

PREFERRED SPECIALTY MANAGEMENT POLICY

POLICY: Inflammatory Conditions Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies

Tumor Necrosis Factor Inhibitors

- Adalimumab Products³
 - adalimumab-aaty subcutaneous injection (Alvotech/Teva)
 - o adalimumab-adaz subcutaneous injection (Sandoz/Novartis)
 - o adalimumab-adbm subcutaneous injection (Boehringer Ingelheim)
 - Cyltezo® (adalimumab-adbm subcutaneous injection Boehringer Ingelheim)
 - Humira[®] (adalimumab subcutaneous injection AbbVie, Cordavis)
 - Hyrimoz[®] (adalimumab-adaz subcutaneous injection Sandoz/Novartis, Cordavis)
 - Simlandi (adalimumab-rykv subcutaneous injection Alvotech/Teva)
- Cimzia® (certolizumab pegol subcutaneous injection UCB)
- Enbrel® (etanercept subcutaneous injection Amgen)
- Simponi® (golimumab subcutaneous injection Janssen Biotech/Johnson & Johnson)
- Zymfentra® (infliximab-dyyb subcutaneous injection Celltrion)

Interleukin-6 Blockers

- Tocilizumab Subcutaneous Products
 - o Actemra® (tocilizumab subcutaneous injection Genentech/Roche)
 - o Tyenne® (tocilizumab-aazg subcutaneous injection Fresenius Kabi)
- Kevzara[™] (sarilumab subcutaneous injection Regeneron)

Interleukin-17 Blockers

- Bimzelx® (bimekizumab subcutaneous injection UCB)
- Cosentyx® (secukinumab subcutaneous injection Novartis)
- Siliq[™] (brodalumab subcutaneous injection Valeant)
- Taltz® (ixekizumab subcutaneous injection Eli Lilly)

Interleukin-23 Blockers

- Ilumya[™] (tildrakizumab-asmn subcutaneous injection Sun/Merck)
- Omvoh® (mirakizumab-mrkz subcutaneous injection Eli Lilly)
- Skyrizi[™] (risankizumab-rzaa subcutaneous injection AbbVie)
- Tremfya[™] (guselkumab subcutaneous injection Janssen Biotech/Johnson & Johnson)

Interleukin 12/23 Blocker

• Stelara® (ustekinumab subcutaneous injection – Janssen Biotech/Johnson & Johnson)

Interleukin-1 Blocker

Kineret[®] (anakinra subcutaneous injection – Swedish Orphan Biovitrim)

T-Cell Costimulation Modulator

Orencia® (abatacept subcutaneous injection – Bristol Myers Squibb)

Integrin Receptor Antagonist

• Entyvio® (vedolizumab subcutaneous injection - Takeda)

Janus Kinases Inhibitors

- Olumiant® (baricitinib tablets Eli Lilly)
- Rinvoq[™] (upadacitinib extended-release tablets AbbVie)
- Rinvog® LO (upadacitinib oral solution AbbVie)
- Xeljanz® (tofacitinib tablets, tofacitinib oral solution Pfizer)
- Xeljanz® XR (tofacitinib extended-release tablets Pfizer)

Phosphodiesterase Type 4 Inhibitor

Otezla® (apremilast tablets – Amgen)

Sphingosine 1-Phosphate Receptor Modulator

- Velsipity[™] (etrasimod tablets Pfizer)
- Zeposia® (ozanimod capsules Celgene)

Tyrosine Kinase 2 Inhibitor

Sotyktu[™] (deucravacitinib tablets – Bristol Myers Squibb)

* For Non-Preferred adalimumab products, refer to the *Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policies for National Preferred, High Performance, and Basic Formularies* or the Choice version of this policy.

REVIEW DATE: 11/22/2023; selected revision 01/03/2024, 01/24/2024,

02/28/2024, 05/08/2024, 05/15/2024, 07/17/2024, 10/02/2024

INSTRUCTIONS FOR USE

The following Coverage Policy applies to health benefit plans administered by Cigna Companies. Certain Cigna Companies and/or lines of business only provide utilization review services to clients and do not make coverage determinations. References to standard benefit plan language and coverage determinations do not apply to those clients. Coverage Policies are intended to provide guidance in interpreting certain standard benefit plans administered by Cigna Companies. Please note, the terms of a customer's particular benefit plan document [Group Service Agreement, Evidence of Coverage, Certificate of Coverage, Summary Plan Description (SPD) or similar plan document] may differ significantly from the standard benefit plans upon which these Coverage Policies are based. For example, a customer's benefit plan document may contain a specific exclusion related to a topic addressed in a Coverage Policy. In the event of a conflict, a customer's benefit plan document always supersedes the information in the Coverage Policies. In the absence of a controlling federal or state coverage mandate, benefits are ultimately determined by the terms of the applicable benefit plan document. Coverage determinations in each specific instance require consideration of 1) the terms of the applicable benefit plan document in effect on the date of service; 2) any applicable laws/regulations; 3) any relevant collateral source materials including Coverage Policies and; 4) the specific facts of the particular situation. Each coverage request should be reviewed on its own merits. Medical directors are expected to exercise clinical judgment and have discretion in making individual coverage determinations. Coverage Policies relate exclusively to the administration of health benefit plans. Coverage Policies are not recommendations for treatment and should never be used as treatment guidelines. In certain markets, delegated vendor guidelines may be used to support medical necessity and other coverage determinations.

Cigna National Formulary Coverage:

OVERVIEW

Several products are available for use in inflammatory conditions such as rheumatoid arthritis, ankylosing spondylitis, juvenile idiopathic arthritis, psoriatic arthritis, plaque psoriasis, Crohn's disease, and ulcerative colitis. This policy involves the use of the products listed above.

The FDA-approved indications for each product listed in this policy are documented in <u>Appendix A</u>. For more information on criteria within a Prior Authorization program by specific condition refer to the respective standard *Prior Authorization Policy*.

POLICY STATEMENT

For all medications, this program requires the patient to meet the respective standard *Prior Authorization Policy* criteria. Additionally, this program requires trial(s) of the Preferred Product(s) according to the table below, when clinically appropriate, prior to the approval of the Non-Preferred Products. There are also situations when trials of Non-Preferred Products will be considered; see criteria below. Other details of the program are as follows:

• **Continuation of Therapy:** Approval for a patient <u>continuing therapy with a</u> **Non-Preferred** subcutaneous or oral Product must be supported with

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verification, noted in the criteria as either [verification in prescription claims history required] or, if not available, as [verification by prescriber required].

- If the patient has at least 130 days of prescription claims history on file, claims history must support that the patient has received the Non-Preferred Product for the specified period of time (90 or 120 days) within a 130-day look-back period; OR
- When 130 days of the patient's prescription claim history file is unavailable for verification, the prescriber must verify that the patient has been receiving the Non-Preferred Product for a specified period of time (90 or 120 days), AND that the patient has been receiving the Non-Preferred Product via paid claims (e.g., patient has <u>not</u> been receiving samples or coupons or other types of waivers in order to obtain access to the Non-Preferred Product).
- o For a patient continuing therapy, other conditions may also apply. Refer to criteria below.
- **Approval Duration:** All approvals for continuation of therapy for Preferred and Non-Preferred Products are provided for 1 year unless noted otherwise below. In cases where the initial approval is authorized in months, 1 month is equal to 30 days.

<u>Documentation</u>: When documentation is required, the prescriber must provide written documentation supporting the trials of these other Products, noted in the criteria as **[documentation required]**. Documentation may include, but is not limited to, chart notes, prescription claims records, and/or prescription receipts.

