COMMERCIAL PHARMACY PROGRAM POLICY ACTIVITY

Provider Notification



Policies Effective: August 15, 2024 Notification Posted: July 1, 2024

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NEW POLICIES DEVELOPED

No new policies for August 15, 2024

POLICIES REVISED

◆ Program Summary: Biologic Immunomodulators Applies to: ☐ Commercial Formularies Type: ☐ Prior Authorization ☐ Quantity Limit ☐ Step Therapy ☐ Coverage / Formulary Exception

POLICY AGENT SUMMARY QUANTITY LIMIT

Wildcard	Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	QL Amount	Dose Form	Days Supply	Duration	Targeted NDCs When Exclusions Exist	Age Limit	Effective Date	Term Date
6627001507F810	Abrilada	adalimumab-afzb prefilled syringe kit	20 MG/0.4ML	2	Syringes	28	DAYS				
6627001507F820	Abrilada	adalimumab-afzb prefilled syringe kit	40 MG/0.8ML	2	Syringes	28	DAYS				
6627001507F520	Abrilada 1-pen kit ; Abrilada 2- pen kit	adalimumab-afzb auto- injector kit	40 MG/0.8ML	2	Pens	28	DAYS				
6650007000E5	Actemra	tocilizumab subcutaneous soln prefilled syringe	162 MG/0.9ML	4	Syringes	28	DAYS				
6650007000D5	Actemra actpen	tocilizumab subcutaneous soln auto- injector	162 MG/0.9ML	4	Pens	28	DAYS				
6627001510D517	Amjevita	adalimumab-atto soln auto-injector	40 MG/0.4ML	2	Pens	28	DAYS				
6627001510D520	Amjevita	adalimumab-atto soln auto-injector	40 MG/0.8ML	2	Pens	28	DAYS				
6627001510D537	Amjevita	adalimumab-atto soln auto-injector	80 MG/0.8ML	2	Pens	28	DAYS				
6627001510E505	Amjevita	adalimumab-atto soln prefilled syringe	10 MG/0.2ML	2	Syringes	28	DAYS				
6627001510E508	Amjevita	adalimumab-atto soln prefilled syringe	20 MG/0.2ML	2	Syringes	28	DAYS				
6627001510E510	Amjevita	adalimumab-atto soln prefilled syringe	20 MG/0.4ML	2	Syringes	28	DAYS				

Wildcard	Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	QL Amount	Dose Form	Days Supply	Duration	Targeted NDCs When Exclusions Exist	Age Limit	Effective Date	Term Date
6627001510E517	Amjevita	adalimumab-atto soln prefilled syringe	40 MG/0.4ML	2	Syringes	28	DAYS				
6627001510E520	Amjevita	adalimumab-atto soln prefilled syringe	40 MG/0.8ML	2	Syringes	28	DAYS				
9025051800D520	Bimzelx	bimekizumab-bkzx subcutaneous soln auto- injector	160 MG/ML	2	Pens	56	DAYS				
9025051800E520	Bimzelx	bimekizumab-bkzx subcutaneous soln prefilled syr	160 MG/ML	2	Syringes	56	DAYS				
525050201064	Cimzia	certolizumab pegol for inj kit	200 MG	2	Kits	28	DAYS				
5250502010F840	Cimzia	Certolizumab Pegol Prefilled Syringe Kit	200 MG/ML	2	Kits	28	DAYS				
5250502010F860	Cimzia starter kit	Certolizumab Pegol Prefilled Syringe Kit	200 MG/ML	1	Kit	180	DAYS				
9025057500E530	Cosentyx	Secukinumab Subcutaneous Pref Syr 150 MG/ML (300 MG Dose)	150 MG/ML	2	Syringes	28	DAYS				
9025057500E510	Cosentyx	Secukinumab Subcutaneous Soln Prefilled Syringe	75 MG/0.5ML	1	Syringe	28	DAYS				
9025057500E520	Cosentyx	Secukinumab Subcutaneous Soln Prefilled Syringe 150 MG/ML	150 MG/ML	1	Syringe	28	DAYS				
9025057500D530	Cosentyx sensoready pen	Secukinumab Subcutaneous Auto-inj 150 MG/ML (300 MG Dose)	150 MG/ML	2	Pens	28	DAYS				
9025057500D520	Cosentyx sensoready pen	Secukinumab Subcutaneous Soln Auto-injector 150 MG/ML	150 MG/ML	1	Pen	28	DAYS				
9025057500D550	Cosentyx unoready	secukinumab subcutaneous soln auto- injector	300 MG/2ML	1	Pen	28	DAYS				
6627001505F520	Cyltezo	adalimumab-adbm auto-injector kit	40 MG/0.8ML	2	Pens	28	DAYS	00597037597; 00597054522; 82009014822			
6627001505F805	Cyltezo	adalimumab-adbm prefilled syringe kit	10 MG/0.2ML	2	Syringes	28	DAYS				
6627001505F810	Cyltezo	adalimumab-adbm prefilled syringe kit	20 MG/0.4ML	2	Syringes	28	DAYS				
6627001505F820	Cyltezo	adalimumab-adbm prefilled syringe kit	40 MG/0.8ML	2	Syringes	28	DAYS				
6627001505F520	Cyltezo starter package f	adalimumab-adbm auto-injector kit	40 MG/0.8ML	1	Kit	180	DAYS	00597037516; 00597054566			

Wildcard	Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	QL Amount	Dose Form	Days Supply	Duration	Targeted NDCs When Exclusions Exist	Age Limit	Effective Date	Term Date
6627001505F520	Cyltezo starter package f	adalimumab-adbm auto-injector kit	40 MG/0.8ML	1	Kit	180	DAYS	00597037523 ; 00597054544			
662900300021	Enbrel	etanercept for subcutaneous inj	25 MG	8	Vials	28	DAYS				
66290030002015	Enbrel	Etanercept Subcutaneous Inj 25 mg/0.5ml	25 MG/0.5ML	8	Vials	28	DAYS				
6629003000E525	Enbrel	Etanercept Subcutaneous Soln Prefilled Syringe 25 MG/0.5ML	25 MG/0.5ML	4	Syringes	28	DAYS				
6629003000E530	Enbrel	Etanercept Subcutaneous Soln Prefilled Syringe 50 MG/ML	50 MG/ML	4	Syringes	28	DAYS				
6629003000E2	Enbrel mini	etanercept subcutaneous solution cartridge	50 MG/ML	4	Cartridg es	28	DAYS				
6629003000D5	Enbrel sureclick	etanercept subcutaneous solution auto-injector	50 MG/ML	4	Pens	28	DAYS				
5250308000D220	Entyvio	vedolizumab soln pen- injector	108 MG/0.68ML	2	Pens	28	DAYS				
6627001520E510	Hadlima	adalimumab-bwwd soln prefilled syringe	40 MG/0.4ML	2	Syringes	28	DAYS				
6627001520E520	Hadlima	adalimumab-bwwd soln prefilled syringe	40 MG/0.8ML	2	Syringes	28	DAYS				
6627001520D510	Hadlima pushtouch	adalimumab-bwwd soln auto-injector	40 MG/0.4ML	2	Pens	28	DAYS				
6627001520D520	Hadlima pushtouch	adalimumab-bwwd soln auto-injector	40 MG/0.8ML	2	Pens	28	DAYS				
6627001535F520	Hulio	adalimumab-fkjp auto- injector kit	40 MG/0.8ML	2	Pens	28	DAYS				
6627001535F810	Hulio	adalimumab-fkjp prefilled syringe kit	20 MG/0.4ML	2	Syringes	28	DAYS				
6627001535F820	Hulio	adalimumab-fkjp prefilled syringe kit	40 MG/0.8ML	2	Syringes	28	DAYS				
6627001500F804	Humira	Adalimumab Prefilled Syringe Kit 10 MG/0.1ML	10 MG/0.1ML	2	Syringes	28	DAYS				
6627001500F809	Humira	Adalimumab Prefilled Syringe Kit 20 MG/0.2ML	20 MG/0.2ML	2	Syringes	28	DAYS				
6627001500F830	Humira	Adalimumab Prefilled Syringe Kit 40 MG/0.4ML	40 MG/0.4ML	2	Syringes	28	DAYS				
6627001500F820	Humira	Adalimumab Prefilled Syringe Kit 40 MG/0.8ML	40 MG/0.8ML	2	Syringes	28	DAYS				

Wildcard	Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	QL Amount	Dose Form	Days Supply	Duration	Targeted NDCs When Exclusions Exist	Age Limit	Effective Date	Term Date
6627001500F840	Humira pediatric crohns d	Adalimumab Prefilled Syringe Kit 80 MG/0.8ML	80 MG/0.8ML	1	Kit	180	DAYS				
6627001500F880	Humira pediatric crohns d	Adalimumab Prefilled Syringe Kit 80 MG/0.8ML & 40 MG/0.4ML	80 MG/0.8ML & 40MG/0.4ML	1	Kit	180	DAYS				
6627001500F440	Humira pen	adalimumab pen- injector kit	80 MG/0.8ML	2	Pens	28	DAYS	00074012402 ; 83457012402			
6627001500F430	Humira pen	Adalimumab Pen- injector Kit 40 MG/0.4ML	40 MG/0.4ML	2	Pens	28	DAYS				
6627001500F440	Humira pen- cd/uc/hs start	adalimumab pen- injector kit	80 MG/0.8ML	1	Kit	180	DAYS	00074012403			
6627001500F420	Humira pen- cd/uc/hs start	Adalimumab Pen- injector Kit ; adalimumab pen- injector kit	40 MG/0.8ML	1	Kit	180	DAYS	00074433906			
6627001500F440	Humira pen- pediatric uc s	adalimumab pen- injector kit	80 MG/0.8ML	4	Pens	180	DAYS	00074012404			
6627001500F420	Humira pen- ps/uv starter	Adalimumab Pen- injector Kit ; adalimumab pen- injector kit	40 MG/0.8ML	1	Kit	180	DAYS	00074433907			
6627001500F450	Humira pen- ps/uv starter	Adalimumab Pen- injector Kit 80 MG/0.8ML & 40 MG/0.4ML	80 MG/0.8ML & 40MG/0.4ML	1	Kit	180	DAYS				
6627001504D515	Hyrimoz	adalimumab-adaz soln auto-injector	40 MG/0.4ML	2	Pens	28	DAYS				
6627001504D515	Hyrimoz	adalimumab-adaz soln auto-injector	40 MG/0.4ML	2	Pens	28	DAYS				
6627001504D520	Hyrimoz	adalimumab-adaz soln auto-injector	40 MG/0.8ML	2	Pens	28	DAYS				
6627001504E508	Hyrimoz	adalimumab-adaz soln prefilled syringe	10 MG/0.1 ML	2	Syringes	28	DAYS				
6627001504E513	Hyrimoz	adalimumab-adaz soln prefilled syringe	20 MG/0.2ML	2	Syringes	28	DAYS				
6627001504E515	Hyrimoz	adalimumab-adaz soln prefilled syringe	40 MG/0.4ML	2	Syringes	28	DAYS				
6627001504E520	Hyrimoz	adalimumab-adaz soln prefilled syringe	40 MG/0.8ML	2	Syringes	28	DAYS				
6627001504D540	Hyrimoz ; Hyrimoz sensoready pens	adalimumab-adaz soln auto-injector	80 MG/0.8ML	2	Pens	28	DAYS	61314045420 ; 83457010701			
6627001504D540	Hyrimoz crohn's disease a ; Hyrimoz	adalimumab-adaz soln auto-injector	80 MG/0.8ML	1	Starter Kit	180	DAYS	61314045436; 83457011301			

Wildcard	Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	QL Amount	Dose Form	Days Supply	Duration	Targeted NDCs When Exclusions Exist	Age Limit	Effective Date	Term Date
	sensoready pens										
6627001504E560	Hyrimoz pediatric crohn's	adalimumab-adaz soln prefilled syr	80 MG/0.8ML & 40MG/0.4ML	2	Syringes	180	DAYS				
6627001504E540	Hyrimoz pediatric crohns	adalimumab-adaz soln prefilled syringe	80 MG/0.8ML	3	Syringes	180	DAYS				
6627001504D560	Hyrimoz plaque psoriasis	adalimumab-adaz soln auto-injector	80 MG/0.8ML & 40MG/0.4ML	1.6	Starter Kit	180	DAYS				
6627001502F540	Idacio (2 pen)	adalimumab-aacf auto- injector kit	40 MG/0.8ML	2	Pens	28	DAYS	65219055408; 65219061299			
6627001502F840	Idacio (2 syringe)	adalimumab-aacf prefilled syringe kit	40 MG/0.8ML	1	Kit	28	DAYS				
6627001502F540	Idacio starter package fo	adalimumab-aacf auto- injector kit	40 MG/0.8ML	1	Kit	180	DAYS	65219055438			
6627001502F540	Idacio starter package fo	adalimumab-aacf auto- injector kit	40 MG/0.8ML	1	Kit	180	DAYS	65219055428			
6650006000E5	Kevzara	sarilumab subcutaneous soln prefilled syringe	150 MG/1.14ML; 200 MG/1.14ML	2	Syringes	28	DAYS				
6650006000D5	Kevzara	sarilumab subcutaneous solution auto-injector	150 MG/1.14ML; 200 MG/1.14ML	2	Pens	28	DAYS				
6626001000E5	Kineret	anakinra subcutaneous soln prefilled syringe	100 MG/0.67ML	28	Syringes	28	DAYS				
90731060100120	Litfulo	ritlecitinib tosylate cap	50 MG	28	Capsule s	28	DAYS				
666030100003	Olumiant	baricitinib tab	1 MG ; 2 MG ; 4 MG	30	Tablets	30	DAYS				
5250405040D520	Omvoh	mirikizumab-mrkz subcutaneous soln auto- injector	100 MG/ML	2	Pens	28	DAYS				
6640001000E520	Orencia	Abatacept Subcutaneous Soln Prefilled Syringe 125 MG/ML	125 MG/ML	4	Syringes	28	DAYS				
6640001000E510	Orencia	Abatacept Subcutaneous Soln Prefilled Syringe 50 MG/0.4ML	50 MG/0.4ML	4	Syringes	28	DAYS				
6640001000E515	Orencia	Abatacept Subcutaneous Soln Prefilled Syringe 87.5 MG/0.7ML	87.5 MG/0.7ML	4	Syringes	28	DAYS				
6640001000D5	Orencia clickject	abatacept subcutaneous soln auto-injector	125 MG/ML	4	Syringes	28	DAYS				

