM BASELINE."				
VECTIBIX	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	
VOTRIENT	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	
ADCIRCA   PEVATIO   SILDENAFII					CARDIOLOGIST OR	12 MONTHS	PEOLIEST FOR A DOING A PROLIBE TRIAL OR
ADCINCA   REVATIO   SILDENATIE	OTHERWISE EXCLUDED FROM PART D.				PULMONOLOGIST	12 WONTHS	CONTRAINDICATION TO REVATIO.
SYLATRON 4-PACK	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	CRITERIA APPLIES TO NEW STARTS ONLY. DURATION LIMITATION OF 5 YEARS OF THERAPY.
PERJETA	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.						
<del>MOZOBIL</del>	OTHERWISE EXCLUDED FROM PART D.		IOSE IN COMBINATION WITH GRANULOCYTE- COLONY STIMULATING FACTOR (G-CSF) TO MOBILIZE HEMATOPOIETIC STEM CELLS TO THE PERIPHERAL BLOOD FOR COLLECTION AND SUBSEQUENT AUTOLOGOUS TRANSPLANTATION IN PATIENTS WITH NON-HODGKIN'S LYMPHOMA AND MULTIPLE MYELOMA				
POMALYST	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	
ICLUSIG	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	
SYMLINPEN 120   SYMLINPEN 60	OTHERWISE EXCLUDED FROM PART D. ALL FDA APPROVED INDICATIONS NOT		TYPE FOR TYPE II DIABETES. REQUIRING INSULIN			12 MONTHS	
	OTHERWISE EXCLUDED FROM PART D.		OR CONTINUOUS INSULIN INFUSION (INSULIN PUMP) FOR GLYCEMIC CONTROL				
QUININE SULFATE	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	
IMOVAX RABIES VACCINE   RABAVERT	PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
STIVARGA	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D					12 MONTHS	"TRIAL OR CONTRAINDICATION TO ANTI-EGFR THERAPY SUCH AS ERBITUX OR VECTIBIX. TRIAL
							OR CONTRAINDICATION TO ANTI-VEGF THERAPY SUCH AS AVASTIN OR ZALTRAP AND A FLUOROPYRMIDE-, OXAPLATIN- AND IRINOTECAN BASED CHEMOTHERAPY SUCH AS FOLFOX, FOLFIRI, CAPEOX, INFUSIONAL 5-FU/LV OR CAPECITABINE, AND FOLFOXIRI."
XIFAXAN	OTHERWISE EXCLUDED FROM PART D.			DIARRHEA: 12 YEARS OR OLDER. HEPATIC ENCEPHALOPATHY: 18 YEARS OR OLDER.		MONTH. HEPATIC ENCEPHALOPATHY: 12	TRAVELERS' DIARRHEA. TRIAL OF CIPROFLOXACIN OR AZITHROMYCIN. HEPATIC ENCEPHALOPATHY: TRIAL OF LACTULOSE MONOTHERAPY.
RITUXAN	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		RENEWAL. ACTIVE RHEUMATOID ARTHRITIS/PSORIATIC ARTHRITIS: GREATER THAN 20% IMPROVEMENT IN TENDER JOINT COUNT AND SWOLLEN JOINT COUNT.		RHEUMATOID ARTHRITIS A RHEUMATOLOGIST. FOR NHL OR CLL AN	YEAR. CLL: 6 MO. WG,	TINITIAL. RHEUMATOID ARTHRITIS. CURRENTLY TAKING OR HAVE A CONTRAINDICATION TO THE USE OF METHOTREXATE AND TRIALIFAILURE OF ONE TNF BLOCKER (ENBREL, HUMIRA, SIMPONI, CIMZIA). NON HODGKIN'S LYMPHOMA/CHRONIC LYMPHOCYTIC LEUKEMIA: USED IN COMBINATION WITH CHEMOTHERAPY. WEGNER'S GRANULOMATOSIS/MICROSCOPIC POLYANGIITIS: CONCURRENT GLUCOCORTICOID USE."