Preferred and Non-Preferred Products.¥

	Rheumatology					Dermatology	Gastroe	nterology
	RA	JIA	AS	nr-axSpA	PsA	Psoriasis	CD	UC
Step 1	• Enbrel	• Enbrel	• Enbrel	• Cimzia	• Enbrel	• Enbrel	 Adalimum 	 Adalimuma
Preferred	 Adalimum 	 Adalimum 	 Adalimum 	Taltz	 Adalimuma 	 Adalimumab 	ab	b
	ab	ab	ab		b	Products^ -	Products [^]	Products [^]
	Products [^]	Products [^]	Products [^]		Products [^]	Humira	– Humira	– Humira
	– Humira	– Humira	– Humira		– Humira	(NDCs starting	(NDCs	(NDCs starting
	(NDCs	(NDCs	(NDCs		(NDCs starting	with <u>00074</u>),	starting with	with <u>00074</u>),
	starting with	starting with	starting with		with <u>00074</u>),	Cyltezo/	<u>00074</u>),	Cyltezo/
	<u>00074</u>),	<u>00074</u>),	<u>00074</u>) ,		Cyltezo/	adalimumab	Cyltezo/	adalimuma
	Cyltezo/	Cyltezo/	Cyltezo/		adalimuma	-adbm,	adalimum	b-adbm,
	adalimum	adalimuma	adalimum		b-adbm,	Hyrimoz	ab-adbm,	Hyrimoz
	ab-adbm,	b-adbm,	ab-adbm,		Hyrimoz	(NDCs starting	Hyrimoz	(NDCs starting
	Hyrimoz	Hyrimoz	Hyrimoz		(NDCs starting	with <u>61314</u>)/	(NDCs	with <u>61314</u>)/
	(NDCs	(NDCs	(NDCs		with <u>61314</u>)/	adalimumab	starting with	adalimuma
	starting with	starting with	starting with		adalimuma	-adaz,	61314)/	b-adaz,
	61314)/ adalimum	61314)/ adalimuma	<u>61314</u>)/		b-adaz,	Simlandi/	adalimum	Simlandi/
	ab-adaz,	b-adaz,	adalimum		Simlandi/	adalimumab	ab-adaz,	adalimuma
			ab-adaz,		adalimuma	-ryvk	Simlandi/	b-ryvk
	Simlandi/	Simlandi/	Simlandi/		b-ryvk	Otezla	adalimum	Skyrizi SC
	adalimum	adalimuma	adalimum		Otezla	Skyrizi SC#	ab-ryvk	(on-body
	ab-ryvk	b-ryvk	ab-ryvk		Skyrizi SC#	Sotyktu	 Skyrizi SC 	injector)
			• Taltz		Stelara SC	 Stelara SC 	(on-body	 Stelara SC
					Taltz	Taltz	injector)	Zymfentra
					Tremfya	Tremfya SC	 Stelara SC 	-
					•	•	 Zymfentra 	
Step 2			Rinvoq	Rinvoq	Rinvoq/		Cimzia	Omvoh SC
Non-	ab SC	SC Products	Directed	Directed	Rinvoq LQ			Rinvoq
Preferred	Products	- Actemra	specifically to	specifically	Directed		adalimumab	Directed to
	- Actemra	SC, Tyenne		to Cimzia.	specifically to		specifically.	

(directed to ONE Step 1 Product)	SC, Tyenne SC Directed to adalimumab specifically. •Rinvoq •Xeljanz tablets/ Xeljanz XR tablets	SC Directed to adalimumab specifically. JIA Step SC is for PJIA. •Rinvoq/Rinvoq LQ •Xeljanz tablets/ Xeljanz oral solution	Enbrel or adalimumab. • Xeljanz tablets/ Xeljanz XR tablets Directed specifically to Enbrel or adalimumab.		Enbrel or adalimumab. • Xeljanz tablets/ Xeljanz XR tablets Directed specifically to Enbrel or adalimumab.		• Rinvoq Directed to adalimumab specifically.	adalimumab specifically. Simponi SC Directed to adalimumab specifically. Xeljanz tablets/ Xeljanz/ XR tablets Directed to adalimumab specifically.
Step 2b Non- Preferred (directed to ONE Step 1 Product)						• Bimzelx		specifically.
Step 3a Non- Preferred (directed to TWO Step 1 or 2a Products) [documenta tion required]*	Cimzia Kevzara Kineret Olumiant Orencia SC Simponi SC	• Kevzara • Orencia SC	• Cimzia • Cosentyx SC • Simponi SC	•Cosentyx SC	• Cimzia • Cosentyx SC • Orencia SC • Simponi SC	• Cimzia • Cosentyx SC • Ilumya • Siliq	• Entyvio SC	• Entyvio SC
Step 3b Non- Preferred (directed to TWO Step 1								• Zeposia Refer to MS and UC – Zeposia PSM Policy

		R	heumatology	Dermatology	Gastroe	nterology		
	RA	JIA	AS	nr-axSpA	PsA	Psoriasis	CD	UC
Step 4 Non- Preferred (directed to								Velsipity
TWO Step 1 or 2 Products AND ONE Step 3b Product) [documenta tion								

For Non-Preferred Products, refer to the *Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies* or the Choice version of that policy. Note that adalimumab-adaz, adalimumab-adbm, and Simlandi/adalimumab-ryvk are Non-Preferred for some plans; RA – Rheumatoid arthritis; ^ A trial of more than one adalimumab product counts as ONE Preferred Product; JIA – Juvenile idiopathic arthritis; AS – Ankylosing spondylitis; nr-axSpA – Nonradiographic axial spondyloarthritis; PSA – Products; CD – Crohn's disease; UC – Ulcerative colitis; SC – Subcutaneous; * Pen and syringe; PJIA – Polyarticular juvenile idiopathic arthritis; * The prescriber must provide written documentation supporting the trial of Preferred Products, noted in the criteria as [documentation required]. Documentation may include, but is not limited to, chart notes, prescription claims records, and/or prescription receipts; PSM – Preferred Specialty Management.

Inflammatory Conditions Preferred Specialty Management Policy non-preferred product(s) is(are) covered as medically necessary when the following non-preferred product exception criteria is(are) met. Any other exception is considered not medically necessary.

Non-Preferred Product Exception Criteria

Non-	Exception Criteria
Preferred	
Product	
Tumor Necre	osis Factor Inhibitors
Cimzia	1. Rheumatoid Arthritis - Initial Therapy.
	A) Approve for 6 months if the patient meets BOTH of the
	following (i <u>and</u> ii):
	i. Patient meets the standard <i>Inflammatory Conditions</i> –
	Cimzia Prior Authorization Policy criteria; AND
	ii. Patient has tried TWO of a tocilizumab subcutaneous
	product, Enbrel, an adalimumab product, Rinvoq, or
	Xeljanz/XR [documentation required].
	Note: Examples of tocilizumab subcutaneous products
	include Actemra subcutaneous and Tyenne subcutaneous.
	A trial of multiple tocilizumab products counts as ONE
	product. Examples of adalimumab products include
	Humira, Abrilada, adalimumab-adaz, adalimumab-adbm,
	adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,
	Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,
	Idacio, Yuflyma, and Yusimry. A trial of multiple
	adalimumab products counts as ONE product. A trial of
	either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as ONE product.
	B) If the patient has met criterion 1Ai (the standard
	Inflammatory Conditions – Cimzia Prior Authorization Policy
	criteria), but criterion 1Aii is not met: offer to review for a
	Step 1 or Step 2 Product (<u>Actemra subcutaneous, Tyenne</u>
	subcutaneous, Enbrel, Humira [NDCs starting with 00074],
	adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with
	61314], adalimumab-adaz, adalimumab-ryvk, Simlandi,
	Rinvog, Xeljanz tablets, or Xeljanz XR) using the respective
	standard Inflammatory Conditions Prior Authorization Policy
	criteria.
	2. Ankylosing Spondylitis – Initial Therapy.
	A) Approve for 6 months if the patient meets BOTH of the
	following (i <u>and</u> ii):
	i. Patient meets the standard Inflammatory Conditions –
	Cimzia Prior Authorization Policy criteria; AND

- ii. Patient has tried TWO of Enbrel, an adalimumab product, Rinvoq, Taltz, or Xeljanz/XR [documentation required].
 - Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product.
- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Cimzia Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Step 1 or Step 2 Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Taltz, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

3. Psoriatic Arthritis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Cimzia Prior Authorization Policy* criteria; AND
 - ii. Patient has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvoq/Rinvoq LQ, Skyrizi subcutaneous, Stelara subcutaneous, Taltz, Tremfya, and Xeljanz/XR [documentation required].
 - Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, adalimumab-adbm, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product.
- B) If the patient has met criterion 3Ai (the standard Inflammatory Conditions Cimzia Prior Authorization Policy criteria), but criterion 3Aii is not met: offer to review for a Step 1 or Step 2 Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Rinvoq, Rinvoq LQ, Skyrizi subcutaneous [pen or syringe], Stelara subcutaneous, Taltz, Tremfya,

<u>Xeljanz tablets, or Xeljanz XR</u>) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

4. Plaque Psoriasis - Initial Therapy.

- **A)** Approve for 3 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Cimzia Prior Authorization Policy* criteria; AND
 - ii. Patient has tried TWO of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya [documentation required].