Wildcard	Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	QL Amount	Dose Form	Days Supply	Duration	Targeted NDCs When Exclusions Exist	Age Limit	Effective Date	Term Date
66603072007530	Rinvoq	Upadacitinib Tab ER	30 MG	30	Tablets	30	DAYS				
66603072007540	Rinvoq	Upadacitinib Tab ER	45 MG	84	Tablets	365	DAYS				
66603072007520	Rinvoq	Upadacitinib Tab ER 24HR 15 MG	15 MG	30	Tablets	30	DAYS				<u> </u>
66603072002020	Rinvoq Iq	upadacitinib oral soln	1	360	mLs	30	DAYS				
9025052000E5	Siliq	brodalumab subcutaneous soln prefilled syringe	210 MG/1.5ML	2	Syringes	28	DAYS				<u> </u>
6627001540F520	Simlandi 1-pen kit ; Simlandi 2-pen kit	adalimumab-ryvk auto- injector kit	40 MG/0.4ML	2	Pens	28	DAYS				
6627004000D540	Simponi	Golimumab Subcutaneous Soln Auto-injector 100 MG/ML	100 MG/ML	1	Syringe	28	DAYS				
6627004000D520	Simponi	Golimumab Subcutaneous Soln Auto-injector 50 MG/0.5ML	50 MG/0.5ML	1	Syringe	28	DAYS				
6627004000E540	Simponi	Golimumab Subcutaneous Soln Prefilled Syringe 100 MG/ML	100 MG/ML	1	Syringe	28	DAYS				
6627004000E520	Simponi	Golimumab Subcutaneous Soln Prefilled Syringe 50 MG/0.5ML	50 MG/0.5ML	1	Syringe	28	DAYS				
9025057070F8	Skyrizi	risankizumab-rzaa sol prefilled syringe	75 MG/0.83ML	1	Вох	84	DAYS				
9025057070E5	Skyrizi	risankizumab-rzaa soln prefilled syringe	150 MG/ML	1	Injection Device	84	DAYS				
5250406070E210	Skyrizi	Risankizumab-rzaa Subcutaneous Soln Cartridge	180 MG/1.2ML	1	Cartridg es	56	DAY				
5250406070E220	Skyrizi	Risankizumab-rzaa Subcutaneous Soln Cartridge	360 MG/2.4ML	1	Cartridg es	56	DAYS				
9025057070D5	Skyrizi pen	risankizumab-rzaa soln auto-injector	150 MG/ML	1	Pen	84	DAYS				
90250524000320	Sotyktu	Deucravacitinib Tab	6 MG	30	Tablets	30	DAYS				·
90250585002020	Stelara	Ustekinumab Inj 45 MG/0.5ML	45 MG/0.5ML	1	Vial	84	DAYS				
9025058500E520	Stelara	Ustekinumab Soln Prefilled Syringe 45 MG/0.5ML	45 MG/0.5ML	1	Syringe	84	DAYS				
9025058500E540	Stelara	Ustekinumab Soln Prefilled Syringe 90 MG/ML	90 MG/ML	1	Syringe	56	DAYS				

Wildcard	Target Brand Agent Name(s)	Target Generic Agent Name(s)	Strength	QL Amount	Dose Form	Days Supply	Duration	Targeted NDCs When Exclusions Exist	Age Limit	Effective Date	Term Date
9025055400D5	Taltz	ixekizumab subcutaneous soln auto- injector	80 MG/ML	1	Syringe	28	DAYS				
9025055400E5	Taltz	ixekizumab subcutaneous soln prefilled syringe	80 MG/ML	1	Syringe	28	DAYS				
9025054200D2	Tremfya	guselkumab soln pen- injector	100 MG/ML	1	Pen	56	DAYS				
9025054200E5	Tremfya	guselkumab soln prefilled syringe	100 MG/ML	1	Syringe	56	DAYS				
52504525100350	Velsipity	etrasimod arginine tab	2 MG	30	Tablets	30	DAYS				
66603065102020	Xeljanz	Tofacitinib Citrate Oral Soln	1 MG/ML	240	mLs	30	DAYS				
66603065100330	Xeljanz	Tofacitinib Citrate Tab 10 MG (Base Equivalent)	10 MG	240	Tablets	365	DAYS				
66603065100320	Xeljanz	Tofacitinib Citrate Tab 5 MG (Base Equivalent)	5 MG	60	Tablets	30	DAYS				
66603065107530	Xeljanz xr	Tofacitinib Citrate Tab ER 24HR 11 MG (Base Equivalent)	11 MG	30	Tablets	30	DAYS				
66603065107550	Xeljanz xr	Tofacitinib Citrate Tab ER 24HR 22 MG (Base Equivalent)	22 MG	120	Tablets	365	DAYS				
6627001503F530	Yuflyma 1-pen kit	adalimumab-aaty auto- injector kit	40 MG/0.4ML	2	Pens	28	DAYS	72606002209 ; 72606003009			
6627001503F560	Yuflyma 1-pen kit	adalimumab-aaty auto- injector kit	80 MG/0.8ML	2	Pens	28	DAYS	72606002304 ; 72606004004			
6627001503F530	Yuflyma 2-pen kit	adalimumab-aaty auto- injector kit	40 MG/0.4ML	2	Pens	28	DAYS	72606002210 ; 72606003010			
6627001503F820	Yuflyma 2- syringe kit	adalimumab-aaty prefilled syringe kit	20 MG/0.2ML	2	Syringes	28	DAYS				
6627001503F830	Yuflyma 2- syringe kit	adalimumab-aaty prefilled syringe kit	40 MG/0.4ML	1	Kit	28	DAYS				
6627001503F560	Yuflyma cd/uc/hs starter	adalimumab-aaty auto- injector kit	80 MG/0.8ML	1	Kit	180	DAYS	72606002307			
6627001509D240	Yusimry	adalimumab-aqvh soln pen-injector	40 MG/0.8ML	2	Pens	28	DAYS				
5250504020F530	Zymfentra 1- pen	infliximab-dyyb soln auto-injector kit	120 MG/ML	2	Pens	28	DAYS	72606002501			
5250504020F530	Zymfentra 2- pen	infliximab-dyyb soln auto-injector kit	120 MG/ML	2	Pens	28	DAYS	72606002502			
5250504020F830	Zymfentra 2- syringe	infliximab-dyyb soln prefilled syringe kit	120 MG/ML	2	Syringes	28	DAYS				

PRIOR AUTHORIZATION CLINICAL CRITERIA FOR APPROVAL

Module	ORIZATION CLINICAL CRI			ria for Approv	<i>r</i> al		
Option A -			Step	Table			
FlexRx, GenRx,		S	tep 1 Step 1b (Directed			Step 3b	
BasicRx, and KeyRx	Disease State	Step 1a	to ONE TNF inhibitor) NOTE: Please see Step 1a for preferred TNF inhibitors	Step 2 (Directed to ONE step 1 agent)	Step 3a (Directed to TWO step 1 agents)	(Directed to TWO agents from step 1 and/or step 2)	Step 3c (Directed to THREE step 1 agents)
			Rheumato	oid Disorders			
	Ankylosing Spondylitis (AS)	SQ: Cosentyx, Enbrel, Hadlima, Humira	Oral: Rinvoq, Xeljanz, Xeljanz XR	N/A	SQ: Cimzia, Simponi, Taltz	N/A	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,
	Nonradiographic Axial Spondyloarthritis (nr- axSpA)	SQ: Cimzia, Cosentyx	Oral: Rinvoq	N/A	SQ: Taltz	N/A	N/A
	Polyarticular Juvenile Idiopathic Arthritis (PJIA)	SQ: Enbrel, Hadlima, Humira	Oral: Rinvoq, Rinvoq LQ, Xeljanz	SQ: Actemra (Hadlima, or Humira is a required Step 1 agent)	N/A	SQ: Orencia	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,
	Psoriatic Arthritis (PsA)	SQ: Cosentyx, Enbrel, Hadlima, Humira, Skyrizi, Stelara, Tremfya	Oral: Rinvoq, Rinvoq LQ, Xeljanz, Xeljanz XR	N/A	SQ: Cimzia, Orencia, Simponi, Taltz	N/A	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**, Yusimry**

Module	Clinical Criteria for Approval										
	Rheumatoid Arthritis	SQ: Enbrel, Hadlima, Humira	Oral: Rinvoq, Xeljanz, Xeljanz XR	SQ: Actemra (H adlima, or Humira is a required Step 1 agent)	Oral: Olumiant SQ: Cimzia, Kevzara, Kineret, Orencia, Simponi	N/A	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**, Yusimry**				
			Dermatolo	gical Disorde	r						
	Hidradenitis Suppurativa (HS)	SQ: Cosentyx, Hadlima, Humira	N/A	N/A	N/A	N/A	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,				
	Psoriasis (PS)	SQ: Cosentyx, Enbrel, Hadlima, Humira, Skyrizi, Stelara, Tremfya Oral: Otezla	N/A	Oral: Sotyktu	SQ: Cimzia, Ilumya	N/A	SQ: Abrilada**, Amjevita**, Bimzelx, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Siliq, Simlandi**, Taltz, Yuflyma**, Yusimry**				
		п	Inflammator	y Bowel Disea	ise		1				
	Crohn's Disease	SQ: Hadlima, Humira, Skyrizi, Stelara	Oral: Rinvoq	N/A	SQ: Cimzia (Hadlima, or Humira is a required Step 1 agent)	N/A	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,				

Module	Clinical Criteria for Approval											
							Yusimry**, Zymfentra					
	Ulcerative Colitis	SQ: Hadlima, Humira, Stelara	Oral: Rinvoq, Xeljanz, Xeljanz XR	SQ: Simponi (Ha dlima, or Humira is a required Step 1 agent)	N/A	SQ: Entyvio Oral: Zeposia (Hadlima, Humira, Rinvoq, Stelara, OR Xeljanz/Xelja nz XR are required Step agents)	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Omvoh, Simlandi**, Yuflyma**, Yusimry**, Zymfentra Oral Velsipity					
			C	ther								
	Uveitis	SQ: Hadlima, Humira	N/A	N/A	N/A	N/A	SQ: Abrilada**, Amjevita**, Cyltezo**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**, Yusimry**					
	I	ndications Wit	hout Prerequisite B	Biologic Immun	omodulator	s Required						
	Alopecia Areata Atopic Dermatitis Deficiency of IL-1 Receptor Antagonist (DIRA) Enthesitis Related Arthritis (ERA) Giant Cell Arteritis (GCA) Juvenile Psoriatic Arthritis (JPsA)	N/A	N/A	N/A	N/A	N/A	N/A					

Module	Clinical Criteria for Approval									
	Neonatal-Onset Multisystem Inflammatory Disease (NOMID)									
	Polymyalgia Rheumatica (PMR)									
	Systemic Juvenile Idiopathic Arthritis (SJIA)									
	Systemic Sclerosis- associated Interstitial Lung Disease (SSc-ILD)									

^{*}Note: For Xeljanz products (Xeljanz and Xeljanz XR) and Rinvoq products (Rinvoq and Rinvoq LQ), a trial of either or both dosage forms collectively counts as ONE product

Note: Branded generic available for Cyltezo, Idacio, Hulio, Hyrimoz, and Yuflyma and are included as a target at same step level in this program

Initial Evaluation

Target Agent(s) will be approved when ALL of the following are met:

- 1. The request is NOT for use of Olumiant in the treatment of coronavirus disease 2019 (COVID-19) in hospitalized adults requiring supplemental oxygen, non-invasive or invasive mechanical ventilation, or extracorporeal membrane oxygenation (ECMO) *NOTE: This indication is not covered under the pharmacy benefit **AND**
- 2. If the request is for use in Alopecia Areata and Alopecia Areata is NOT restricted from coverage under the patient's benefit **AND**
- 3. ONE of the following:
 - A. The requested agent is eligible for continuation of therapy AND ONE of the following:

Agents Eligible for Continuation of Therapy
All target agents EXCEPT the following are eligible for
ontinuation of therapy:
Abrilada
Amjevita
Cyltezo, Adalimumab-adbm
lulio, Adalimumab-fkjp
lyrimoz, Adalimumab-adaz
dacio, Adalimumab-aacf
Omvoh
imlandi
uflyma, Adalimumab-aaty
'usimry
/ymfentra

^{**}Note: Hadlima and Humira are required Step 1 agents

Module	Clinical Criteria for Approval
	 The patient has been treated with the requested agent (starting on samples is not approvable) within the past 90 days OR The prescriber states the patient has been treated with the requested agent (starting on samples is not approvable) within the past 90 days AND is at risk if therapy is changed OR
	samples is not approvable) within the past 90 days AND is at risk if therapy is changed OR B. ALL of the following: 1. The patient has an FDA labeled indication or an indication supported in compendia for the requested agent and route of administration AND ONE of the following: A. The patient has a diagnosis of moderately to severely active rheumatoid arthritis (RA) AND BOTH of the following: 1. ONE of the following: A. The patient has tried and had an inadequate response to maximally tolerated methotrexate (e.g., titrated to 25 mg weekly) after at least a 3-month duration of therapy OR B. The patient has tried and had an inadequate response to another conventional agent (i.e., hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA after at least a 3-month duration of therapy OR C. The patient has an intolerance or hypersensitivity to ONE of the following conventional agents (i.e., maximally tolerated methotrexate, hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA OR D. The patient has an FDA labeled contraindication to ALL of the following conventional agents (i.e., methotrexate, hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA OR E. The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of RA OR F. The patient is currently being treated with the requested agent
	F. The patient is currently being treated with the requested agent as indicated by ALL of the following: 1. A statement by the prescriber that the patient is currently taking the requested agent AND 2. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND 3. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR G. The prescriber has provided documentation that ALL conventional agents (i.e., methotrexate, hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm AND 2. If the request is for Simponi, ONE of the following: A. The patient will be taking the requested agent in combination with methotrexate OR B. The patient has an intolerance, FDA labeled contraindication, or hypersensitivity to methotrexate OR