The second secon	ADCIRCA   REVATIO   SILDENAFIL  SYLATRON 4-PACK  PERJETA  AOZOBIL  OMALYST  CLUSIG  SYMLINPEN 120   SYMLINPEN 60  WINNE SULFATE  MOVAX RABIES VACCINE   RABAVERT	OTHERWISE EXCLUDED FROM PART D ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.  ALL FROM APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	DESILING REDUCTION NO GAL OR RINHALED CORTICOSTEROID USE FROM BASELINE."  ALL MEDICALLY ACCEPTED ROBATION NOT OTHERWISE EXCLLIDED FROM PART D.  OTHERWISE EXCLLIDED FROM PART D.  ALL MEDICALLY ACCEPTED ROBATION NOT OTHERWISE EXCLLIDED FROM PART D.  ALL MEDICALLY ACCEPTED ROBATION NOT OTHERWISE EXCLLIDED FROM PART D.  ALL FOR AFFROVED ROBATION NOT OTHERWISE EXCLIDED FROM PART D.  ALL FOR AFFROVED ROBATION NOT OTHERWISE EXCLIDED FROM PART D.  ALL FOR AFFROVED ROBATION NOT OTHERWISE EXCLIDED FROM PART D.  ALL FOR AFFROVED ROBATION NOT OTHERWISE EXCLIDED FROM PART D.  ALL FOR AFFROVED ROBATION NOT OTHERWISE EXCLIDED FROM PART D.  ALL FOR AFFROM PART D.  ALL FOR AFFROM PART D.  ALL FOR AFFROM PART D	RETHINK  ALL MEDICALLY ACCEPTED ROBICATIONS NOT  OTHERWISE EXCLUDED PROMPARE D  OTHERWISE EXC	DASSELNE, SELECTION POR GAL OR STRIALED CONTROL STRONG USE FROM BASELINE.*  OTHERWISE EXCLUDED FROM PART D.  OTHERWISE EXCLUDED FROM	RASSING REPORTED NOR OR SHALED.  17 MOTOR AT ACCOPTION NOT CONTROL SERVINGS FROM ASSELAN.*  18 MOTOR ACCOUNTS FROM ASSELAN.*  18 MOTOR ACCOUNTS FROM ASSELAN.*  19 MOTOR ACCOUNTS FROM ASSELANCE ACC

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Prior Authorization	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
Group Description COMIDEPSIN	ISTODAX	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	TRIAL OF OR CONTRAINDICATION TO VORINOSTAT (ZOLINZA) AND NOT ABLE TO TOLERATE ORAL MEDICATIONS, OR IS ABLE TO TOLERATE ORAL MEDICATIONS AND HAS TRIED AT LEAST ONE SYSTEMIC THERAPY (RETINOID, INTERFERON, EXTRACORPOREAL PHOTOPHERESIS, DENILEUKIN DIFTITOX, METHOTREXATE, LIPOSOMAL DOXORUBICIN, GEMCITABINE, CHLORAMBUCIL, PENTOSTATIN, ETOPOSIDE, CYCLOPHOSPHAMIDE, TEMOZOLOMIDE, BORTEZOMIB)."
RUXOLITINIB	JAKAFI	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		"RENEWAL: IMPROVEMENT OR MAINTENANCE OF SYMPTOM IMPROVEMENT SUCH AS A 50% OR GREATER REDUCTION IN TOTAL SYMPTOM SCORE ON THE MODIFIED MYELOFIBROSIS SYMPTOM ASSESSMENT FORM (MFSAF) V2.0 OR 50% OR GREATER REDUCTION IN PALPABLE SPLEEN LENGTH, OR REDUCTION OF 35% OR GREATER FROM BASELINE SPLEEN VOLUME AFTER 6 MONTHS OF THERAPY."			INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS	
SOMATROPIN - SEROSTIM	SEROSTIM	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	"ATHLETIC ENHANCEMENT, ANTI-AGING PURPOSES"	"HIV/WASTING: MEETS CRITERA OF WEIGHT LOSS: 10% UNINTENTIONAL WEIGHT LOSS OVER 12 MONTHS, OR 7.5% OVER 6 MONTHS, OR 5% BODY CELL MASS (BCM) LOSS WITHIN 6 MONTHS, OR A BCM LESS THAN 35% (MEN), 23% (WOMEN) OF TOTAL BODY WEIGHT AND A BODY MASS INDEX (BMI) LESS THAN 27 KG PER METER SQUARED, OR BMI LESS THAN 20 KG PER METER SQUARED."			HIV/AIDS: 3 MONTHS.	"HIV/WASTING: CURRENTLY ON ANTIRETROVIRATHERAPY. IF CURRENTLY ON GROWTH HORMON PATIENT HAS SHOWN CLINICAL BENEFIT IN MUSCLE MASS AND WEIGHT OR IF NOT ON GROWTH HORMONE, PATIENT HAS HAD INADEQUATE RESPONSE TO PREVIOUS THERAPY (I.E. EXERCISE TRAINING, NUTRITIONAL SUPPLEMENTS, APPETITE STIMULANTS OR ANABOLIC STEROIDS)."