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, adalimumab-adbm, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product.

B) If the patient has met criterion 4Ai (the standard Inflammatory Conditions – Cimzia Prior Authorization Policy criteria), but criterion 4Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu, Stelara subcutaneous, Taltz, or Tremfya) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

5. Crohn's Disease - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Cimzia Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one adalimumab product.

 Note: Examples of adalimumab products include Humira,
 Abrilada, adalimumab-adaz, adalimumab-fkjp,
 adalimumab-aaty, adalimumab-ryvk, Simlandi,
 adalimumab-adbm, Amjevita, Cyltezo, Hadlima, Hulio,
 Hyrimoz, Idacio, Yuflyma, and Yusimry.
- B) If the patient has met criterion 5Ai (the standard Inflammatory Conditions Cimzia Prior Authorization Policy criteria), but criterion 5Aii is not met: offer to review for a Preferred Product (Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous [on-body injector], Stelara subcutaneous, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

- 6. Rheumatoid Arthritis, Ankylosing Spondylitis, Psoriatic Arthritis, Plaque Psoriasis, or Crohn's Disease Patient is Currently Receiving Cimzia.
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Cimzia Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following (a, b, c, d, e, or f):
 - a) Patient has Rheumatoid Arthritis and has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq, or Xeljanz/XR [documentation required]; OR
 Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as ONE product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as ONE product. A trial of either or both
 - b) Patient has Ankylosing Spondylitis and has tried TWO of Enbrel, an adalimumab product, Rinvoq, Taltz, or Xeljanz/XR [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as ONE product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as ONE product.

Xeljanz products (Xeljanz and Xeljanz XR) collectively

c) Patient has <u>Psoriatic Arthritis</u> and has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvoq/Rinvoq LQ, Skyrizi subcutaneous, Stelara subcutaneous, Taltz, Tremfya, or Xeljanz/XR [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both

counts as **ONE** product.

- Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product.
- d) Patient has <u>Plaque Psoriasis</u> and has tried TWO of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumabfkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, adalimumab-adbm, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product.
- e) Patient has <u>Crohn's Disease</u> and has tried one adalimumab product; OR

 <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry.
- f) Patient has been established on Cimzia for at least 90 days <u>and</u> prescription claims history indicates <u>at least a 90-day supply of Cimzia was dispensed within the past 130 days</u> [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Cimzia for at least 90 days AND the patient has been receiving Cimzia via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Cimzia).

- **B)** If the patient has met criterion 6Ai (the standard *Inflammatory Conditions Cimzia Prior Authorization Policy* criteria), but criterion 6Aii is not met: offer to review for one of the following Products using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria:
 - i. Rheumatoid Arthritis: Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvog, Xeljanz tablets, or Xeljanz XR.

	ii. Ankylosing Spondylitis: Enbrel, Humira (NDCs starting
	with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs
	starting with 61314), adalimumab-adaz, adalimumab-
	ryvk, Simlandi, Rinvoq, Taltz, Xeljanz tablets, or Xeljanz
	XR.
	iii. Psoriatic Arthritis: Enbrel, Humira (NDCs starting with
	00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs
	starting with 61314), adalimumab-adaz, adalimumab-
	ryvk, Simlandi, Otezla, Rinvog, Rinvog LQ, Skyrizi
	subcutaneous (pen or syringe), Stelara subcutaneous,
	Taltz, Tremfya, Xeljanz tablets, or Xeljanz XR.
	iv. Plaque Psoriasis: Enbrel, Humira (NDCs starting with
	00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs
	starting with 61314), adalimumab-adaz, adalimumab-
	ryvk, Simlandi, Otezla, Skyrizi subcutaneous (pen or
	syringe), Sotyktu, Stelara subcutaneous, Taltz, or
	Tremfya.
	v. Crohn's Disease: Humira (NDCs starting with 00074),
	adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with
	61314), adalimumab-adaz, adalimumab-ryvk, Simlandi,
	Skyrizi subcutaneous (on-body injector), Stelara
	subcutaneous, or Zymfentra.
	7. Other Conditions. Approve Cimzia (initial therapy for a
	duration as directed or <u>1 year</u> for a patient continuing therapy)
	if the patient meets the standard <i>Inflammatory Conditions</i> –
	Cimzia Prior Authorization Policy criteria.
Enbrel	All Conditions. Approve Enbrel (initial therapy for a duration as
	directed or <u>1 year</u> for a patient continuing therapy) if the patient
	meets the standard <i>Inflammatory Conditions – Enbrel Prior</i>
	Authorization Policy criteria.
Humira	All Conditions. Approve (initial therapy for a duration as directed
(NDCs starting	or <u>1 year</u> for a patient continuing therapy) if the patient meets the
with 00074)	standard Inflammatory Conditions – Adalimumab Products Prior
Adalimuma	Authorization Policy criteria.
b-adaz	Note: Adalimumab-adaz, adalimumab-adbm, and
Adalimuma	Simlandi/adalimumab-ryvk Non-Preferred for some plans. Refer to
b-adbm	respective Inflammatory Conditions – Adalimumab Products
Cyltezo	Preferred Specialty Management Policy for National Preferred, High
Hyrimoz	Performance, and Basic Formularies Policies or the Choice version
(NDCs starting	of that policy.
with 61314) Simlandi	r/
adalimuma	
b-ryvk	4 Dhamasaid Authritis Tuitis Theorem
Simponi	1. Rheumatoid Arthritis – Initial Therapy.
Subcutane	A) Approve for 6 months if the patient meets BOTH of the
ous	following (i <u>and</u> ii):

- Patient meets the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria; AND
- ii. Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq, or Xeljanz/XR [documentation required]; OR Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as ONE product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as ONE product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as ONE product.
- B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. Ankylosing Spondylitis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria; AND
 - ii. Patient has tried TWO of Enbrel, an adalimumab product, Rinvoq, Taltz, or Xeljanz/XR [documentation required].
 - Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product.
- **B)** If the patient has met criterion 2Ai (the standard *Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy* criteria), but criterion 2Aii is not met: offer to review for a Step 1 or Step 2 Product (Enbrel,

Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Taltz, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria.

3. Psoriatic Arthritis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i and ii):
 - Patient meets the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria; AND
 - ii. Patient has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvoq/Rinvoq LQ, Skyrizi subcutaneous, Stelara subcutaneous, Taltz, Tremfya, or Xeljanz/XR [documentation required].

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product.

B) If the patient has met criterion 3Ai (the standard Inflammatory Conditions – Simponi Subcutaneous Prior Authorization Policy criteria), but criterion 3Aii is not met: offer to review for a Step 1 or Step 2 Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Rinvoq, Rinvoq LQ, Skyrizi subcutaneous [pen or syringe], Stelara subcutaneous, Taltz, Tremfya, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria.

4. <u>Ulcerative Colitis – Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria; AND
 - **ii.** Patient has tried one adalimumab product.