Module	Clinical Criteria for Approval
	B. The patient has a diagnosis of active psoriatic arthritis (PsA) AND ONE of the following:
	 The patient has tried and had an inadequate response to ONE conventional agent (i.e., cyclosporine, leflunomide, methotrexate, sulfasalazine) used in the treatment of PsA after at least a 3-month duration of therapy OR The patient has an intolerance or hypersensitivity to ONE of the conventional agents used in the treatment of PsA OR
	3. The patient has an FDA labeled contraindication to ALL of the conventional agents used in the treatment of PsA OR
	4. The patient has severe active PsA (e.g., erosive disease, elevated markers of inflammation [e.g., ESR, CRP] attributable to PsA, long-term damage that interferes with function [i.e., joint deformities], rapidly progressive) OR
	5. The patient has concomitant severe psoriasis (PS) (e.g., greater than 10% body surface area involvement, occurring on select locations [i.e., hands, feet, scalp, face, or genitals], intractable pruritus, serious emotional consequences) OR
	6. The patient's medication history indicates use of another biologic immunomodulator agent OR Otezla that is FDA labeled or supported in compendia for the treatment of PsA OR
	 7. The patient is currently being treated with the requested agent as indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking the requested agent AND B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested
	agent AND C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	8. The prescriber has provided documentation that ALL conventional agents (i.e., cyclosporine, leflunomide, methotrexate, sulfasalazine) used in the treatment of PsA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonabl functional ability in performing daily activities or cause physical or menta
	harm OR C. The patient has a diagnosis of moderate to severe plaque psoriasis (PS) AND ONE
	of the following:
	 The patient has tried and had an inadequate response to ONE conventional agent (i.e., acitretin, anthralin, calcipotriene, calcitriol, coal tar products, cyclosporine, methotrexate, pimecrolimus, PUVA [phototherapy], tacrolimus, tazarotene, topical corticosteroids) used in the treatment of PS after at least a 3-month duration of therapy OR
	 The patient has an intolerance or hypersensitivity to ONE conventional agent used in the treatment of PS OR
	 The patient has an FDA labeled contraindication to ALL conventional agents used in the treatment of PS OR
	4. The patient has severe active PS (e.g., greater than 10% body surface are involvement, occurring on select locations [i.e., hands, feet, scalp, face, o genitals], intractable pruritus, serious emotional consequences) OR

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	5. 6.	The patient has concomitant severe psoriatic arthritis (PsA) (e.g., erosive disease, elevated markers of inflammation [e.g., ESR, CRP] attributable to PsA, long-term damage that interferes with function [i.e., joint deformities], rapidly progressive) OR The patient's medication history indicates use of another biologic
		immunomodulator agent OR Otezla that is FDA labeled or supported in compendia for the treatment of PS OR
	7.	The patient is currently being treated with the requested agent as indicated by ALL of the following:
		A. A statement by the prescriber that the patient is currently taking the requested agent AND
		 A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
		C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	8.	The prescriber has provided documentation that ALL conventional agents (i.e., acitretin, anthralin, calcipotriene, calcitriol, coal tar products,
		cyclosporine, methotrexate, pimecrolimus, PUVA [phototherapy],
		tacrolimus, tazarotene, topical corticosteroids) used in the treatment of
		PS cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of
		the patient to achieve or maintain reasonable functional ability in
		performing daily activities or cause physical or mental harm OR
	-	ient has a diagnosis of moderately to severely active Crohn's disease (CD)
		NE of the following:
	1.	The patient has tried and had an inadequate response to ONE
		conventional agent (i.e., 6-mercaptopurine, azathioprine, corticosteroids [e.g., prednisone, budesonide EC capsule], methotrexate) used in the
		treatment of CD after at least a 3-month duration of therapy OR
	2.	The patient has an intolerance or hypersensitivity to ONE of the
		conventional agents used in the treatment of CD OR
	3.	The patient has an FDA labeled contraindication to ALL of the
	,	conventional agents used in the treatment of CD OR
	4.	The patient's medication history indicates use of another biologic
		immunomodulator agent that is FDA labeled or supported in compendia for the treatment of CD OR
	5.	The patient is currently being treated with the requested agent as
		indicated by ALL of the following:
		A. A statement by the prescriber that the patient is currently taking the requested agent AND
		B. A statement by the prescriber that the patient is currently
		receiving a positive therapeutics outcome on requested agent AND
		 The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6.	The prescriber has provided documentation that ALL conventional
		agents (i.e., 6-mercaptopurine, azathioprine, corticosteroids [e.g.,
		prednisone, budesonide EC capsule], methotrexate) used in the
		treatment of CD cannot be used due to a documented medical condition
		or comorbid condition that is likely to cause an adverse reaction,
		decrease ability of the patient to achieve or maintain reasonable

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	functional ability in performing daily activities or cause physical or mental
	harm OR
	E. The patient has a diagnosis of moderately to severely active ulcerative colitis (UC)
	AND ONE of the following:
	1. The patient has tried and had an inadequate response to ONE
	conventional agent (i.e., 6-mercaptopurine, azathioprine, balsalazide,
	corticosteroids, cyclosporine, mesalamine, sulfasalazine) used in the treatment of UC after at least a 3-month duration of therapy OR
	2. The patient has severely active ulcerative colitis OR
	3. The patient has an intolerance or hypersensitivity to ONE of the
	conventional agents used in the treatment of UC OR
	4. The patient has an FDA labeled contraindication to ALL of the
	conventional agents used in the treatment of UC OR
	5. The patient's medication history indicates use of another biologic
	immunomodulator agent that is FDA labeled or supported in compendia
	for the treatment of UC OR
	6. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be
	ineffective or cause harm OR
	7. The prescriber has provided documentation that ALL conventional
	agents (i.e., 6-mercaptopurine, azathioprine, balsalazide, corticosteroids,
	cyclosporine, mesalamine, sulfasalazine) used in the treatment of UC
	cannot be used due to a documented medical condition or comorbid
	condition that is likely to cause an adverse reaction, decrease ability of
	the patient to achieve or maintain reasonable functional ability in
	performing daily activities or cause physical or mental harm OR
	 F. The patient has a diagnosis of non-infectious intermediate uveitis, posterior uveitis, or panuveitis AND ONE of the following:
	1. BOTH of the following:
	A. ONE of the following:
	1. The patient has tried and had an inadequate response
	to oral corticosteroids used in the treatment of non-
	infectious intermediate uveitis, posterior uveitis, or
	panuveitis after at least a 2-week duration of
	therapy OR
	2. The patient has tried and had an inadequate response
	to periocular or intravitreal corticosteroid injections in
	the treatment of non-infectious intermediate uveitis,
	posterior uveitis, or panuveitis OR
	3. The patient has an intolerance or hypersensitivity to
	oral corticosteroids OR periocular or intravitreal corticosteroid injections used in the treatment of non-
	infectious intermediate uveitis, posterior uveitis, or
	panuveitis OR
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	 The patient has an FDA labeled contraindication to BOTH oral corticosteroids and periocular/intravitreal corticosteroids OR The patient is currently being treated with the
	requested agent as indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking the requested agent AND
	B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OF
	6. The prescriber has provided documentation that BOTH oral corticosteroids and periocular/intravitreal
	corticosteroids cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the
	patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or
	mental harm AND B. ONE of the following:
	1. The patient has tried and had an inadequate response
	to ONE conventional systemic agent (i.e., azathioprine,
	mycophenolate, methotrexate, cyclosporine,
	tacrolimus) used in the treatment of non-infectious
	intermediate uveitis, posterior uveitis, or panuveitis
	after at least a 3-month duration of therapy OR 2. The patient has an intolerance or hypersensitivity to
	ONE conventional systemic agent used in the treatmen
	of non-infectious intermediate uveitis, posterior uveitis
	or panuveitis OR
	3. The patient has an FDA labeled contraindication to ALL
	conventional systemic agents used in the treatment of
	non-infectious intermediate uveitis, posterior uveitis, o
	panuveitis OR
	 The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient
	is currently taking the requested agent AND
	B. A statement by the prescriber that the patient
	is currently receiving a positive therapeutics
	outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OF
	5. The prescriber has provided documentation that ALL
	conventional systemic agents used in the treatment of
	non-infectious intermediate uveitis, posterior uveitis, o
	panuveitis cannot be used due to a documented
	medical condition or comorbid condition that is likely to
	cause an adverse reaction, decrease ability of the
	patient to achieve or maintain reasonable functional

Module	Clinical Criteria for Approval
	ability in performing daily activities or cause physical or mental harm OR 2. The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of non-infectious intermediate uveitis, posterior uveitis,
	or panuveitis OR
	 G. The patient has a diagnosis of giant cell arteritis (GCA) AND ONE of the following: The patient has tried and had an inadequate response to systemic corticosteroids (e.g., prednisone, methylprednisolone) used in the treatment of GCA after at least a 7-10 day duration of therapy OR
	 The patient has an intolerance or hypersensitivity to systemic corticosteroids used in the treatment of GCA OR
	3. The patient has an FDA labeled contraindication to ALL systemic corticosteroids OR
	 The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of GCA OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	 B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL systemic corticosteroids (e.g., prednisone, methylprednisolone) used in the treatment of GCA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm OR
	H. The patient has a diagnosis of active ankylosing spondylitis (AS) AND ONE of the
	following: 1. The patient has tried and had an inadequate response to TWO different
	NSAIDs used in the treatment of AS after at least a 4-week total trial OR 2. The patient has an intolerance or hypersensitivity to TWO different NSAIDs used in the treatment of AS OR
	 The patient has an FDA labeled contraindication to ALL NSAIDs used in the treatment of AS OR
	4. The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of AS OR
	 The patient is currently being treated with the requested agent as indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking the requested agent AND B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND

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	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL NSAIDs used in the treatment of AS cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm OR
	I. The patient has a diagnosis of active non-radiographic axial spondyloarthritis (nr-
	axSpA) AND ONE of the following:
	1. The patient has tried and had an inadequate response to TWO different NSAIDs used in the treatment of nr-axSpA after at least a 4-week total trial OR
	 The patient has an intolerance or hypersensitivity to TWO different NSAIDs used in the treatment of nr-axSpA OR
	 The patient has an FDA labeled contraindication to ALL NSAIDs used in the treatment of nr-axSpA OR
	 The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of nr-axSpA OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	 B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL NSAIDs used in the treatment of nr-axSpA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable
	functional ability in performing daily activities or cause physical or mental harm OR
	J. The patient has a diagnosis of moderately to severely active polyarticular juvenile
	idiopathic arthritis (PJIA) AND ONE of the following:
	 The patient has tried and had an inadequate response to ONE conventional agent (i.e., methotrexate, leflunomide) used in the treatment of PJIA after at least a 3-month duration of therapy OR
	The patient has an intolerance or hypersensitivity to ONE conventional agent used in the treatment of PJIA OR
	 The patient has an FDA labeled contraindication to ALL of the conventional agents used in the treatment of PJIA OR
	4. The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia
	for the treatment of PJIA OR 5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND

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	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL conventional agents
	(i.e., methotrexate, leflunomide) used in the treatment of PJIA cannot be used due to a documented medical condition or comorbid condition that
	is likely to cause an adverse reaction, decrease ability of the patient to
	achieve or maintain reasonable functional ability in performing daily
	activities or cause physical or mental harm OR K. The patient has a diagnosis of moderate to severe hidradenitis suppurativa (HS)
	AND ONE of the following:
	1. The patient has tried and had an inadequate response to ONE
	conventional agent (i.e., oral tetracyclines [doxycycline, minocycline,
	tetracycline]; oral contraceptives [females only]; metformin [females
	only]; finasteride [females only]; spironolactone [females only]; intralesional corticosteroids [triamcinolone]; clindamycin in combination
	with rifampin; combination of rifampin, moxifloxacin, and metronidazole;
	cyclosporine, oral retinoids) used in the treatment of HS after at least a 3-
	month duration of therapy OR
	2. The patient has an intolerance or hypersensitivity to ONE conventional
	agent used in the treatment of HS OR 3. The patient has an FDA labeled contraindication to ALL conventional
	agents used in the treatment of HS OR
	4. The patient's medication history indicates use of another biologic
	immunomodulator agent that is FDA labeled or supported in compendia
	for the treatment of HS OR
	The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking
	the requested agent AND
	B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested
	agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL conventional agents
	(i.e., oral tetracyclines [doxycycline, minocycline, tetracycline]; oral
	contraceptives [females only]; metformin [females only]; finasteride
	[females only]; spironolactone [females only]; intralesional corticosteroids [triamcinolone]; clindamycin in combination with
	rifampin; combination of rifampin, moxifloxacin, and metronidazole;
	cyclosporine, oral retinoids) used in the treatment of HS cannot be used
	due to a documented medical condition or comorbid condition that is
	likely to cause an adverse reaction, decrease ability of the patient to
	achieve or maintain reasonable functional ability in performing daily
	activities or cause physical or mental harm OR L. BOTH of the following:
	The patient has a diagnosis of systemic sclerosis associated interstitial
	lung disease (SSc-ILD) AND