SOMATROPIN - ZORBTIVE	ZORBTIVE	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	"ATHLETIC ENHANCEMENT, ANTI-AGING PURPOSES"	SHORT-BOWEL SYNDROME: CURRENTLY ON SPECIALIZED NUTRITIONAL SUPPORT.			SHORT BOWEL: 4 WEEK ONCE.	
SORAFENIB TOSYLATE	NEXAVAR	ALL MEDICALLY ACCEPTED INDICATIONS NOT	TON OSES	SI LEMELLED NOTHING ME SOTTOKT.			12 MONTHS	
SUNITINIB MALATE	SUTENT	OTHERWISE EXCLUDED FROM PART D. ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	GASTROINTESTIONAL STROMAL TUMORS (GIST) TRIAL OF OR CONTRAINDICATION TO GLEEVEC.
TELAPREVIR	INCIVEK	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	COMPLETED PRIOR COURSE OF THERAPY WITH TELAPREVIR (INCIVEK) OR BOCEPREVIR (VICTRELIS) AND DID NOT ACHIEVE A SUSTAINED VIROLOGIC RESPONSE. CURRENTLY TAKING RIFAMPIN OR HAS A CO-INFECTION WITH HEPATITIS B.	"CHRONIC HEPATITIS C, GENOTYPE 1. HCV RNA LEVEL/VIRAL LOAD OF LESS THAN 1,000 IU/ML AT 4 WEEKS OF TELAPREVIR THERAPY."	PATIENT 18 YEARS OF AGE OR OLDER.	"GASTROENTEROLOGIS T, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (E.G. HEPATOLOGIST) OR SPECIALLY TRAINED GROUP (E.G. EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES). HEP C AND ORGAN TRANSPLANT: TRANSPLANT CENTER AND TRANSPLANT PHYSICIAN."		HEP C: CONCURRENT USE OF RIBAVIRIN AND PEGINTERFERON ALFA.
TERIFLUNOMIDE	AUBAGIO	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	"TRIAL OF OR CONTRAINDICATION TO ONE INTERFERON THERAPY (SUCH AS AVONEX, BETASERON, EXTAVIA, OR REBIF) AND TO COPAXONE."
TERIPARATIDE	FORTEO	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	GREATER THAN 24 MONTHS OF THERAPY.	"A PATIENT WITH EITHER A DIAGNOSIS OF SEVERI OSTEOPOROSIS (T-SCORE LESS THAN -2-5 WITH FRAGILITY FRACTURE) OR A T SCORE EQUAL TO OR LESS THAN -2-5 AND MULTIPLE RISK FACTORS FOR FRACTURE (E.G. HISTORY OF MULTIPLE RECENT LOW TRAUMA FRACTURES, CORTICOSTEROID USE, OR USE OF GNRH ANALOGS), OR FAILED AN ADEQUATE TRIAL OF BISPHOSPHONATES, IS INTOLERANT, OR HAS A CONTRAINDICATION TO BISPHOSPHONATES."			12 MONTHS	

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Prior Authorization Group Description	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
TESTOSTERONE	ANDRODERM   ANDROGEL   AXIRON   TESTOSTERONE CYPIONATE   TESTOSTERONE ENANTHATE	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		"MALE HYPOGONADISM CONFIRMED BY EITHER: 1) LAB CONFIRMED TOTAL SERUM TESTOSTERONE LEVEL OF LESS THAN 300 NG/DL OR 2) A LOW TOTAL SERUM TESTOSTERONE LEVEL AS INDICATED BY A LAB RESULT WITH A REFERENCE RANGE OBTAINED WITHIN 90 DAYS, OR 3) A FREE SERUM TESTOSTERONE LEVEL OF LESS THAN 50 NG/L."			LIFETIME OF MEMBERSHIP IN PLAN	
TETANUS TOXOID VACCINE BVD DETERMINATION	TETANUS TOXOID ADSORBED	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
TETRABENAZINE	XENAZINE	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.				NEUROLOGIST	12 MONTHS	
THALIDOMIDE	THALOMID	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ADDITIONAL COVERAGE CONSIDERATION FOR ANEMIA DUE TO MYELODYSPLASTIC SYNDROME AND WALDENSTROM'S MACROGLOBULINEMIA.					12 MONTHS	
THIAZOLIDINEDIONE	AVANDAMET   AVANDARYL   AVANDIA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	"APPLIES TO NEW STARTS ONLY. TRIAL OR CONTRAINDICATION TO METFORMIN, METFORMIN ER, GLYBURIDE/METFORMIN, GLIPIZIDE/METFORMIN OR A SULFONYLUREA AND PIOGLITAZONE."