 <u>Note</u>: Examples of adalimumab products include Humira,
 Abrilada, adalimumab-adaz, adalimumab-adbm,
 adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,
 Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,
 Idacio, Yuflyma, and Yusimry.

- B) If the patient has met criterion 4Ai (the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria), but criterion 4Aii is not met: offer to review for a Preferred Product (Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 5. Rheumatoid Arthritis, Ankylosing Spondylitis, Psoriatic Arthritis, or Ulcerative Colitis Patient is Currently Receiving Simponi Subcutaneous or Aria.
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria; AND
 - **ii.** Patient meets ONE of the following (a, b, c, d, e, or f):
 - a) Patient has <u>Rheumatoid Arthritis</u> and has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq, or Xeljanz/XR [documentation required]; OR
 - Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product.
 - b) Patient has Ankylosing Spondylitis and has tried TWO of Enbrel, an adalimumab product, Rinvoq, Taltz, or Xeljanz/XR [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as ONE product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as ONE product.

- c) Patient has <u>Psoriatic Arthritis</u> and has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvoq/Rinvoq LQ, Skyrizi subcutaneous, Stelara subcutaneous, Taltz, Tremfya, or Xeljanz/XR [documentation required]; OR

 <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively
- d) Patient has <u>Ulcerative Colitis</u> and has tried one adalimumab product; OR <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumabadbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry.

counts as **ONE** product.

- **e)** According to the prescriber, the patient has been established on Simponi Aria for at least 90 days; OR
- f) Patient has been established on Simponi subcutaneous for at least 90 days <u>and</u> prescription claims history indicates <u>at least a 90-day supply of Simponi subcutaneous was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].</u>

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Simponi subcutaneous for at least 90 days AND the patient has been receiving Simponi subcutaneous via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Simponi subcutaneous).

B) If the patient has met criterion 5Ai (the standard *Inflammatory Conditions – Simponi Subcutaneous Prior Authorization Policy* criteria), but criterion 5Aii is not met: offer to review for one of the following Products using the

respective standard *Inflammatory Conditions – Prior Authorization Policy* criteria:

- i. Rheumatoid Arthritis: Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR.
- ii. Ankylosing Spondylitis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Taltz, Xeljanz tablets, or Xeljanz XR.
- iii. Psoriatic Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Rinvoq, Rinvoq LQ, Skyrizi subcutaneous (pen or syringe), Stelara subcutaneous, Taltz, Tremfya, Xeljanz tablets, or Xeljanz XR.
- iv. Ulcerative Colitis: Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra.
- **6.** Other Conditions. Approve Simponi subcutaneous (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Simponi Subcutaneous Prior Authorization Policy criteria.

Zymfentra

<u>All Conditions</u>. Approve <u>Zymfentra</u> (initial therapy for a duration as directed or <u>1 year</u> for a patient continuing therapy) if the patient meets the standard <u>Inflammatory Conditions</u> – <u>Zymfentra Prior Authorization Policy</u> criteria.

Interleukin-6 Blockers

Actemra Subcutane ous Tyenne Subcutane ous

- 1. <u>Polyarticular Juvenile Idiopathic Arthritis Initial</u> Therapy.
 - **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Tocilizumab Subcutaneous Prior Authorization Policy criteria; AND
 - ii. Patient meets ONE of the following (a or b):
 - a) Patient has tried one adalimumab product; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and

- Yusimry. A trial of Enbrel, an infliximab product (e.g., Remicade, biosimilars), or Simponi Aria also counts.
- **b)** According to the prescriber, the patient has heart failure or a previously treated lymphoproliferative disorder.
- B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Tocilizumab Subcutaneous Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. Rheumatoid Arthritis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Tocilizumab Subcutaneous Prior Authorization Policy* criteria; AND
 - **ii.** Patient meets ONE of the following (a <u>or</u> b):
 - a) Patient has tried one adalimumab product; OR
 Note: Examples of adalimumab products include
 Humira, Abrilada, adalimumab-adaz, adalimumabadbm, adalimumab-fkjp, adalimumab-aaty,
 adalimumab-ryvk, Simlandi, Amjevita, Cyltezo,
 Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and
 Yusimry. A trial of Cimzia, Enbrel, an infliximab
 product (e.g., Remicade, biosimilars), or Simponi (Aria
 or subcutaneous) also counts.
 - **b)** According to the prescriber, the patient has heart failure or a previously treated lymphoproliferative disorder.
- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Tocilizumab Subcutaneous Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 3. Polyarticular Juvenile Idiopathic Arthritis or Rheumatoid Arthritis Patient is Currently Receiving Tocilizumab Subcutaneous or Intravenous.
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):

- i. Patient meets the standard *Inflammatory Conditions Tocilizumab Subcutaneous Policy* criteria; AND
- ii. Patient meets ONE of the following (a, b, c, d, or e):
 - a) Patient has <u>Polyarticular Juvenile Idiopathic Arthritis</u> and has tried one adalimumab product; OR <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Enbrel, an infliximab product (e.g., Remicade, biosimilars), or Simponi Aria also counts.
 - b) Patient has Rheumatoid Arthritis and has tried one adalimumab product; OR
 Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, Enbrel, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
 - c) According to the prescriber, the patient has heart failure or a previously treated lymphoproliferative disorder; OR
 - **d)** According to the prescriber, the patient has been established on tocilizumab intravenous for at least 90 days; OR
 - e) Patient has been established on tocilizumab subcutaneous for at least 90 days <u>and</u> prescription claims history indicates <u>at least a 90-day supply of tocilizumab subcutaneous was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].</u>

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving tocilizumab subcutaneous for at least 90 days AND the patient has been receiving tocilizumab subcutaneous via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to tocilizumab subcutaneous).

B) If the patient has met criterion 3Ai (the standard *Inflammatory Conditions – Tocilizumab Subcutaneous Prior*

Authorization Policy criteria), but criterion 3Aii is not met: offer to review for a Preferred Product using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria:

- i. Polyarticular Juvenile Idiopathic Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, or Simlandi.
- ii. Rheumatoid Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-rvyk, or Simlandi.
- **4.** <u>All Other Conditions</u> (including systemic juvenile idiopathic arthritis). Approve <u>tocilizumab subcutaneous</u> (initial therapy for a duration as directed or <u>1 year</u> for a patient continuing therapy) if the patient meets the standard *Inflammatory Conditions Tocilizumab Subcutaneous Prior Authorization Policy* criteria.

Kevzara

1. Rheumatoid Arthritis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Kevzara Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following (a or b):
 - a) Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvog, or Xeljanz/XR [documentation required]; OR Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xelianz products (Xelianz and Xelianz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, an infliximab product (e.g., Remicade, biosimilars), Orencia (intravenous or subcutaneous), or Simponi (Aria or subcutaneous) also counts [documentation required].
 - **b)** According to the prescriber, the patient has heart failure or a previously treated lymphoproliferative disorder.

B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions – Kevzara Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. <u>Juvenile Idiopathic Arthritis/Juvenile Rheumatoid</u> Arthritis – Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Kevzara Prior Authorization Policy* criteria; AND
 - **ii.** Patient meets ONE of the following conditions (a <u>or</u> b):
 - a) Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq/Rinvoq LQ, or Xeljanz [documentation required]; OR

Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Rinvog products (Rinvog and Rinvog LQ) collectively counts as **ONE** product. A trial of a tocilizumab intravenous product (Actemra intravenous, biosimilar), Orencia intravenous or subcutaneous, an infliximab product (e.g., Remicade, biosimilars), or Simponi Aria also counts [documentation required].

- **b)** According to the prescriber, the patient has heart failure, a previously treated lymphoproliferative disorder, a previous serious infection, OR a demyelinating disorder.
- **B)** If the patient has met criterion 2Ai (the standard Inflammatory Conditions –Kevzara Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with

61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Rinvoq LQ, or Xeljanz tablets) using the respective standard *Inflammatory Conditions – Prior Authorization Policy* criteria.