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		2.	The patient's diagnosis has been confirmed on high-resolution computed tomography (HRCT) or chest radiography scans OR
		The pati the follo	cient has a diagnosis of active enthesitis related arthritis (ERA) and ONE of
		1.	The patient has tried and had an inadequate response to TWO different
			NSAIDs used in the treatment of ERA after at least a 4-week total trial OR
		2.	The patient has an intolerance or hypersensitivity to TWO different
			NSAIDs used in the treatment of ERA OR
		3.	The patient has an FDA labeled contraindication to ALL NSAIDs used in
		4.	the treatment of ERA OR The patient's medication history indicates use of another biologic
		4.	immunomodulator agent that is FDA labeled or supported in compendia
			for the treatment of ERA OR
		5.	The patient is currently being treated with the requested agent as
			indicated by ALL of the following:
			A. A statement by the prescriber that the patient is currently taking
			the requested agent AND B. A statement by the prescriber that the patient is currently
			receiving a positive therapeutics outcome on requested
			agent AND
			C. The prescriber states that a change in therapy is expected to be
			ineffective or cause harm OR
		6.	The prescriber has provided documentation that ALL NSAIDs used in the treatment of ERA cannot be used due to a documented medical condition
			or comorbid condition that is likely to cause an adverse reaction,
			decrease ability of the patient to achieve or maintain reasonable
			functional ability in performing daily activities or cause physical or mental
			harm OR
	N.	-	cient has a diagnosis of moderate-to-severe atopic dermatitis (AD) AND ALL
		of the fo	ollowing: ONE of the following:
		1.	A. The patient has at least 10% body surface area involvement OR
			B. The patient has involvement of body sites that are difficult to
			treat with prolonged topical corticosteroid therapy (e.g., hands,
			feet, face, neck, scalp, genitals/groin, skin folds) OR
			C. The patient has an Eczema Area and Severity Index (EASI) score
			of greater than or equal to 16 OR D. The patient has an Investigator Global Assessment (IGA) score of
			greater than or equal to 3 AND
		2.	ONE of the following:
			A. The patient has tried and had an inadequate response to at least
			a mid- potency topical steroid used in the treatment of AD after
			at least a 4-week duration of therapy AND a topical calcineurin inhibitor (e.g., Elidel/pimecrolimus, Protopic/tacrolimus) used in
			the treatment of AD after at least a 6-week duration of
			the treatment of Ab after at least a 0-week duration of
			B. The patient has an intolerance or hypersensitivity to at least a
			mid- potency topical steroid AND a topical calcineurin inhibitor
			(e.g., Elidel/pimecrolimus, Protopic/tacrolimus) used in the
			treatment of AD OR

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	 The patient has an FDA labeled contraindication to ALL mid-, high-, and super-potency topical steroids AND topical calcineurin inhibitors used in the treatment of AD OR
	D. The patient is currently being treated with the requested agent as indicated by ALL of the following: 1. A statement by the prescriber that the patient is currently taking the requested agent AND 2. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND 3. The prescriber states that a change in therapy is
	expected to be ineffective or cause harm OR E. The prescriber has provided documentation that ALL mid-, high-, and super-potency topical steroids AND topical calcineurin inhibitors used in the treatment of AD cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm AND
	3. The prescriber has documented the patient's baseline pruritus and other symptom severity (e.g., erythema, edema, xerosis, erosions/excoriations,
	oozing and crusting, and/or lichenification) AND 4. BOTH of the following: A. The patient is currently treated with topical emollients and
	practicing good skin care AND B. The patient will continue the use of topical emollients and good skin care practices in combination with the requested agent OR
	O. BOTH of the following:
	1. The patient has a diagnosis of severe alopecia areata (AA) AND
	The patient has at least 50% scalp hair loss that has lasted 6 months or more OR
	P. The patient has a diagnosis of polymyalgia rheumatica (PMR) AND ONE of the following:
	 The patient has tried and had an inadequate response to systemic corticosteroids at a dose equivalent to at least 7.5 mg/day of prednisone used in the treatment of PMR after at least an 8-week duration of therapy OR
	 The patient is currently treated with systemic corticosteroids at a dose equivalent to at least 7.5 mg/day of prednisone and cannot tolerate a corticosteroid taper OR
	 The patient is currently being treated with the requested agent as indicated by ALL of the following:
	 A. A statement by the prescriber that the patient is currently taking the requested agent AND B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	4. The prescriber has provided documentation that ALL systemic corticosteroids used in the treatment of PMR cannot be used due to a documented medical condition or comorbid condition that is likely to
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Module	Clinical Criteria for Approval
	cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm OR
	Q. The patient has a diagnosis of juvenile psoriatic arthritis (JPsA) AND ONE of the following:
	1. The patient has tried and had an inadequate response to ONE
	conventional agent (i.e., methotrexate, leflunomide, sulfasalazine) used in
	the treatment of JPsA after at least a 3-month duration of therapy OR
	2. The patient has an intolerance or hypersensitivity to ONE conventional
	agent used in the treatment of JPsA OR
	 The patient has an FDA labeled contraindication to methotrexate OR
	4. The patient has severe active JPsA (e.g., erosive disease, elevated markers
	of inflammation [e.g., ESR, CRP] attributable to JPsA, long-term damage
	that interferes with function [i.e., joint deformities], rapidly progressive) OR
	5. The patient has concomitant severe psoriasis (PS) (e.g., greater than 10%
	body surface area involvement, occurring on select locations [i.e., hands,
	feet, scalp, face, or genitals], intractable pruritus, serious emotional
	consequences) OR
	6. The patient's medication history indicates use of another biologic
	immunomodulator agent that is FDA labeled or supported in compendia
	for the treatment of JPsA OR
	7. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested
	agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	8. The prescriber has provided documentation that ALL conventional agent
	(i.e., methotrexate, leflunomide, sulfasalazine) used in the treatment of
	JPsA cannot be used due to a documented medical condition or comorbid
	condition that is likely to cause an adverse reaction, decrease ability of
	the patient to achieve or maintain reasonable functional ability in
	performing daily activities or cause physical or mental harm OR
	R. The patient has a diagnosis not mentioned previously AND2. ONE of the following (reference Step Table):
	A. The requested indication does NOT require any prerequisite biologic
	immunomodulator agents OR
	B. The requested agent is a Step 1a agent for the requested indication OR
	C. If the requested agent is a Step 1b agent for the requested indication, then ONE of
	the following:
	 The patient has tried and had an inadequate response to ONE Tumor
	Necrosis Factor (TNF) inhibitor for the requested indication after at least a
	3-month duration of therapy (See Step 1a for preferred TNF inhibitors)
	OR
	 The patient has an intolerance (defined as an intolerance to the drug or its excipients, not to the route of administration) or hypersensitivity to therapy with a TNF inhibitor for the requested indication OR

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	3.	The patient has an FDA labeled contraindication to ALL TNF inhibitors for
		the requested indication OR
	4.	BOTH of the following:
		A. ALL TNF inhibitors are not clinically appropriate for the patient
		AND
		B. The prescriber has provided a complete list of previously tried
	_	agents for the requested indication OR
	5.	The patient is currently being treated with the requested agent as indicated by ALL of the following:
		A. A statement by the prescriber that the patient is currently taking
		the requested agent AND
		B. A statement by the prescriber that the patient is currently
		receiving a positive therapeutics outcome on requested
		agent AND
		C. The prescriber states that a change in therapy is expected to be
	_	ineffective or cause harm OR
	6.	The prescriber has provided documentation that ALL TNF inhibitors for
		the requested indication cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse
		reaction, decrease ability of the patient to achieve or maintain reasonable
		functional ability in performing daily activities or cause physical or mental
		harm OR
	D. If the re	quested agent is a Step 2 agent for the requested indication, then ONE of
	the follo	owing:
	1.	The patient has tried and had an inadequate response to ONE of the
		required Step 1 agents for the requested indication after at least a 3-
	2	month duration of therapy (See Step 2) OR
	2.	The patient has an intolerance (defined as an intolerance to the drug or its excipients, not to the route of administration) or hypersensitivity to
		ONE of the required Step 1 agents for the requested indication OR
	3.	The patient has an FDA labeled contraindication to ALL required Step 1
		agents for the requested indication OR
	4.	BOTH of the following:
		A. ALL of the required Step 1 agents are not clinically appropriate
		for the patient AND
		B. The prescriber has provided a complete list of previously tried
	5.	agents for the requested indication OR The patient is currently being treated with the requested agent as
	3.	indicated by ALL of the following:
		A. A statement by the prescriber that the patient is currently taking
		the requested agent AND
		B. A statement by the prescriber that the patient is currently
		receiving a positive therapeutics outcome on requested
		agent AND
		C. The prescriber states that a change in therapy is expected to be
		ineffective or cause harm OR
	6.	The prescriber has provided documentation that ALL required Step 1
		agents for the requested indication cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse
		reaction, decrease ability of the patient to achieve or maintain reasonable
		functional ability in performing daily activities or cause physical or mental
		harm OR
L		

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	E. If the requested agent is a Step 3a agent for the requested indication, then ONE of
	the following (chart notes required):
	 The patient has tried and had an inadequate response to TWO of the Step
	1 agents for the requested indication after at least a 3-month trial per
	agent (See Step 3a) OR
	2. The patient has an intolerance (defined as an intolerance to the drug or
	its excipients, not to the route of administration or hypersensitivity to
	TWO of the Step 1 agents for the requested indication OR 3. The patient has an FDA labeled contraindication to ALL of the Step 1
	agents for the requested indication OR
	4. BOTH of the following:
	A. ALL of the Step 1 agents are not clinically appropriate for the
	patient AND
	B. The prescriber has provided a complete list of previously tried
	agents for the requested indication OR
	The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking
	the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested
	agent AND Con The prescriber states that a shange in the rank is expected to be
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL of the Step 1 agents
	for the requested indication cannot be used due to a documented
	medical condition or comorbid condition that is likely to cause an adverse
	reaction, decrease ability of the patient to achieve or maintain reasonable
	functional ability in performing daily activities or cause physical or mental
	harm OR
	F. If the requested agent is a Step 3b agent for the requested indication, then ONE of
	the following (chart notes required):
	1. The patient has tried and had an inadequate response to TWO agents
	from Step 1 and/or Step 2 for the requested indication after at least a 3-
	month trial per agent (See Step 3b) OR 2. The patient has an intolerance (defined as an intolerance to the drug or
	its excipients, not to the route of administration) or hypersensitivity to
	TWO agents from Step 1 and/or Step 2 for the requested indication OR
	3. The patient has an FDA labeled contraindication to ALL of the Step 1 AND
	Step 2 agents for the requested indication OR
	4. BOTH of the following:
	A. ALL of the Step 1 AND Step 2 agents are not clinically appropriate
	for the patient AND
	B. The prescriber has provided a complete list of previously tried
	agents for the requested indication OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking
	the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested
	agent AND

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	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR 6. The prescriber has provided documentation that ALL of the Step 1 AND Step 2 agents for the requested indication cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or
	cause physical or mental harm OR
	G. If the requested agent is a Step 3c agent for the requested indication, then ONE of
	the following (chart notes required):
	 The patient has tried and had an inadequate response to THREE of the Step 1 agents for the requested indication after at least a 3-month trial per agent (See Step 3c) OR
	2. The patient has an intolerance (defined as an intolerance to the drug or
	its excipients, not to the route of administration) or hypersensitivity to THREE of the Step 1 agents for the requested indication OR
	 The patient has an FDA labeled contraindication to ALL of the Step 1 agents for the requested indication OR
	4. BOTH of the following: A. ALL of the Step 1 agents are not clinically appropriate for the patient AND
	B. The prescriber has provided a complete list of previously tried agents for the requested indication OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	 A statement by the prescriber that the patient is currently taking the requested agent AND
	 A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	 The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL of the Step 1 agents for the requested indication cannot be used due to a documented
	medical condition or comorbid condition that is likely to cause an adverse
	reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm AND
	3. If Cosentyx 300 mg is requested as maintenance dosing, ONE of the following:
	A. The patient has a diagnosis of moderate to severe plaque psoriasis with or without coexistent active psoriatic arthritis AND the requested dose is 300 mg every 4 weeks OR
	B. The patient has a diagnosis of hidradenitis suppurativa AND ONE of the following:
	1. The requested dose is 300 mg every 4 weeks OR
	2. The requested dose is 300 mg every 2 weeks AND the patient has tried and had an inadequate response to Cosentyx 300 mg every 4 weeks
	after at least a 3-month duration of therapy OR C. The patient has a diagnosis of active psoriatic arthritis or active ankylosing
	spondylitis AND has tried and had an inadequate response to Cosentyx 150 mg
	every 4 weeks after at least a 3-month duration of therapy AND
	 If Omvoh is requested for the treatment of ulcerative colitis, ONE of the following: A. the patient has received Omvoh IV for induction therapy OR