TOFACITINIB	XELJANZ	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		RENEWAL: RHEUMATOID ARTHRITIS: EXPERIENCED OR MAINTAINED 20 PERCENT IMPROVEMENT IN TENDER OR SWOLLEN JOINT COUNT WHILE ON THERAPY.		RHEUMATOLOGIST		RHEUMATOID ARTHRITIS INITIAL: TRIAL OR CONTRAINDICATION TO HUMIRA AND CIMZIA.
TOPICAL TRETINOIN	AVITA   TRETINOIN	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.	"WRINKLES, PHOTOAGING, MELASMA."				12 MONTHS	BRAND TRETINON WILL REQUIRE TRIAL OF GENERIC TOPICAL TRETINOIN.
TOTAL PARENTARAL NUTRITION AGENT BVD DETERMINATION	AMINOSYN   AMINOSYN II   AMINOSYN M   AMINOSYN-HBC   AMINOSYN-PF   CLINIMIX   CLINIMIX E   CLINISOL   DEXTROSE IN WATER   FREAMINE III   HEPATAMINE   HEPATASOL   INTRALIPID   LIPOSYN III   NEPHRAMINE   PREMASOL   PROCALAMINE   PROSOL   TRAVASOL   TROPHAMINE	THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION.						
TRAMETINIB DIMETHYL SULFOXIDE	MEKINIST	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	
TRASTUZUMAB	HERCEPTIN	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		"BREAST CANCER, METASTATIC BREAST CANCER, GASTRIC CANCER: HER2 POSITIVE"			12 MONTHS	B VS D COVERAGE CONSIDERATION. BREAST CANCER: USED IN COMBINATION WITH CHEMOTHERAPY (EXAMPLES INCLUDE: DOXORUBICIN AND CYCLOPHOSPHAMIDE FOLLOWED BY PACLITAXEL OR DOCETAXEL AND CARBOPLATIN OR DOCETAXEL FOLLOWED BY FLUOROURICIL/EPIRUBICIN/CYCLOPHOSPHAMIDE OR DOXORUBICIN/CYCLOPHOSPHAMIDE FOLLOWED BY DOCETAXEL OR PACLITAXEL. GASTRIC CANCER: USED IN COMBINATION WITH CHEMOTHERAPY (EXAMPLES INCLUDE: CISPLATIN AND FLUOROPYRIMIDINE.
USTEKINUMAB	STELARA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		INITIAL: PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 10 PERCENT BODY SURFACE AREA OR PASI SCORE GREATER THAN OR EQUAL TO 12. PATIENT'S WEIGHT.		DERMATOLOGIST OR RHEUMATOLOGIST	INITIAL: 4 MONTHS. RENEWAL: 12 MONTHS	"FOR SEVERE PLAQUE PSORIASIS COVERING 10% BSA: TRIAL/FAILURE/INTOLERABLE SIDE AFFECTS TO AT LEAST ONE PREFERRED THERAPY (PUVA, UVB, ACITRETIN, METHOTIEXATE OR CYCLOSPORINE). RENEWAL: PHYSICIAN'S GLOBAL ASSESMENT EQUAL TO ZERO OR ONE OR A DECREASE OF PASI OF AT LEAST 50% OR GREATER."
VANDETANIB	CAPRELSA	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	CRITERIA APPLIES TO NEW STARTS ONLY.
VEMURAFENIB	ZELBORAF	ALL MEDICALLY ACCEPTED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.		BRAFV600E MUTATION			12 MONTHS	
VILAZODONE	VIIBRYD	ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D.					12 MONTHS	"TRIAL OF OR CONTRAINDICATION TO A SSRI (PAROXETINE, SERTARLINE, CITALOPRAM, FLUOXETINE, OR ESCITALOPRAM) AND A SECOND AGENT (BUPROPION HCL (IR, SR, OR XL), MIRTAZAPINE, OR VENLAFAXINE (IR OR XR))."



Prior Authorization	Drug Name	Covered Uses	Exclusion Criteria	Required Medical Information	Age Restrictions	Prescriber Restrictions	Coverage Duration	Other Criteria
Group Description								
VISMODEGIB	ERIVEDGE	ALL MEDICALLY ACCEPTED INDICATIONS NOT					12 MONTHS	
		OTHERWISE EXCLUDED FROM PART D.						

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