- 3. <u>Juvenile Idiopathic Arthritis or Rheumatoid Arthritis –</u>
 <u>Patient is Currently Receiving Kevzara.</u>
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Kevzara Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following (a, b, c, or d):
 - a) Patient has Rheumatoid Arthritis and has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq, or Xeljanz/XR [documentation required]; OR Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xelianz products (Xelianz and Xelianz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, an infliximab product (e.g., Remicade, biosimilars), Orencia (intravenous or subcutaneous), or Simponi (Aria or subcutaneous) also counts [documentation
 - **b)** Patient has Juvenile Idiopathic Arthritis and has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvog, Rinvog LQ, or Xeljanz [documentation required]; OR Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Rinvog products (Rinvog and Rinvog LQ) collectively counts as **ONE** product. A trial of a tocilizumab

required].

intravenous product (Actemra intravenous, biosimilar), Orencia intravenous or subcutaneous, an infliximab product (e.g., Remicade, biosimilars), or Simponi Aria also counts [documentation required].

- According to the prescriber, the patient has heart failure or a previously treated lymphoproliferative disorder; OR
- d) Patient has been established on Kevzara for at least 90 days and prescription claims history indicates at least a 90-day supply of Kevzara was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Kevzara for at least 90 days AND the patient has been receiving Kevzara via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Kevzara).

- **B)** If the patient has met criterion 3Ai (the standard *Inflammatory Conditions Kevzara Prior Authorization Policy* criteria), but criterion 3Aii is not met: offer to review for one of the following Products using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria.
 - i. Rheumatoid Arthritis: Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR.
 - ii. Juvenile Idiopathic Arthritis: Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Rinvoq LQ, or Xeljanz tablets.
- **3.** Other Conditions. Approve Kevzara (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Kevzara Prior Authorization Policy criteria.

Interleukin-17 Blockers

Bimzelx

1. Plaque Psoriasis - Initial Therapy.

A) Approve for 3 months if the patient meets BOTH of the following (i <u>and</u> ii):

- i. Patient meets the standard *Inflammatory Conditions Bimzelx Prior Authorization Policy* criteria for plaque psoriasis; AND
- ii. Patient has tried ONE of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya subcutaneous. Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry.
- B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Bimzelx Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu, Stelara subcutaneous, Taltz, or Tremfya subcutaneous) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. Plaque Psoriasis - Patient is Currently Receiving Bimzelx.

- **A)** Approve for 1 year if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Bimzelx Prior Authorization Policy* criteria; AND
 - **ii.** Patient meets ONE of the following (a <u>or</u> b):
 - a) Patient has tried ONE of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya subcutaneous; OR

 Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumabadbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry.
 - b) Patient has been established on Bimzelx for at least 90 days and prescription claims history indicates at least a 90-day supply of Bimzelx was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

<u>Note</u>: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been

receiving Bimzelx for at least 90 days AND the patient has been receiving Bimzelx via paid claims (e.g., patient has <u>not</u> been receiving samples or coupons or other types of waivers in order to obtain access to Bimzelx).

- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Bimzelx Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu, Stelara subcutaneous, Taltz, or Tremfya subcutaneous) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- **3.** Other Conditions. Approve Bimzelx (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Bimzelx Prior Authorization Policy criteria.

Cosentyx SC

1. Ankylosing Spondylitis – Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Cosentyx Subcutaneous Prior Authorization Policy* criteria; AND
 - ii. Patient has tried TWO of Enbrel, an adalimumab product, Rinvoq, Taltz, or Xeljanz/XR [documentation required].

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of Cimzia, an infliximab product (e.g. Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts [documentation required].

B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions – Cosentyx Subcutaneous Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Step 1 or Step 2 Product (Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Enbrel, Rinvoq, Taltz, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria.

2. Non-Radiographic Spondyloarthritis (nr-axSpA) - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Cosentyx Subcutaneous Prior Authorization Policy* criteria; AND
 - ii. Patient has tried TWO of Cimzia, Taltz, or Rinvoq [documentation required].

Note: A trial of Enbrel, an adalimumab product (e.g., Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry), an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts [documentation required]. A trial of multiple adalimumab products counts as ONE product.

B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions – Cosentyx Subcutaneous Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Step 1 or Step 2 Product (Cimzia, Taltz, or Rinvoq) using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria.

3. Plaque Psoriasis - Initial Therapy.

- **A)** Approve for 3 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Cosentyx Subcutaneous Prior Authorization Policy* criteria; AND
 - ii. Patient has tried TWO of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya [documentation required].

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product.

If the patient has met criterion 3Ai (the standard Inflammatory Conditions – Cosentyx Subcutaneous Prior Authorization Policy criteria), but criterion 3Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu, Stelara subcutaneous, Taltz, or

<u>Tremfya</u>) using the respective standard *Inflammatory Conditions – Prior Authorization Policy* criteria.

4. <u>Psoriatic Arthritis – Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Cosentyx Subcutaneous Prior Authorization Policy criteria; AND
 - ii. Patient meets ONE of the following (a or b):
 - a) Patient is ≥ 18 years of age AND has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvoq/Rinvoq LQ, Skyrizi subcutaneous, Stelara subcutaneous, Taltz, Tremfya, or Xeljanz/XR [documentation required]; OR
 - b) Patient is < 18 years of age AND has tried ONE of Enbrel, Rinvoq/Rinvoq LQ, or Stelara SC [documentation required].

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumabadbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (subcutaneous or Aria) also counts toward a trial of a TNFi [documentation required]. For a patient < 18 years of age, a trial of another TNFi counts towards a trial of Enbrel [documentation **required**]. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvog products (Rinvog and Rinvog LQ) collectively counts as **ONE** product.

- If the patient has met criterion 4Ai (the standard Inflammatory Conditions Cosentyx Subcutaneous Prior Authorization Policy criteria), but criterion 4Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Rinvoq, Rinvoq LQ, Skyrizi subcutaneous [pen or syringe], Stelara subcutaneous, Taltz, Tremfya, Xeljanz, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 5. Ankylosing Spondylitis; nr-axSpA; Plaque Psoriasis; or Psoriatic Arthritis Patient is Currently Receiving Cosentyx (Subcutaneous or Intravenous).

- **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Cosentyx Subcutaneous Prior Authorization Policy* criteria; AND
 - **ii.** Patient meets ONE of the following (a, b, c, d, e, f, or g):
 - a) Patient has Ankylosing Spondylitis and has tried TWO of Enbrel, an adalimumab product, Rinvoq, Taltz, or Xeljanz/XR [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as ONE product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as ONE product. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts [documentation required].
 - b) Patient has nr-axSpA and has tried TWO of Cimzia, Taltz, or Rinvoq [documentation required]; OR Note: A trial of Enbrel, an adalimumab product (e.g., Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry), an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts [documentation required]. A trial of multiple adalimumab products counts as ONE product.
 - c) Patient has <u>Plaque Psoriasis</u> and has tried TWO of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product.
 - d) Patient is ≥ 18 years of age with <u>Psoriatic Arthritis</u> and has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvoq/Rinvoq LQ, Skyrizi subcutaneous, Stelara subcutaneous, Taltz, Tremfya, or Xeljanz/XR [documentation required]; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts **Idocumentation required**].

- e) Patient is < 18 years of age with <u>Psoriatic Arthritis</u> and has tried ONE of Enbrel, Rinvoq/Rinvoq LQ, or Stelara SC [documentation required]; OR <u>Note</u>: A trial of another TNFi counts towards a trial of Enbrel [documentation required]. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product.
- f) According to the prescriber, the patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, or Psoriatic Arthritis has been established on Cosentyx intravenous for at least 90 days; OR
- g) Patient has been established on Cosentyx subcutaneous for at least 90 days and prescription claims history indicates at least a 90-day supply of Cosentyx SC was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Cosentyx SC for at least 90 days AND the patient has been receiving Cosentyx SC via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Cosentyx SC).