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	B. The patient is new to therapy and will receive Omvoh IV for induction therapy
	AND
	5. If Entyvio is requested for the treatment of ulcerative colitis, ONE of the following:
	A. The patient has received at least 2 doses of Entyvio IV therapy OR
	B. The patient is new to therapy and will receive 2 doses of Entyvio IV therapy AND 6. If Skyrizi is requested for the treatment of Crohn's disease, ONE of the following
	 If Skyrizi is requested for the treatment of Crohn's disease, ONE of the following A. The patient received Skyrizi IV for induction therapy OR
	B. The patient is new to therapy and will receive Skyrizi IV for induction therapy AND
	7. If an ustekinumab product is requested for the treatment of Crohn's disease or ulcerative
	colitis, ONE of the following:
	A. The patient received an ustekinumab IV product for induction therapy OR
	B. The patient is new to therapy and will receive an ustekinumab IV product for
	induction therapy AND
	8. If Zymfentra is requested for the treatment of Crohn's disease or ulcerative colitis, then
	ONE of the following:
	A. The patient received an infliximab IV product for induction therapy OR
	B. The patient is new to therapy and will receive an infliximab IV product for induction therapy AND
	9. If the patient has an FDA labeled indication, then ONE of the following:
	A. The patient's age is within FDA labeling for the requested indication for the
	requested agent OR
	B. There is support for using the requested agent for the patient's age for the
	requested indication AND
	4. If an ustekinumab 90 mg product is requested, then ONE of the following:
	A. The patient has a diagnosis of psoriasis AND weighs >100kg OR
	B. The patient has a dual diagnosis of psoriasis AND psoriatic arthritis AND the patient is >100kg OR
	 C. The patient has a diagnosis of Crohn's disease or ulcerative colitis AND If Actemra is requested for a diagnosis of systemic sclerosis associated interstitial lung disease, the request is
	for the Actemra syringe (NOTE: Actemra ACTpen is not approvable for SSc-ILD) AND
	6. The prescriber is a specialist in the area of the patient's diagnosis (e.g., rheumatologist for JIA, PsA, RA;
	gastroenterologist for CD, UC; dermatologist for PS, AD; pulmonologist, radiologist, pathologist,
	rheumatologist for SSc-ILD; allergist, immunologist for AD) or has consulted with a specialist in the area of
	the patient's diagnosis AND
	7. ONE of the following (Please refer to "Agents NOT to be used Concomitantly" table):
	A. The patient will NOT be using the requested agent in combination with another immunomodulatory
	agent (e.g., TNF inhibitors, JAK inhibitors, IL-4 inhibitors) OR B. The patient will be using the requested agent in combination with another immunomodulatory
	agent AND BOTH of the following:
	1. The prescribing information for the requested agent does NOT limit the use with another
	immunomodulatory agent AND
	2. There is support for the use of combination therapy (copy of support required, i.e., clinical
	trials, phase III studies, guidelines) AND
	8. The patient does NOT have any FDA labeled contraindications to the requested agent AND
	9. The patient has been tested for latent tuberculosis (TB) when required by the prescribing information for the
	requested agent AND if positive the patient has begun therapy for latent TB
	Compendia Allowed: AHFS, DrugDex 1 or 2a level of evidence, or NCCN 1 or 2a recommended use
	Length of Approval: 12 months for all agents EXCEPT adalimumab containing products for ulcerative colitis (UC),
	Rinvoq for atopic dermatitis (AD), Siliq for plaque psoriasis (PS), Xeljanz and Xeljanz XR for induction therapy for UC,
	and the agents with indications that require loading doses for new starts. NOTE: For agents that require a loading
	dose for a new start, approve the loading dose based on FDA labeling AND the maintenance dose for the remainder of

Module **Clinical Criteria for Approval** the 12 months. Adalimumab containing products for UC may be approved for 12 weeks, Rinvoq for AD may be approved for 6 months, Silig for PS may be approved for 16 weeks, and Xeljanz and Xeljanz XR for UC may be approved for 16 weeks. **NOTE: Cosentyx for the diagnoses of AS, nr-axSpA, and PSA loading doses are not approvable. NOTE: If Quantity Limit applies, please refer to Quantity Limit Criteria. Renewal Evaluation **Target Agent(s)** will be approved when ALL of the following are met: 1. The request is NOT for use of Olumiant or Actemra in the treatment of coronavirus disease 2019 (COVID-19) in hospitalized adults requiring supplemental oxygen, non-invasive or invasive mechanical ventilation, or extracorporeal membrane oxygenation (ECMO) *NOTE: This indication is not covered under the pharmacy benefit AND 2. The request is for use in Alopecia Areata and Alopecia Areata is NOT restricted from coverage under the patient's benefit AND 3. The patient has been previously approved for the requested agent through the plan's Prior Authorization process (*please note ustekinumab product renewal must be for the same strength as the initial approval) [Note: patients not previously approved for the requested agent will require initial evaluation review] AND 4. ONE of the following: The patient has a diagnosis of moderate to severe atopic dermatitis AND BOTH of the following: 1. The patient has had a reduction or stabilization from baseline (prior to therapy with the requested agent) of ONE of the following: A. Affected body surface area OR B. Flares OR C. Pruritus, erythema, edema, xerosis, erosions/excoriations, oozing and crusting, and/or lichenification OR D. A decrease in the Eczema Area and Severity Index (EASI) score OR E. A decrease in the Investigator Global Assessment (IGA) score AND 2. The patient will continue standard maintenance therapies (e.g., topical emollients, good skin care practices) in combination with the requested agent OR В. The patient has a diagnosis of polymyalgia rheumatica AND BOTH of the following: 1. The patient has had clinical benefit with the requested agent AND 2. If the requested agent is Kevzara, the patient does NOT have any of the following: A. Neutropenia (ANC less than 1,000 per mm³ at the end of the dosing interval) AND B. Thrombocytopenia (platelet count is less than 100,000 per mm^3) AND C. AST or ALT elevations 3 times the upper limit of normal **OR**

- the area of the patient's diagnosis **AND**6. ONE of the following (Please refer to "Agents NOT to be used Concomitantly" table):
 - 1. The patient will NOT be using the requested agent in combination with another immunomodulatory agent (e.g., TNF inhibitors, JAK inhibitors, IL-4 inhibitors) **OR**

The patient has a diagnosis other than moderate to severe atopic dermatitis or polymyalgia

rheumatologist for SSc-ILD; allergist, immunologist for AD) or the prescriber has consulted with a specialist in

rheumatica AND the patient has had clinical benefit with the requested agent **AND**5. The prescriber is a specialist in the area of the patient's diagnosis (e.g., rheumatologist for JIA, PsA, RA; gastroenterologist for CD, UC; dermatologist for PS, AD; pulmonologist, radiologist, pathologist,

C.

Module			Clinical Crite	eria for Approva	al			
	 The patient will be using the requested agent in combination with another immunomodulatory agent AND BOTH of the following: The prescribing information for the requested agent does NOT limit the use with another immunomodulatory agent AND There is support for the use of combination therapy (copy of support required, i.e., clinical trials, phase III studies, guidelines) AND If Cosentyx 300 mg is requested as maintenance dosing, ONE of the following: The patient has a diagnosis of moderate to severe plaque psoriasis with or without coexistent active psoriatic arthritis AND the requested dose is 300 mg every 4 weeks OR The patient has a diagnosis of hidradenitis suppurativa AND ONE of the following:							
	C. The patient has a diagnosis of active psoriatic arthritis or active ankylosing spondylitis AND has tried and had an inadequate response to Cosentyx 150 mg every 4 weeks after at least a 3-month duration of therapy AND 8. If Actemra is requested for a diagnosis of systemic sclerosis associated interstitial lung disease, the request is for the Actemra syringe (NOTE: Actemra ACTpen is not approvable for SSc-ILD) AND 9. The patient does NOT have any FDA labeled contraindications to the requested agent Compendia Allowed: AHFS, DrugDex 1 or 2a level of evidence, or NCCN 1 or 2a recommended use Length of Approval: 12 months							
Option B -	**NOTE: Cosentyx for the diagnoses of AS, nr-axSpA, and PSA loading doses are not approvable. NOTE: If Quantity Limit applies, please refer to Quantity Limit Criteria. Step Table							
Focus Rx								
	Disease State	Step 1a	ep 1 Step 1b (Directed to ONE TNF inhibitor) NOTE: Please see Step 1a for preferred TNF inhibitors	Step 2 (Directed to ONE step 1 agent)	Step 3a (Directed to TWO step 1 agents)	Step 3b (Directed to TWO agents from step 1 and/or step 2)	Step 3c (Directed to THREE step 1 agents)	
	Rheumatoid Disorders							
	Ankylosing Spondylitis (AS)	SQ: Cosentyx, Cyltezo, Enbrel, Humira	Oral: Rinvoq, Xeljanz, Xeljanz XR	N/A	SQ: Cimzia, Simponi, Taltz	N/A	SQ: Abrilada**, Amjevita**, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,	

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	Nonradiographic Axial Spondyloarthritis (nr-axSpA)	SQ: Cimzia, Cosentyx	Oral: Rinvoq	N/A	SQ: Taltz	N/A	N/A
	Polyarticular Juvenile Idiopathic Arthritis (PJIA)	SQ: Cyltezo, Enbrel, Humira	Oral: Rinvoq, Rinvoq LQ, Xeljanz	SQ: Actemra (Cyltezo or Humira a is required Step 1 agent)	N/A	SQ: Orencia	SQ: Abrilada**, Amjevita**, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,
	Psoriatic Arthritis (PsA)	SQ: Cosentyx, Cyltezo, Enbrel, Humira, Skyrizi, Stelara, Tremfya Oral: Otezla	Oral: Rinvoq, Rinvoq LQ, Xeljanz, Xeljanz XR	N/A	SQ: Cimzia, Orencia, Simponi, Taltz	N/A	SQ: Abrilada**, Amjevita*, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**, Yusimry**
	Rheumatoid Arthritis	SQ: Cyltezo, Enbrel, Humira	Oral: Rinvoq, Xeljanz, Xeljanz XR	SQ: Actemra (Cylt ezo or Humira is a required Step 1 agent)	Oral: Olumiant SQ: Cimzia, Kevzara, Kineret, Orencia, Simponi	N/A	SQ: Abrilada**, Amjevita**, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,
			Dermato	logical Disorder			
	Hidradenitis Suppurativa (HS)	SQ: Cosentyx, Cyltezo, Humira	N/A	N/A	N/A	N/A	SQ: Abrilada**, Amjevita**, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**,

Module	Clinical Criteria for Approval						
	Psoriasis (PS)	SQ: Cosentyx, Cyltezo, Enbrel, Humira, Skyrizi, Stelara, Tremfya	N/A	Oral: Sotyktu	SQ: Cimzia, Ilumya	N/A	SQ: Abrilada**, Amjevita**, Bimzelx, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Siliq, Simlandi**, Taltz, Yuflyma**, Yusimry**
			Inflammato	ory Bowel Diseas	se		
	Crohn's Disease	SQ: Cyltezo, Humira, Skyrizi, Stelara	Oral: Rinvoq	N/A	SQ: Cimzia (Cyltezo, or Humira is a required Step 1 agent)	N/A	SQ: Abrilada**, Amjevita**, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Simlandi**, Yuflyma**, Yusimry**, Zymfentra
	Ulcerative Colitis	SQ: Cyltezo, Humira, Stelara	Oral: Rinvoq, Xeljanz, Xeljanz XR	SQ: Simponi (Cylte zo, or Humira is required Step 1 agent)	N/A	SQ: Entyvio Oral: Zeposia (Cyltezo, Hu mira, Rinvoq, Stelara, OR Xeljanz/Xelja nz XR are required Step agents)	SQ: Abrilada**, Amjevita**, Hadlima**, Hulio**, Hyrimoz**, Idacio**, Omvoh, Simlandi**, Yuflyma**, Yusimry**, Zymfentra Oral: Velsipity
	Other						
		50:					SQ:
	Uveitis	SQ: Cyltezo, Humira	N/A	N/A	N/A	N/A	Abrilada**, Amjevita**, Hadlima**, Hulio**,

Vlodule			Clinical Crite	eria for Approv	al		
							Hyrimoz**, Idacio**, Simlandi**, Yuflyma**, Yusimry**
		Indications W	ithout Prerequisite	Biologic Immu	nomodulators	Required	
	Alopecia Areata						
	Atopic Dermatitis						
	Deficiency of IL-1 Receptor Antagonist (DIRA)						
	Enthesitis Related Arthritis (ERA)						
	Giant Cell Arteritis (GCA)						
	Juvenile Psoriatic Arthritis (JPsA)	N/A	N/A	N/A	N/A	N/A	N/A
	Neonatal-Onset Multisystem Inflammatory Disease (NOMID)						
	Polymyalgia Rheumatica (PMR)						
	Systemic Juvenile Idiopathic Arthritis (SJIA)						
	Systemic Sclerosis- associated Interstitial Lung Disease (SSc-ILD)						

^{*}Note: For Xeljanz products (Xeljanz and Xeljanz XR) and Rinvoq products (Rinvoq and Rinvoq LQ), a trial of either or both dosage forms collectively counts as ONE product

Note: Branded generic available for Cyltezo, Idacio, Hulio, Hyrimoz, and Yuflyma and are included as a target at same step level in this program

Initial Evaluation

Target Agent(s) will be approved when ALL of the following are met:

^{**}Note: Cyltezo and Humira are required Step 1 agents

Module **Clinical Criteria for Approval** 1. The request is NOT for use of Olumiant in the treatment of coronavirus disease 2019 (COVID-19) in hospitalized adults requiring supplemental oxygen, non-invasive or invasive mechanical ventilation, or extracorporeal membrane oxygenation (ECMO) *NOTE: This indication is not covered under the pharmacy benefit AND 2. If the request is for use in Alopecia Areata and Alopecia Areata is NOT restricted from coverage under the patient's benefit AND 3. ONE of the following: The requested agent is eligible for continuation of therapy AND ONE of the following: **Agents Eligible for Continuation of Therapy** All target agents EXCEPT the following are eligible for continuation of therapy: Abrilada Amjevita Hadlima Hulio, Adalimumab-fkjp Hyrimoz, Adalimumab-adaz Idacio, Adalimumab-aacf Omvoh Simlandi Yuflyma, Adalimumab-aaty Yusimry Zymfentra 1. The patient has been treated with the requested agent (starting on samples is not approvable) within the past 90 days OR The prescriber states the patient has been treated with the requested agent (starting on samples is not approvable) within the past 90 days AND is at risk if therapy is changed OR В. ALL of the following: 1. The patient has an FDA labeled indication or an indication supported in compendia for the requested agent and route of administration AND ONE of the following: A. The patient has a diagnosis of moderately to severely active rheumatoid arthritis (RA) AND BOTH of the following: 1. ONE of the following: A. The patient has tried and had an inadequate response to maximally tolerated methotrexate (e.g., titrated to 25 mg weekly) after at least a 3-month duration of therapy OR B. The patient has tried and had an inadequate response to another conventional agent (i.e., hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA after at least a 3month duration of therapy **OR** C. The patient has an intolerance or hypersensitivity to ONE of the following conventional agents (i.e., maximally tolerated methotrexate, hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA OR D. The patient has an FDA labeled contraindication to ALL of the following conventional agents (i.e., methotrexate, hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA OR