B) If the patient has met criterion 5Ai (the standard *Inflammatory Conditions – Cosentyx Subcutaneous Prior Authorization Policy* criteria), but criterion 5Aii is not met: offer to review for one of the following Products using the

respective standard *Inflammatory Conditions – Prior Authorization Policy* criteria:

- i. Ankylosing Spondylitis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Taltz, Xeljanz tablets, or Xeljanz XR.
- ii. nr-axSpA: Cimzia, Taltz, or Rinvoq.
- iii. Plaque Psoriasis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous (pen or syringe), Sotyktu, Stelara subcutaneous, Taltz, or Tremfya.
- iv. Psoriatic Arthritis in a Patient ≥ 18 years of age:

 Enbrel, Humira (NDCs starting with 00074), adalimumabadbm, Cyltezo, Hyrimoz (NDCs starting with 61314),
 adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla,
 Rinvoq, Rinvoq LQ, Skyrizi subcutaneous (pen or
 syringe), Stelara subcutaneous, Taltz, Tremfya, Xeljanz,
 or Xeljanz XR.
- v. Psoriatic Arthritis in a Patient < 18 years of age: Enbrel, Rinvog, Rinvog LQ, or Stelara SC.
- **6.** Other Conditions. Approve Cosentyx SC (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Cosentyx Subcutaneous Prior Authorization Policy criteria.

Siliq

1. Plaque Psoriasis - Initial Therapy.

- **A)** Approve for 3 months if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Siliq Prior Authorization Policy* criteria for plaque psoriasis; AND
 - ii. Patient has tried TWO of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya [documentation required].

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product.

B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions – Siliq Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting

with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu, Stelara subcutaneous, Taltz, or Tremfya) using the respective standard *Inflammatory Conditions – Prior Authorization Policy* criteria.

- 2. <u>Plaque Psoriasis Patient is Currently Receiving Siliq</u>.
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Silig Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following (a or b):
 - a) Patient has tried TWO of Enbrel, an adalimumab product, Otezla, Skyrizi subcutaneous, Sotyktu, Stelara subcutaneous, Taltz, or Tremfya [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product.
 - b) Patient has been established on Siliq for at least 90 days and prescription claims history indicates at least a 90-day supply of Siliq was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Siliq for at least 90 days AND the patient has been receiving Siliq via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Siliq).

B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions – Siliq Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu, Stelara subcutaneous, Taltz, or Tremfya) using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria.

	3. Other Conditions. Approve Siliq (initial therapy for a duration						
	as directed or $\frac{1}{year}$ for a patient continuing therapy) if the						
	patient meets the standard <i>Inflammatory Conditions – Siliq</i>						
	Prior Authorization Policy criteria.						
Taltz	All Conditions. Approve <u>Taltz</u> (initial therapy for a duration as						
	directed or <u>1 year</u> for a patient continuing therapy) if the patient						
	meets the standard <i>Inflammatory Conditions – Taltz Prior</i>						
	Authorization Policy criteria.						
	-23 Blockers						
Ilumya	1. Plaque Psoriasis - Initial Therapy.						
	A) Approve for 3 months if the patient meets BOTH of the						
	following (i <u>and</u> ii):						
	i. Patient meets the standard <i>Inflammatory Conditions</i> –						
	Ilumya Prior Authorization Policy criteria; AND						
	ii. Patient has tried TWO of Enbrel, an adalimumab product,						
	Otezla, Skyrizi subcutaneous, Sotyktu, Stelara						
	subcutaneous, Taltz, or Tremfya [documentation						
	required].						
	Note: Examples of adalimumab products include Humira,						
	Abrilada, adalimumab-adaz, adalimumab-adbm,						
	adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,						
	Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,						
	Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as ONE product.						
	B) If the patient has met criterion 1Ai (the standard						
	Inflammatory Conditions – Ilumya Prior Authorization Policy						
	criteria), but criterion 1Aii is not met: offer to review for a						
	Preferred Product (Enbrel, Humira [NDCs starting with						
	00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting						
	with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi,						
	Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu,						
	Stelara subcutaneous, Taltz, or Tremfya) using the						
	respective standard <i>Inflammatory Conditions – Prior</i>						
	Authorization Policy criteria.						
	2. Plaque Psoriasis - Patient is Currently Receiving Ilumya.						
	A) Approve for 1 year if the patient meets BOTH of the						
	following (i and ii):						
	i. Patient meets the standard Inflammatory Conditions –						
	Ilumya Prior Authorization Policy criteria; AND						
	ii. Patient meets ONE of the following (a or b):						
	a) Patient has plaque psoriasis and has tried TWO of						
	Enbrel, an adalimumab product, Otezla, Skyrizi						
	subcutaneous, Sotyktu, Stelara subcutaneous, Taltz,						
	or Tremfya [documentation required]; OR						
	Note: Examples of adalimumab products include						
	Humira, Abrilada, adalimumab-adaz, adalimumab-						
	adbm, adalimumab-fkjp, adalimumab-aaty,						
	adalimumab-ryvk, Simlandi, Amjevita, Cyltezo,						

- Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product.
- b) Patient has been established on Ilumya for at least 90 days and prescription claims history indicates at least a 90-day supply of Ilumya was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Ilumya for at least 90 days AND the patient has been receiving Ilumya via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Ilumya).

- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Ilumya Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Sotyktu, Stelara subcutaneous, Taltz, or Tremfya) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- **3.** Other Conditions. Approve Ilumya (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Ilumya Prior Authorization Policy criteria.

Omvoh SC

1. <u>Ulcerative Colitis – Initial Therapy.</u>

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Omvoh Subcutaneous Prior Authorization Policy criteria; AND
 - ii. Patient meets ONE of the following (a or b):
 - Patient has tried one of an adalimumab product, Skyrizi subcutaneous, Stelara subcutaneous, or Zymfentra; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-aacf, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio,

- Yuflyma, and Yusimry. A trial of an infliximab intravenous product (e.g., Remicade, biosimilars), Simponi subcutaneous, Skyrizi intravenous, or Stelara intravenous also counts.
- **b)** According to the prescriber, the patient has already started on or is currently undergoing induction therapy with Omvoh intravenous.
- If the patient has met criterion 1Ai (the standard Inflammatory Conditions Omvoh Subcutaneous Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. <u>Ulcerative Colitis - Patient is Currently Receiving Omvoh</u> Subcutaneous.

- **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - Patient meets the standard Inflammatory Conditions Omvoh Subcutaneous Prior Authorization Policy criteria;
 AND
 - **ii.** Patient meets ONE of the following (a <u>or</u> b):
 - **a)** Patient has tried one of an adalimumab product, Skyrizi subcutaneous, Stelara subcutaneous, or Zymfentra; OR
 - Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-aacf, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab intravenous product (e.g., Remicade, biosimilars), Simponi subcutaneous, Skyrizi intravenous, or Stelara intravenous also counts.
 - b) Patient has been established on Omvoh subcutaneous for at least 90 days and prescription claims history indicates at least a 90-day supply of Omvoh subcutaneous was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

<u>Note</u>: In cases where 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if

the prescriber has verified that the patient has been receiving Omvoh subcutaneous for at least 90 days AND the patient has been receiving Omvoh subcutaneous via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Omvoh subcutaneous). **B)** If the patient has met criterion 2Ai (the standard Inflammatory Conditions - Omvoh Subcutaneous Prior Authorization Policy criteria), but criterion 2Aii is not met, offer to review for a Preferred Product (Humira [NDCs starting with 00074], adalimumab-adaz, adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumabryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria. **3.** Other Conditions. Approve Omvoh subcutaneous (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions - Omvoh Subcutaneous Prior Authorization Policy criteria. **All Conditions.** Approve Skyrizi subcutaneous (initial therapy for Skyrizi Subcutane a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions - Skyrizi ous Subcutaneous Prior Authorization Policy criteria. Tremfya **All Conditions.** Approve Tremfya (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard *Inflammatory Conditions – Tremfya Prior* Authorization Policy criteria. IL-12/23 Blocker Stelara All Conditions. Approve Stelara subcutaneous (initial therapy for Subcutane a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions - Stelara ous Subcutaneous Prior Authorization Policy criteria. **Integrin Receptor Antagonist** 1. Crohn's Disease - Initial Therapy. **Entyvio SC** A) Approve for 6 months if the patient meets BOTH of the following (i and ii): i. Patient meets the standard *Inflammatory Conditions* – Entyvio Subcutaneous Prior Authorization Policy criteria; AND ii. Patient meets ONE of the following (a or b): a) Patient has tried TWO of an adalimumab product, Skyrizi subcutaneous, Stelara subcutaneous, Zymfentra, Cimzia, or Rinvoq [documentation required]; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-aacf, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of an infliximab intravenous product (e.g., Remicade, biosimilars), Skyrizi intravenous, or Stelara intravenous also counts [documentation required].

- **b)** According to the prescriber, the patient has already started on or is currently undergoing induction therapy with Entvyio IV.
- B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Entyvio Subcutaneous Prior Authorization Policy criteria), but criterion 1Aii is not met, offer to review for a Step 1 or Step 2 Product (Humira [NDCs starting with 00074], adalimumab-adaz, adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-ryvk, Simlandi, Skyrizi subcutaneous, Stelara subcutaneous, Rinvoq, Cimzia, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. <u>Ulcerative Colitis - Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Entyvio Subcutaneous Prior Authorization Policy criteria; AND
 - ii. Patient meets ONE of the following (a or b):
 - a) Patient has tried TWO of an adalimumab product, Skyrizi subcutaneous, Stelara subcutaneous, Zymfentra, Omvoh subcutaneous, Rinvoq, Simponi subcutaneous, or Xeljanz/XR [documentation required]; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-aacf, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of an infliximab intravenous product (e.g., Remicade, biosimilars), Omvoh intravenous, Skyrizi intravenous, or Stelara intravenous also counts [documentation required].

- **b)** According to the prescriber, the patient has already started on or is currently undergoing induction therapy with Entvyio IV.
- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Entyvio Subcutaneous Prior Authorization Policy criteria), but criterion 2Aii is not met, offer to review for a Step 1 or Step 2 Product (Humira [NDCs starting with 00074], adalimumab-adaz, adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-ryvk, Simlandi, Stelara subcutaneous, Omvoh subcutaneous, Rinvoq, Simponi SC, Skyrizi subcutaneous (on-body injector), Xeljanz/XR, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 3. <u>Crohn's Disease and Ulcerative Colitis Patient is Currently Receiving Entyvio Subcutaneous or Intravenous.</u>
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Entyvio Subcutaneous Prior Authorization Policy criteria; AND
 - ii. Patient meets ONE of the following conditions (a, b, c, or d):

a) Patient has Crohn's Disease and has tried TWO of an

- adalimumab product, Skyrizi subcutaneous, Stelara subcutaneous, Zymfentra, Cimzia, or Rinvoq [documentation required]; OR

 Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-aacf, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of an infliximab intravenous product (e.g., Remicade, biosimilars), Skyrizi intravenous, or Stelara intravenous also counts [documentation required].
- b) Patient has <u>Ulcerative Colitis</u> and has tried TWO of an adalimumab product, Skyrizi subcutaneous, Stelara subcutaneous, Zymfentra, Omvoh subcutaneous, Rinvoq, Simponi subcutaneous, or Xeljanz/XR [documentation required]; OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-aacf, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio,

Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of an infliximab intravenous product (e.g., Remicade, biosimilars), Omvoh intravenous, Skyrizi intravenous, or Stelara intravenous also counts **[documentation required]**.

- c) According to the prescriber, the patient has been established on Entyvio intravenous for at least 90 days; OR
- d) Patient has been established on Entyvio subcutaneous for at least 90 days and prescription claims history indicates at least a 90-day supply of Entyvio subcutaneous was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases where 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Entyvio subcutaneous for at least 90 days AND the patient has been receiving Entyvio subcutaneous via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Entyvio subcutaneous).

- **B)** If the patient has met criterion 3Ai (the standard *Inflammatory Conditions Entyvio Subcutaneous Prior Authorization Policy* criteria), but criterion 3Aii is not met, offer to review for one of the following Products using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria.
 - i. Crohn's Disease: Humira (NDCs starting with 00074), adalimumab-adaz, adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-ryvk, Simlandi, Skyrizi subcutaneous, Stelara subcutaneous, Rinvoq, Cimzia, or Zymfentra.
 - ii. Ulcerative Colitis: Humira (NDCs starting with 00074), adalimumab-adaz, adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, Omvoh subcutaneous, Rinvoq, Simponi SC, Xeljanz/XR, or Zymfentra.
- **4.** <u>Other Conditions</u>. Approve <u>Entyvio subcutaneous</u> (initial therapy for a duration as directed or <u>1 year</u> for a patient

continuing therapy) if the patient meets the standard Inflammatory Conditions – Entyvio Subcutaneous Prior Authorization Policy criteria.

Interleukin-1 Blocker

Kineret

1. Rheumatoid Arthritis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Kineret Prior Authorization Policy* criteria; AND
 - ii. Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvog, or Xeljanz/XR [documentation required]. Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkip, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, Orencia (subcutaneous or intravenous), an infliximab product (e.g., Remicade, biosimilar), Kevzara, or Simponi (Aria or subcutaneous) also counts [documentation required]
- Inflammatory Conditions Kineret Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. Rheumatoid Arthritis - Patient is Currently Receiving Kineret.

- **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Kineret Prior Authorization Policy* criteria; AND
 - **ii.** Patient meets ONE of the following (a <u>or</u> b):
 - a) Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq, or Xeljanz/XR [documentation required]; OR

Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, Orencia (subcutaneous or intravenous), an infliximab product (e.g., Remicade, biosimilar), Kevzara, or Simponi (Aria or subcutaneous) also counts [documentation required].

b) Patient has been established on Kineret at least 90 days <u>and</u> prescription claims history indicates <u>at least a 90-day supply of Kineret was dispensed within the past 130 days</u> [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Kineret for at least 90 days AND the patient has been receiving Kineret via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Kineret).

- If the patient has met criterion 2Ai (the standard Inflammatory Conditions Kineret Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- **3.** Other Conditions. Approve Kineret (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Kineret Prior Authorization Policy criteria.

<u>Note</u>: This includes Cryopyrin-Associated Periodic Syndromes (CAPS), Systemic Juvenile Idiopathic Arthritis.

T-Cell Costimulation Modulator

Orencia Subcutane ous

1. Rheumatoid Arthritis – Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy criteria; AND
 - **ii.** Patient meets ONE of the following (a <u>or</u> b):
 - a) Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvog, or Xeljanz/XR [documentation required]; OR Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, an infliximab product (e.g., Remicade, biosimilars), Kevzara, or Simponi (Aria or subcutaneous) also counts [documentation required].
 - **b)** According to the prescriber, the patient has heart failure, a previously treated lymphoproliferative disorder, a previous serious infection, OR a demyelinating disorder.
- B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 2. <u>Juvenile Idiopathic Arthritis/Juvenile Rheumatoid</u>
 <u>Arthritis Initial Therapy</u>.
 - **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):

- Patient meets the standard Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy criteria; AND
- ii. Patient meets ONE of the following (a or b):
 - a) Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq/Rinvoq LQ, or Xeljanz [documentation required]; OR

Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz tablets and Xeljanz oral solution) collectively counts as **ONE** product. A trial of either or both Rinvog products (Rinvog and Rinvog LQ) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Kevzara, Orencia intravenous, an infliximab product (e.g., Remicade, biosimilar), or Simponi Aria also counts [documentation] required].