Module	Clinical Criteria for Approval						
Module	E. The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of RA OR F. The patient is currently being treated with the requested agent as indicated by ALL of the following: 1. A statement by the prescriber that the patient is currently taking the requested agent AND 2. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND 3. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR G. The prescriber has provided documentation that ALL conventional agents (i.e., methotrexate, hydroxychloroquine, leflunomide, sulfasalazine) used in the treatment of RA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease						
	ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm AND 2. If the request is for Simponi, ONE of the following: A. The patient will be taking the requested agent in combination						
	with methotrexate OR B. The patient has an intolerance, FDA labeled contraindication, or hypersensitivity to methotrexate OR B. The patient has a diagnosis of active psoriatic arthritis (PsA) AND ONE of the						
	following:						
	1. The patient has tried and had an inadequate response to ONE conventional agent (i.e., cyclosporine, leflunomide, methotrexate, sulfasalazine) used in the treatment of PsA after at least a 3-month duration of therapy OR						
	2. The patient has an intolerance or hypersensitivity to ONE of the						
	conventional agents used in the treatment of PsA OR 3. The patient has an FDA labeled contraindication to ALL of the						
	conventional agents used in the treatment of PsA OR 4. The patient has severe active PsA (e.g., erosive disease, elevated markers of inflammation [e.g., ESR, CRP] attributable to PsA, long-term damage that interferes with function [i.e., joint deformities], rapidly progressive) OR						
	5. The patient has concomitant severe psoriasis (PS) (e.g., greater than 10% body surface area involvement, occurring on select locations [i.e., hands, feet, scalp, face, or genitals], intractable pruritus, serious emotional						
	consequences) OR 6. The patient's medication history indicates use of another biologic immunomodulator agent OR Otezla that is FDA labeled or supported in compendia for the treatment of PsA OR						
	7. The patient is currently being treated with the requested agent as indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking the requested agent AND						

Module	Clinical Criteria for Approval
	 B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND C. The prescriber states that a change in therapy is expected to be
	ineffective or cause harm OR
	8. The prescriber has provided documentation that ALL conventional agents (i.e., cyclosporine, leflunomide, methotrexate, sulfasalazine) used in the treatment of PsA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or menta
	harm OR
	C. The patient has a diagnosis of moderate to severe plaque psoriasis (PS) AND ONE
	of the following:
	1. The patient has tried and had an inadequate response to ONE conventional agent (i.e., acitretin, anthralin, calcipotriene, calcitriol, coal tar products, cyclosporine, methotrexate, pimecrolimus, PUVA [phototherapy], tacrolimus, tazarotene, topical corticosteroids) used in the treatment of PS after at least a 3-month duration of therapy OR
	 The patient has an intolerance or hypersensitivity to ONE conventional agent used in the treatment of PS OR
	 The patient has an FDA labeled contraindication to ALL conventional agents used in the treatment of PS OR
	4. The patient has severe active PS (e.g., greater than 10% body surface are involvement, occurring on select locations [i.e., hands, feet, scalp, face, o genitals], intractable pruritus, serious emotional consequences) OR
	5. The patient has concomitant severe psoriatic arthritis (PsA) (e.g., erosive disease, elevated markers of inflammation [e.g., ESR, CRP] attributable to PsA, long-term damage that interferes with function [i.e., joint deformities], rapidly progressive) OR
	6. The patient's medication history indicates use of another biologic immunomodulator agent OR Otezla that is FDA labeled or supported in compendia for the treatment of PS OR
	7. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	 B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	8. The prescriber has provided documentation that ALL conventional agents (i.e., acitretin, anthralin, calcipotriene, calcitriol, coal tar products)
	cyclosporine, methotrexate, pimecrolimus, PUVA [phototherapy], tacrolimus, tazarotene, topical corticosteroids) used in the treatment of PS cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of
	the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm OR
	 D. The patient has a diagnosis of moderately to severely active Crohn's disease (CD) AND ONE of the following:

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	1. The patient has tried and had an inadequate response to ONE conventional agent (i.e., 6-mercaptopurine, azathioprine, corticosteroids [e.g., prednisone, budesonide EC capsule], methotrexate) used in the
	treatment of CD after at least a 3-month duration of therapy OR 2. The patient has an intolerance or hypersensitivity to ONE of the
	conventional agents used in the treatment of CD OR 3. The patient has an FDA labeled contraindication to ALL of the
	conventional agents used in the treatment of CD OR 4. The patient's medication history indicates use of another biologic
	immunomodulator agent that is FDA labeled or supported in compendia for the treatment of CD OR
	 The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL conventional
	agents (i.e., 6-mercaptopurine, azathioprine, corticosteroids [e.g.,
	prednisone, budesonide EC capsule], methotrexate) used in the
	treatment of CD cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction,
	decrease ability of the patient to achieve or maintain reasonable
	functional ability in performing daily activities or cause physical or mental harm OR
	E. The patient has a diagnosis of moderately to severely active ulcerative colitis (UC)
	AND ONE of the following:
	1. The patient has tried and had an inadequate response to ONE
	conventional agent (i.e., 6-mercaptopurine, azathioprine, balsalazide,
	corticosteroids, cyclosporine, mesalamine, sulfasalazine) used in the treatment of UC after at least a 3-month duration of therapy OR
	2. The patient has severely active ulcerative colitis OR
	3. The patient has an intolerance or hypersensitivity to ONE of the
	conventional agents used in the treatment of UC OR
	4. The patient has an FDA labeled contraindication to ALL of the
	conventional agents used in the treatment of UC OR
	 The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia
	for the treatment of UC OR
	 The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR

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	7. The prescriber has provided documentation that ALL conventional agents (i.e., 6-mercaptopurine, azathioprine, balsalazide, corticosteroids, cyclosporine, mesalamine, sulfasalazine) used in the treatment of UC cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in
	performing daily activities or cause physical or mental harm OR F. The patient has a diagnosis of non-infectious intermediate uveitis, posterior uveitis, or panuveitis AND ONE of the following:
	1. BOTH of the following:
	A. ONE of the following: 1. The patient has tried and had an inadequate response to oral corticosteroids used in the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis after at least a 2-week duration of therapy OR
	2. The patient has tried and had an inadequate response to periocular or intravitreal corticosteroid injections in the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis OR
	3. The patient has an intolerance or hypersensitivity to oral corticosteroids OR periocular or intravitreal corticosteroid injections used in the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis OR
	 The patient has an FDA labeled contraindication to BOTH oral corticosteroids and periocular/intravitreal corticosteroids OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking the requested agent AND B. A statement by the prescriber that the patient is currently receiving a positive the requested.
	is currently receiving a positive therapeutics outcome on requested agent AND C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that BOTH oral corticosteroids and periocular/intravitreal corticosteroids cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or
	mental harm AND
	B. ONE of the following: 1. The patient has tried and had an inadequate response to ONE conventional systemic agent (i.e., azathioprine, mycophenolate, methotrexate, cyclosporine, tacrolimus) used in the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis after at least a 3-month duration of therapy OR

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	2. The patient has an intolerance or hypersensitivity to ONE conventional systemic agent used in the treatment of non-infectious intermediate uveitis, posterior uveitis,
	or panuveitis OR 3. The patient has an FDA labeled contraindication to ALL conventional systemic agents used in the treatment of non-infectious intermediate uveitis, posterior uveitis, or
	panuveitis OR 4. The patient is currently being treated with the requested agent as indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking the requested agent AND B. A statement by the prescriber that the patient is currently receiving a positive therapeutics
	outcome on requested agent AND C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	5. The prescriber has provided documentation that ALL conventional systemic agents used in the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the
	patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm OR
	 The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis OR
	G. The patient has a diagnosis of giant cell arteritis (GCA) AND ONE of the following:
	1. The patient has tried and had an inadequate response to systemic corticosteroids (e.g., prednisone, methylprednisolone) used in the treatment of GCA after at least a 7-10 day duration of therapy OR
	2. The patient has an intolerance or hypersensitivity to systemic
	corticosteroids used in the treatment of GCA OR 3. The patient has an FDA labeled contraindication to ALL systemic
	 The patient has an FDA labeled contraindication to ALL systemic corticosteroids OR
	4. The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of GCA OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested agent AND C. The prescriber states that a change in therapy is expected to be
	ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL systemic corticosteroids (e.g., prednisone, methylprednisolone) used in the

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	treatment of GCA cannot be used due to a documented medical condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or men
	harm OR
	H. The patient has a diagnosis of active ankylosing spondylitis (AS) AND ONE of the
	following:
	 The patient has tried and had an inadequate response to TWO different NSAIDs used in the treatment of AS after at least a 4-week total trial OR The patient has an intolerance or hypersensitivity to TWO different
	NSAIDs used in the treatment of AS OR
	3. The patient has an FDA labeled contraindication to ALL NSAIDs used in the treatment of AS OR
	4. The patient's medication history indicates use of another biologic
	immunomodulator agent that is FDA labeled or supported in compending for the treatment of AS OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested agent AND
	 The prescriber states that a change in therapy is expected to b ineffective or cause harm OR
	 The prescriber has provided documentation that ALL NSAIDs used in the treatment of AS cannot be used due to a documented medical condition
	or comorbid condition that is likely to cause an adverse reaction,
	decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or men
	harm OR
	I. The patient has a diagnosis of active non-radiographic axial spondyloarthritis (nr axSpA) AND ONE of the following:
	 The patient has tried and had an inadequate response to TWO different NSAIDs used in the treatment of nr-axSpA after at least a 4-week total trial OR
	 The patient has an intolerance or hypersensitivity to TWO different NSAIDs used in the treatment of nr-axSpA OR
	3. The patient has an FDA labeled contraindication to ALL NSAIDs used in
	the treatment of nr-axSpA OR
	 The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendication for the treatment of nr-axSpA OR
	5. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking
	the requested agent AND
	B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested
	agent AND C. The prescriber states that a change in therapy is expected to b ineffective or cause harm OR

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	6. The prescriber has provided documentation that ALL NSAIDs used in the treatment of nr-axSpA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental
	harm OR J. The patient has a diagnosis of moderately to severely active polyarticular juvenile idiopathic arthritis (PJIA) AND ONE of the following: 1. The patient has tried and had an inadequate response to ONE conventional agent (i.e., methotrexate, leflunomide) used in the treatment of PJIA after at least a 3-month duration of therapy OR 2. The patient has an intolerance or hypersensitivity to ONE conventional agent used in the treatment of PJIA OR 3. The patient has an FDA labeled contraindication to ALL of the conventional agents used in the treatment of PJIA OR 4. The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of PJIA OR 5. The patient is currently being treated with the requested agent as indicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking the requested agent AND
	B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR 6. The prescriber has provided documentation that ALL conventional agents (i.e., methotrexate, leflunomide) used in the treatment of PJIA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm OR
	 K. The patient has a diagnosis of moderate to severe hidradenitis suppurativa (HS) AND ONE of the following: The patient has tried and had an inadequate response to ONE conventional agent (i.e., oral tetracyclines [doxycycline, minocycline, tetracycline]; oral contraceptives [females only]; metformin [females only]; finasteride [females only]; spironolactone [females only]; intralesional corticosteroids [triamcinolone]; clindamycin in combination with rifampin; combination of rifampin, moxifloxacin, and metronidazole; cyclosporine, oral retinoids) used in the treatment of HS after at least a 3-month duration of therapy OR
	 The patient has an intolerance or hypersensitivity to ONE conventional agent used in the treatment of HS OR The patient has an FDA labeled contraindication to ALL conventional agents used in the treatment of HS OR The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of HS OR The patient is currently being treated with the requested agent as
	indicated by ALL of the following:

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Wiodule	• • • • • • • • • • • • • • • • • • • •
	 A. A statement by the prescriber that the patient is currently taking the requested agent AND B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL conventional agents (i.e., oral tetracyclines [doxycycline, minocycline, tetracycline]; oral contraceptives [females only]; metformin [females only]; finasteride [females only]; spironolactone [females only]; intralesional corticosteroids [triamcinolone]; clindamycin in combination with rifampin; combination of rifampin, moxifloxacin, and metronidazole; cyclosporine, oral retinoids) used in the treatment of HS cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily
	activities or cause physical or mental harm OR
	L. BOTH of the following:
	The patient has a diagnosis of systemic sclerosis associated interstitial lung disease (SSc-ILD) AND
	 The patient's diagnosis has been confirmed on high-resolution computed tomography (HRCT) or chest radiography scans OR
	M. The patient has a diagnosis of active enthesitis related arthritis (ERA) and ONE of
	the following: 1. The patient has tried and had an inadequate response to TWO different NSAIDs used in the treatment of ERA after at least a 4-week total trial OR
	2. The patient has an intolerance or hypersensitivity to TWO different NSAIDs used in the treatment of ERA OR
	3. The patient has an FDA labeled contraindication to ALL NSAIDs used in the treatment of ERA OR
	 The patient's medication history indicates use of another biologic immunomodulator agent that is FDA labeled or supported in compendia for the treatment of ERA OR
	5. The patient is currently being treated with the requested agent as indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking the requested agent AND
	 B. A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	 The prescriber states that a change in therapy is expected to be ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL NSAIDs used in the treatment of ERA cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm OR
	 N. The patient has a diagnosis of moderate-to-severe atopic dermatitis (AD) AND ALL of the following: 1. ONE of the following:

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	 A. The patient has at least 10% body surface area involvement OR B. The patient has involvement of body sites that are difficult to treat with prolonged topical corticosteroid therapy (e.g., hands, feet, face, neck, scalp, genitals/groin, skin folds) OR C. The patient has an Eczema Area and Severity Index (EASI) score
	of greater than or equal to 16 OR D. The patient has an Investigator Global Assessment (IGA) score of
	greater than or equal to 3 AND
	2. ONE of the following:
	A. The patient has tried and had an inadequate response to at least a mid- potency topical steroid used in the treatment of AD after at least a 4-week duration of therapy AND a topical calcineurin inhibitor (e.g., Elidel/pimecrolimus, Protopic/tacrolimus) used in the treatment of AD after at least a 6-week duration of therapy OR
	 B. The patient has an intolerance or hypersensitivity to at least a mid- potency topical steroid AND a topical calcineurin inhibitor (e.g., Elidel/pimecrolimus, Protopic/tacrolimus) used in the treatment of AD OR
	C. The patient has an FDA labeled contraindication to ALL mid-, high-, and super-potency topical steroids AND topical calcineurin
	inhibitors used in the treatment of AD OR
	 The patient is currently being treated with the requested agent as indicated by ALL of the following:
	1. A statement by the prescriber that the patient is
	currently taking the requested agent AND 2. A statement by the prescriber that the patient is
	currently receiving a positive therapeutics outcome on requested agent AND
	3. The prescriber states that a change in therapy is
	expected to be ineffective or cause harm OR
	E. The prescriber has provided documentation that ALL mid-, high-, and super-potency topical steroids AND topical calcineurin
	inhibitors used in the treatment of AD cannot be used due to a
	documented medical condition or comorbid condition that is
	likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in
	performing daily activities or cause physical or mental harm AND
	3. The prescriber has documented the patient's baseline pruritus and other
	symptom severity (e.g., erythema, edema, xerosis, erosions/excoriations,
	oozing and crusting, and/or lichenification) AND 4. BOTH of the following:
	1. The patient is currently treated with topical emollients and
	practicing good skin care AND
	2. The patient will continue the use of topical emollients and good
	skin care practices in combination with the requested agent OR O. BOTH of the following:
	1. The patient has a diagnosis of severe alopecia areata (AA) AND
	2. The patient has at least 50% scalp hair loss that has lasted 6 months or
	more OR
	P. The patient has a diagnosis of polymyalgia rheumatica (PMR) AND ONE of the following:
	Tollowing.

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	corticoste prednisor	nt has tried and had an inadequate response to systemic eroids at a dose equivalent to at least 7.5 mg/day of ne used in the treatment of PMR after at least an 8-week of therapy OR
	2. The patie equivaler	nt is currently treated with systemic corticosteroids at a dose at to at least 7.5 mg/day of prednisone and cannot tolerate a deroid taper OR
	3. The patie	nt is currently being treated with the requested agent as by ALL of the following:
	A. /	A statement by the prescriber that the patient is currently taking the requested agent AND
	B. /	A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	C. i	The prescriber states that a change in therapy is expected to be neffective or cause harm OR
	corticoste documen cause an maintain	criber has provided documentation that ALL systemic eroids used in the treatment of PMR cannot be used due to a ted medical condition or comorbid condition that is likely to adverse reaction, decrease ability of the patient to achieve or reasonable functional ability in performing daily activities or
		ysical or mental harm OR diagnosis of juvenile psoriatic arthritis (JPsA) AND ONE of the
	following:	diagnosis of Juvernie psoriatic artificis (JPSA) AND ONE of the
	1. The patie convention	nt has tried and had an inadequate response to ONE onal agent (i.e., methotrexate, leflunomide, sulfasalazine) used in ment of JPsA after at least a 3-month duration of therapy OR
		nt has an intolerance or hypersensitivity to ONE conventional ed in the treatment of JPsA OR
		nt has an FDA labeled contraindication to methotrexate OR
	of inflam	nt has severe active JPsA (e.g., erosive disease, elevated markers mation [e.g., ESR, CRP] attributable to JPsA, long-term damage feres with function [i.e., joint deformities], rapidly progressive)
		nt has concomitant severe psoriasis (PS) (e.g., greater than 10%
	· ·	ace area involvement, occurring on select locations [i.e., hands, p, face, or genitals], intractable pruritus, serious emotional ences) OR
	6. The patie immunon	nt's medication history indicates use of another biologic nodulator agent that is FDA labeled or supported in compendia eatment of JPsA OR
	7. The patie indicated	nt is currently being treated with the requested agent as by ALL of the following:
	t	A statement by the prescriber that the patient is currently taking the requested agent AND
		A statement by the prescriber that the patient is currently receiving a positive therapeutics outcome on requested agent AND
	i	The prescriber states that a change in therapy is expected to be neffective or cause harm OR
		criber has provided documentation that ALL conventional agents ne treatment of JPsA cannot be used due to a documented

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	1	medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental marm OR
		nt has a diagnosis not mentioned previously AND
		ng (reference Step Table):
		ested indication does NOT require any prerequisite biologic
		nodulator agents OR ested agent is a Step 1a agent for the requested indication OR
		uested agent is a Step 1b agent for the requested indication, then ONE of
	the follow	
	1	The patient has tried and had an inadequate response to ONE Tumor Necrosis Factor (TNF) inhibitor for the requested indication after at least a 3-month duration of therapy (See Step 1a for preferred TNF inhibitors)
		The patient has an intolerance (defined as an intolerance to the drug or ts excipients, not to the route of administration) or hypersensitivity to
		herapy with a TNF inhibitor for the requested indication OR
		The patient has an FDA labeled contraindication to ALL TNF inhibitors for the requested indication OR
		BOTH of the following:
	ı.	A. ALL TNF inhibitors are not clinically appropriate for the patient AND
		B. The prescriber has provided a complete list of previously tried
		agents for the requested indication OR
		The patient is currently being treated with the requested agent as
	'	ndicated by ALL of the following: A. A statement by the prescriber that the patient is currently taking
		the requested agent AND
		B. A statement by the prescriber that the patient is currently
		receiving a positive therapeutics outcome on requested agent AND
		C. The prescriber states that a change in therapy is expected to be
		ineffective or cause harm OR
		The prescriber has provided documentation that ALL TNF inhibitors for
		the requested indication cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse
		reaction, decrease ability of the patient to achieve or maintain reasonable
		functional ability in performing daily activities or cause physical or mental
		narm OR
	-	uested agent is a Step 2 agent for the requested indication, then ONE of
	the follow	ving: The patient has tried and had an inadequate response to ONE of the
		required Step 1 agents for the requested indication after at least a 3-
		month duration of therapy (See Step 2) OR
		The patient has an intolerance (defined as an intolerance to the drug or
		ts excipients, not to the route of administration) or hypersensitivity to DNE of the required Step 1 agents for the requested indication OR
		The patient has an FDA labeled contraindication to ALL required Step 1
		agents for the requested indication OR
	4. 1	BOTH of the following:

A. ALL of the required Step 1 agents are not clinically for the patient AND B. The prescriber has provided a complete list of preagents for the requested indication OR 5. The patient is currently being treated with the requested a indicated by ALL of the following: A. A statement by the prescriber that the patient is a the requested agent AND B. A statement by the prescriber that the patient is a receiving a positive therapeutics outcome on requagent AND C. The prescriber states that a change in therapy is expected in the complex of the prescriber or cause harm OR	eviously tried agent as currently taking currently uested
6. The prescriber has provided documentation that ALL requiagents for the requested indication cannot be used due to medical condition or comorbid condition that is likely to careaction, decrease ability of the patient to achieve or main functional ability in performing daily activities or cause phyharm OR	a documented use an adverse tain reasonable
E. If the requested agent is a Step 3a agent for the requested indication	on, then ONE of
the following (chart notes required):	
1. The patient has tried and had an inadequate response to T 1 agents for the requested indication after at least a 3-mo agent (See Step 3a) OR	•
2. The patient has an intolerance (defined as an intolerance to its excipients, not to the route of administration or hypers TWO of the Step 1 agents for the requested indication OR	_
3. The patient has an FDA labeled contraindication to ALL of agents for the requested indication OR	the Step 1
4. BOTH of the following: A. ALL of the Step 1 agents are not clinically appropring patient AND B. The prescriber has provided a complete list of prescriber has provided and the state of prescriber has provided as complete list of prescriber has pre	
agents for the requested indication OR	
5. The patient is currently being treated with the requested a indicated by ALL of the following:	
A. A statement by the prescriber that the patient is of the requested agent AND	
B. A statement by the prescriber that the patient is or receiving a positive therapeutics outcome on requagent AND	uested
C. The prescriber states that a change in therapy is e ineffective or cause harm OR	expected to be
6. The prescriber has provided documentation that ALL of the	e Step 1 agents
for the requested indication cannot be used due to a docu	mented
medical condition or comorbid condition that is likely to ca reaction, decrease ability of the patient to achieve or main functional ability in performing daily activities or cause physical conditions.	tain reasonable
harm OR F. If the requested agent is a Step 3b agent for the requested indication	on then ONE of
the following (chart notes required):	ni, uien ONE OI

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	1. The patient has tried and had an inadequate response to TWO agents
	from Step 1 and/or Step 2 for the requested indication after at least a 3-
	month trial per agent (See Step 3b) OR
	2. The patient has an intolerance (defined as an intolerance to the drug or
	its excipients, not to the route of administration) or hypersensitivity to
	TWO agents from Step 1 and/or Step 2 for the requested indication OR
	3. The patient has an FDA labeled contraindication to ALL of the Step 1 AND
	Step 2 agents for the requested indication OR 4. BOTH of the following:
	A. ALL of the Step 1 AND Step 2 agents are not clinically appropriate
	for the patient AND
	B. The prescriber has provided a complete list of previously tried
	agents for the requested indication OR
	5. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking
	the requested agent AND
	B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested agent AND
	C. The prescriber states that a change in therapy is expected to be
	ineffective or cause harm OR
	6. The prescriber has provided documentation that ALL of the Step 1 AND
	Step 2 agents for the requested indication cannot be used due to a
	documented medical condition or comorbid condition that is likely to
	cause an adverse reaction, decrease ability of the patient to achieve or
	maintain reasonable functional ability in performing daily activities or
	cause physical or mental harm OR
	G. If the requested agent is a Step 3c agent for the requested indication, then ONE of
	the following (chart notes required):
	 The patient has tried and had an inadequate response to THREE of the Step 1 agents for the requested indication after at least a 3-month trial
	per agent (See Step 3c) OR
	2. The patient has an intolerance (defined as an intolerance to the drug or
	its excipients, not to the route of administration) or hypersensitivity to
	THREE of the Step 1 agents for the requested indication OR
	3. The patient has an FDA labeled contraindication to ALL of the Step 1
	agents for the requested indication OR
	4. BOTH of the following:
	 A. ALL of the Step 1 agents are not clinically appropriate for the patient AND
	B. The prescriber has provided a complete list of previously tried
	agents for the requested indication OR
	5. The patient is currently being treated with the requested agent as
	indicated by ALL of the following:
	A. A statement by the prescriber that the patient is currently taking
	the requested agent AND B. A statement by the prescriber that the patient is currently
	receiving a positive therapeutics outcome on requested
	agent AND
	C. The prescriber states that a change in therapy is expected to be
	ineffective or cause harm OR
	menective of cause namin on

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	6. The prescriber has provided documentation that ALL of the Step 1 agents for the requested indication cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm AND
	 If Cosentyx 300 mg is requested as maintenance dosing, ONE of the following: A. The patient has a diagnosis of moderate to severe plaque psoriasis with or without coexistent active psoriatic arthritis AND the requested dose is 300 mg every 4 weeks OR B. The patient has a diagnosis of hidradenitis suppurativa AND ONE of the following:
	1. The requested dose is 300 mg every 4 weeks OR 2. The requested dose is 300 mg every 2 weeks AND the patient has tried and had an inadequate response to Cosentyx 300 mg every 4 weeks after at least a 3-month duration of therapy OR
	C. The patient has a diagnosis of active psoriatic arthritis or active ankylosing spondylitis AND has tried and had an inadequate response to Cosentyx 150 mg every 4 weeks after at least a 3-month duration of therapy AND
	 4. If Omvoh is requested for the treatment of ulcerative colitis, ONE of the following: A. The patient has received Omvoh IV for induction therapy OR B. The patient is new to therapy and will receive Omvoh IV for induction therapy AND
	 If Entyvio is requested for the treatment of ulcerative colitis, ONE of the following: A. The patient has received at least 2 doses of Entyvio IV therapy OR B. The patient is new to therapy and will receive at least 2 doses of Entyvio IV therapy AND
	 6. If Skyrizi is requested for the treatment of Crohn's disease, ONE of the following: A. The patient received Skyrizi IV for induction therapy OR B. The patient is new to therapy and will receive Skyrizi IV for induction therapy AND
	 7. If an ustekinumab product is requested for the treatment of Crohn's disease or ulcerative colitis, ONE of the following: A. The patient received an ustekinumab IV product for induction therapy OR B. The patient is new to therapy and will receive an ustekinumab IV product for induction therapy AND
	8. If Zymfentra is requested for the treatment of Crohn's disease or ulcerative colitis, then ONE of the following: A. The patient received an infliximab IV product for induction therapy OR B. The patient is new to therapy and will receive an infliximab IV product for
	induction therapy AND
	 9. If the patient has an FDA labeled indication, then ONE of the following: A. The patient's age is within FDA labeling for the requested indication for the requested agent OR
	 B. There is support for using the requested agent for the patient's age for the requested indication AND
	4. If an ustekinumab 90 mg product is requested, then ONE of the following:A. The patient has a diagnosis of psoriasis AND weighs >100kg OR
	B. The patient has a dual diagnosis of psoriasis AND psoriatic arthritis AND the patient is >100kg OR
	 C. The patient has a diagnosis of Crohn's disease or ulcerative colitis AND 5. If Actemra is requested for a diagnosis of systemic sclerosis associated interstitial lung disease, the request is
	for the Actemra syringe (NOTE: Actemra ACTpen is not approvable for SSc-ILD) AND
	6. The prescriber is a specialist in the area of the patient's diagnosis (e.g., rheumatologist for JIA, PsA, RA; gastroenterologist for CD, UC; dermatologist for PS, AD; pulmonologist, radiologist, pathologist,

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rheumatologist for SSc-ILD; allergist, immunologist for AD) or has consulted with a specialist in the area of the patient's diagnosis **AND**

- 7. ONE of the following (Please refer to "Agents NOT to be used Concomitantly" table):
 - A. The patient will NOT be using the requested agent in combination with another immunomodulatory agent (e.g., TNF inhibitors, JAK inhibitors, IL-4 inhibitors) **OR**
 - B. The patient will be using the requested agent in combination with another immunomodulatory agent AND BOTH of the following:
 - 1. The prescribing information for the requested agent does NOT limit the use with another immunomodulatory agent **AND**
 - 2. There is support for the use of combination therapy (copy of support required, i.e., clinical trials, phase III studies, guidelines) **AND**
- 8. The patient does NOT have any FDA labeled contraindications to the requested agent AND
- 9. The patient has been tested for latent tuberculosis (TB) when required by the prescribing information for the requested agent AND if positive the patient has begun therapy for latent TB

Compendia Allowed: AHFS, DrugDex 1 or 2a level of evidence, or NCCN 1 or 2a recommended use

Length of Approval: 12 months for all agents EXCEPT adalimumab containing products for ulcerative colitis (UC), Rinvoq for atopic dermatitis (AD), Siliq for plaque psoriasis (PS), Xeljanz and Xeljanz XR for induction therapy for UC, and the agents with indications that require loading doses for new starts. NOTE: For agents that require a loading dose for a new start, approve the loading dose based on FDA labeling AND the maintenance dose for the remainder of the 12 months. Adalimumab containing products for UC may be approved for 12 weeks, Rinvoq for AD may be approved for 6 months, Siliq for PS may be approved for 16 weeks, and Xeljanz and Xeljanz XR for UC may be approved for 16 weeks.

**NOTE: Cosentyx for the diagnoses of AS, nr-axSpA, and PSA loading doses are not approvable.

NOTE: If Quantity Limit applies, please refer to Quantity Limit Criteria.

Renewal Evaluation

Target Agent(s) will be approved when ALL of the following are met:

- The request is NOT for use of Olumiant or Actemra in the treatment of coronavirus disease 2019 (COVID-19)
 in hospitalized adults requiring supplemental oxygen, non-invasive or invasive mechanical ventilation, or
 extracorporeal membrane oxygenation (ECMO) *NOTE: This indication is not covered under the pharmacy
 benefit AND
- 2. The request is for use in Alopecia Areata and Alopecia Areata is NOT restricted from coverage under the patient's benefit **AND**
- 3. The patient has been previously approved for the requested agent through the plan's Prior Authorization process (*please note ustekinumab product renewal must be for the same strength as the initial approval) [Note: patients not previously approved for the requested agent will require initial evaluation review] **AND**
- 4. ONE of the following:
 - A. The patient has a diagnosis of moderate to severe atopic dermatitis AND BOTH of the following:
 - 1. The patient has had a reduction or stabilization from baseline (prior to therapy with the requested agent) of ONE of the following:
 - A. Affected body surface area OR
 - B. Flares **OR**

Module	Clinical Criteria for Approval
	C. Pruritus, erythema, edema, xerosis, erosions/excoriations, oozing and crusting, and/or lichenification OR
	D. A decrease in the Eczema Area and Severity Index (EASI) score OR
	E. A decrease in the Investigator Global Assessment (IGA) score AND
	2. The patient will continue standard maintenance therapies (e.g., topical emollients, good
	skin care practices) in combination with the requested agent OR
	B. The patient has a diagnosis of polymyalgia rheumatica AND BOTH of the following:
	1. The patient has had clinical benefit with the requested agent AND
	2. If the requested agent is Kevzara, the patient does NOT have any of the following:
	 A. Neutropenia (ANC less than 1,000 per mm³ at the end of the dosing interval) AND
	B. Thrombocytopenia (platelet count is less than 100,000 per mm^3) AND
	C. AST or ALT elevations 3 times the upper limit of normal OR
	C. The patient has a diagnosis other than moderate to severe atopic dermatitis or polymyalgia
	rheumatica AND the patient has had clinical benefit with the requested agent AND
	5. The prescriber is a specialist in the area of the patient's diagnosis (e.g., rheumatologist for JIA, PsA, RA;
	gastroenterologist for CD, UC; dermatologist for PS, AD; pulmonologist, radiologist, pathologist,
	rheumatologist for SSc-ILD; allergist, immunologist for AD) or the prescriber has consulted with a specialist in
	the area of the patient's diagnosis AND 6. ONE of the following (Please refer to "Agents NOT to be used Concomitantly" table):
	1. The patient will NOT be using the requested agent in combination with another immunomodulatory
	agent (e.g., TNF inhibitors, JAK inhibitors, IL-4 inhibitors) OR
	2. The patient will be using the requested agent in combination with another immunomodulatory
	agent AND BOTH of the following:
	1. The prescribing information for the requested agent does NOT limit the use with another
	immunomodulatory agent AND
	2. There is support for the use of combination therapy (copy of support required, i.e., clinical
	trials, phase III studies, guidelines) AND
	7. If Cosentyx 300 mg is requested as maintenance dosing, ONE of the following:
	A. The patient has a diagnosis of moderate to severe plaque psoriasis with or without coexistent active
	psoriatic arthritis AND the requested dose is 300 mg every 4 weeks OR
	B. The patient has a diagnosis of hidradenitis suppurativa AND ONE of the following:
	 The requested dose is 300 mg every 4 weeks OR
	2. The requested dose is 300 mg every 2 weeks AND the patient has tried and had an
	inadequate response to Cosentyx 300 mg every 4 weeks after at least a 3-month duration
	of therapy OR
	C. The patient has a diagnosis of active psoriatic arthritis or active ankylosing spondylitis AND has tried
	and had an inadequate response to Cosentyx 150 mg every 4 weeks after at least a 3-month
	duration of therapy AND
	8. If Actemra is requested for a diagnosis of systemic sclerosis associated interstitial lung disease, the request is
	for the Actemra syringe (NOTE: Actemra ACTpen is not approvable for SSc-ILD) AND
	9. The patient does NOT have any FDA labeled contraindications to the requested agent
	Compendia Allowed: AHFS, DrugDex 1 or 2a level of evidence, or NCCN 1 or 2a recommended use
	Length of Approval: 12 months
	**NOTE: Cosentyx for the diagnoses of AS, nr-axSpA, and PSA loading doses are not approvable.
	NOTE: If Quantity Limit applies, please refer to Quantity Limit Criteria.

Module	Clinical Criteria for Approval
QL All	Quantity limit for the Target Agent(s) will be approved when ONE of the following is met:
Program	
Type	1. The requested quantity (dose) does NOT exceed the program quantity limit OR
	2. The requested quantity (dose) exceeds the program quantity limit AND ONE of the following:
	A. The requested agent is Xeljanz/Xeljanz XR for a diagnosis of ulcerative colitis, AND BOTH of the
	following:
	 There is support for therapy for the dose exceeding the quantity limit [e.g., patient has lost response to the FDA labeled maintenance dose (i.e., 5 mg twice daily or 11 mg once daily) during maintenance treatment; requires restart of induction therapy] (medical records required) AND
	2. The requested quantity (dose) cannot be achieved with a lower quantity of a higher
	strength and/or package size that does not exceed the program quantity limit OR
	B. The requested agent is Xeljanz oral solution for a diagnosis of polyarticular course juvenile idiopathic arthritis, AND ONE of the following:
	1. BOTH of the following:
	 A. The requested quantity (dose) does not exceed the maximum FDA labeled dose (i.e., 5 mg twice daily) NOR the maximum compendia supported dose AND B. There is support why the patient cannot take Xeljanz 5 mg tablets OR
	2. The requested quantity (dose) exceeds the maximum FDA labeled dose but does NOT
	exceed the maximum compendia supported dose for the requested indication OR
	3. BOTH of the following:
	A. The requested quantity (dose) exceeds the maximum FDA labeled dose AND the
	maximum compendia supported dose for the requested indication AND
	B. There is support for therapy with a higher dose or shortened dosing interval for
	the requested indication (submitted copy of clinical trials, phase III studies, guidelines required) OR
	C. The requested agent is NOT Xeljanz/Xeljanz XR for a diagnosis of ulcerative colitis or polyarticular
	course juvenile idiopathic arthritis, AND ONE of the following:
	1. The patient has an FDA labeled indication for the requested agent, AND ONE of the
	following:
	A. BOTH of the following:
	The requested quantity (dose) does NOT exceed the maximum FDA Indicated the seasons.
	labeled dose AND 2. The requested quantity (dose) cannot be achieved with a lower quantity
	of a higher strength and/or package size that does NOT exceed the
	program quantity limit OR
	B. ALL of the following:
	 The requested quantity (dose) exceeds the FDA maximum labeled dose AND
	2. The patient has tried and had an inadequate response to at least a 3
	month duration of therapy at the maximum FDA labeled dose (medical
	records required) AND
	3. ONE of the following:
	A. BOTH of the following: 1. The requested quantity (dose) does NOT exceed the
	maximum compendia supported dose for the requested indication AND
	2. The requested quantity (dose) cannot be achieved with
	a lower quantity of a higher strength/and or package
	size that does NOT exceed the program quantity limit
	OR .
	B. BOTH of the following:

odule	Clinical Criteria for Approval
	 The requested quantity (dose) exceeds the maximum FDA labeled dose AND the maximum compendia supported dose for the requested indication AND There is support for therapy with a higher dose or shortened dosing interval for the requested indication (submitted copy of clinical trials, phase III studies, guidelines required) OR
	The patient has a compendia supported indication for the requested agent, AND ONE of the following:
	A. BOTH of the following:
	 The requested quantity (dose) does NOT exceed the maximum
	compendia supported dose for the requested indication AND
	2. The requested quantity (dose) cannot be achieved with a lower quantity
	of a higher strength/and or package size that does NOT exceed the program quantity limit OR
	B. BOTH of the following:
	1. The requested quantity (dose) exceeds the maximum compendia
	supported dose for the requested indication AND
	 There is support for therapy with a higher dose or shortened dosing interval for the requested indication (submitted copy of clinical trials,
	phase III studies, guidelines required) OR
	3. The patient does NOT have an FDA labeled indication NOR a compendia supported
	indication for the requested agent AND BOTH of the following: A. The requested quantity (dose) cannot be achieved with a lower quantity of a
	higher strength and/or package size that does not exceed the program quantity limit AND
	 B. There is support for therapy with a higher dose or shortened dosing interval for the requested indication (submitted copy of clinical trials, phase III studies, guidelines required)
	Compendia Allowed: AHFS, DrugDex 1 or 2a level of evidence, or NCCN 1 or 2a recommended use
	Length of Approval:
	Initial Approval with PA: up to 12 months for all agents EXCEPT adalimumab containing products for ulcerative colitis (UC), Rinvoq for atopic dermatitis (AD), Siliq for plaque psoriasis (PS), Xeljanz and Xeljanz XR for induction therapy for UC, and the agents with indications that require loading doses for new starts. NOTE: For agents that require a loading dose for a new start, approve the loading dose based on FDA labeling AND the maintenance dose for the remainder of the 12 months. Adalimumab containing products for UC may be approved for up to 12 weeks, Rinvoq for AD may be approved for up to 6 months, Siliq for PS may be approved for up to 16 weeks, and Xeljanz and Xeljanz XR for UC may be approved for up to 16 weeks.
	Renewal Approval with PA: up to 12 months
	Standalone QL approval: up to 12 months or through the remainder of an existing authorization, whichever is shorter
	**NOTE: Cosentyx for the diagnoses of AS, nr-axSpA, and PSA loading doses are not approvable.

CONTRAINDICATION AGENTS

Contraindicated as Concomitant Therapy

Agents NOT to be used Concomitantly

Abrilada (adalimumab-afzb)

Actemra (tocilizumab)

Adalimumab

Adbry (tralokinumab-ldrm)

Amjevita (adalimumab-atto)

Arcalyst (rilonacept)

Avsola (infliximab-axxq)

Benlysta (belimumab)

Bimzelx (bimekizumab-bkzx)

Cibingo (abrocitinib)

Cimzia (certolizumab)

Cinqair (reslizumab)

Cosentyx (secukinumab)

Cyltezo (adalimumab-adbm)

Dupixent (dupilumab)

Enbrel (etanercept)

Entyvio (vedolizumab)

Fasenra (benralizumab)

Hadlima (adalimumab-bwwd)

Hulio (adalimumab-fkjp)

Humira (adalimumab)

Hyrimoz (adalimumab-adaz)

Idacio (adalimumab-aacf)

Ilaris (canakinumab)

Ilumya (tildrakizumab-asmn)

Inflectra (infliximab-dyyb)

Infliximab

Kevzara (sarilumab)

Kineret (anakinra)

Litfulo (ritlecitinib)

Nucala (mepolizumab)

Olumiant (baricitinib)

Omvoh (mirikizumab-mrkz)

Opzelura (ruxolitinib)

Orencia (abatacept)

Otezla (apremilast)

Remicade (infliximab)

Renflexis (infliximab-abda)

Riabni (rituximab-arrx)

Rinvoq (upadacitinib)

Rituxan (rituximab)

Rituxan Hycela (rituximab/hyaluronidase human)

Ruxience (rituximab-pvvr)

Siliq (brodalumab)

Simlandi (adalimumab-ryvk)

Simponi (golimumab)

Simponi ARIA (golimumab)

Skyrizi (risankizumab-rzaa)

Sotyktu (deucravacitinib)

Spevigo (spesolimab-sbzo)

Contraindicated as Concomitant Therapy

Stelara (ustekinumab)

Taltz (ixekizumab)

Tezspire (tezepelumab-ekko)

Tofidence (tocilizumab-bavi)

Tremfya (guselkumab)

Truxima (rituximab-abbs)

Tyenne (tocilizumab-aazg)

Tysabri (natalizumab)

Velsipity (etrasimod)

Wezlana (ustekinumab-auub)

Xeljanz (tofacitinib)

Xeljanz XR (tofacitinib extended release)

Xolair (omalizumab)

Yuflyma (adalimumab-aaty)

Yusimry (adalimumab-aqvh)

Zeposia (ozanimod)

Zymfentra (infliximab-dyyb)