- **b)** According to the prescriber, the patient has heart failure, a previously treated lymphoproliferative disorder, a previous serious infection, OR a demyelinating disorder.
- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Rinvoq LQ, Xeljanz tablets, or Xeljanz oral solution) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 3. <u>Psoriatic Arthritis Initial Therapy</u>.
 - **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - Patient meets the standard Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy criteria;
 AND
 - **ii.** Patient meets ONE of the following (a, b, or c):

- a) Patient is ≥ 18 years of age AND has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvog/Rinvog LQ, Skyrizi subcutaneous, Stelara subcutaneous, Taltz, Tremfya, or Xeljanz/XR **[documentation required]:** OR Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumabadbm, adalimumab-fkip, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts [documentation required]
- b) Patient is < 18 years of age AND has tried ONE of Enbrel, Rinvoq/Rinvoq LQ, or Stelara SC [documentation required]; OR Note: A trial of another TNFi counts towards a trial of Enbrel [documentation required]. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as ONE product.
- **c)** According to the prescriber, the patient has heart failure, a previously treated lymphoproliferative disorder, a previous serious infection, OR a demyelinating disorder.
- B) If the patient has met criterion 3Ai (the standard Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy criteria), but criterion 3Aii is not met: offer to review for a Step 1 or Step 2 Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Rinvoq, Rinvoq LQ, Skyrizi subcutaneous (pen or syringe), Stelara subcutaneous, Taltz, Tremfya, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 4. Rheumatoid Arthritis, Juvenile Idiopathic Arthritis, or Psoriatic Arthritis Patient is Currently Receiving Orencia (Subcutaneous or Intravenous).
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Orencia Subcutaneous Policy* criteria; AND

- ii. Patient meets ONE of the following (a, b, c, d, e, f, or g):
 - a) Patient has <u>Rheumatoid Arthritis</u> and has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq, or Xeljanz/XR [documentation required]; OR

Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, an infliximab product (e.g., Remicade, biosimilars), Kevzara, or Simponi (Aria or subcutaneous) also counts [documentation required].

- b) Patient has <u>Juvenile Idiopathic Arthritis</u> and has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq/Rinvoq LQ, or Xeljanz tablets or oral solution [documentation required]; OR
 - Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz tablets and Xeljanz oral solution) collectively counts as **ONE** product. A trial of either or both Rinvog products (Rinvog and Rinvog LQ) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Kevzara, Orencia intravenous, an infliximab product (e.g., Remicade, biosimilars), or Simponi Aria also counts [documentation required].
- c) Patient is ≥ 18 years of age with <u>Psoriatic Arthritis</u> AND has tried TWO of Enbrel, an adalimumab product, Otezla, Rinvoq/Rinvoq LQ, Skyrizi subcutaneous,

Stelara subcutaneous, Taltz, Tremfya, or Xeljanz/XR [documentation required]; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts [documentation required].

- d) Patient is < 18 years of age with <u>Psoriatic Arthritis</u> AND has tried ONE of Enbrel, Rinvoq/Rinvoq LQ, or Stelara SC [documentation required]; OR <u>Note</u>: A trial of another TNFi counts towards a trial of Enbrel [documentation required]. A trial of either or both Rinvoq products (Rinvoq and Rinvoq LQ) collectively counts as **ONE** product.
- e) According to the prescriber, the patient has been established on Orencia intravenous for at least 90 days: OR
- **f)** According to the prescriber, the patient has heart failure, a previously treated lymphoproliferative disorder, a previous serious infection, OR a demyelinating disorder; OR
- g) Patient has been established on Orencia subcutaneous for at least 90 days and prescription claims history indicates at least a 90-day supply of Orencia subcutaneous was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Orencia subcutaneous for at least 90 days AND the patient has been receiving Orencia subcutaneous via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Orencia subcutaneous).

- **B)** If the patient has met criterion 4Ai (the standard *Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy* criteria), but criterion 4Aii is not met, offer to review for one of the following Products using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria.
 - i. Rheumatoid Arthritis: Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR.
 - ii. Juvenile Idiopathic Arthritis: Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Rinvoq LQ, Xeljanz tablets, or Xeljanz oral solution.
 - iii. Psoriatic Arthritis in a Patient ≥ 18 Years of Age:

 Enbrel, Humira (NDCs starting with 00074), adalimumabadbm, Cyltezo, Hyrimoz (NDCs starting with 61314),
 adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla,
 Rinvoq, Rinvoq LQ, Skyrizi subcutaneous (pen or
 syringe), Stelara subcutaneous, Taltz, Tremfya, Xeljanz
 tablets, or Xeljanz XR.
 - iv. Psoriatic Arthritis in a Patient < 18 Years of Age: Enbrel, Rinvoq, Rinvoq LQ, or Stelara SC.
- **5.** Other Conditions. Approve Orencia subcutaneous (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Orencia Subcutaneous Prior Authorization Policy criteria.

Janus Kinases Inhibitors

Olumiant

- 1. Rheumatoid Arthritis Initial Therapy.
 - **A)** Approve for 6 months if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Olumiant Prior Authorization Policy* criteria; AND
 - ii. Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvoq, or Xeljanz/XR [documentation required].
 Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as ONE product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple

adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, an infliximab product (e.g., Remicade, biosimilars), Kevzara, Orencia (intravenous or subcutaneous), or Simponi (Aria or subcutaneous) also counts **[documentation required]**.

B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions – Olumiant Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. <u>Rheumatoid Arthritis – Patient is Currently Receiving</u> Olumiant.

- **A)** Approve for 1 year if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Olumiant Prior Authorization Policy* criteria; AND
 - **ii.** Patient meets ONE of the following (a <u>or</u> b):
 - a) Patient has tried TWO of a tocilizumab subcutaneous product, Enbrel, an adalimumab product, Rinvog, and Xeljanz/XR [documentation required]; OR Note: Examples of tocilizumab subcutaneous products include Actemra subcutaneous and Tyenne subcutaneous. A trial of multiple tocilizumab products counts as **ONE** product. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumabaaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of tocilizumab intravenous (Actemra intravenous, biosimilar), Cimzia, an infliximab product (e.g., Remicade, biosimilars), Kevzara, Orencia (intravenous or subcutaneous), or Simponi (Aria or subcutaneous) also counts [documentation required].
 - **b)** Patient has been established on Olumiant for at least 90 days <u>and</u> prescription claims history indicates <u>at least a 90-day supply of Olumiant was dispensed</u>

within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Olumiant for at least 90 days AND the patient has been receiving Olumiant via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Olumiant).

- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Olumiant Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Step 1 or Step 2 Product (Actemra subcutaneous, Tyenne subcutaneous, Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Rinvoq, Xeljanz tablets, or Xeljanz XR) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- **3.** Other Conditions. Approve Olumiant (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Olumiant Prior Authorization Policy criteria.

Rinvoq

1. Ankylosing Spondylitis – Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.

B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions – Rinvoq/LQ Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi,

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<u>or Taltz</u>) using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria.

2. <u>Crohn's Disease – Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvog/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one adalimumab product.

 Note: Examples of adalimumab products include Humira,
 Abrilada, adalimumab-adaz, adalimumab-adbm,
 adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,
 Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,
 Idacio, Yuflyma, and Yusimry. A trial of an infliximab
 product (e.g., Remicade, biosimilars; Zymfentra) or
 Cimzia also counts.
- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Preferred Product (Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous [on-body injector], Stelara subcutaneous, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

3. <u>Juvenile Idiopathic Arthritis – Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvog/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR
 - Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.
- B) If the patient has met criterion 3Ai (the standard Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy criteria), but criterion 3Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 4. Non-Radiographic Spondyloarthritis (nr-axSpA) Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient has tried Cimzia.

Note: A trial of Enbrel, an adalimumab product, an infliximab product (Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry.

B) If the patient has met criterion 4Ai (the standard Inflammatory Conditions – Rinvoq/LQ Prior Authorization Policy criteria), but criterion 4Aii is not met: offer to review for a Preferred Product (Cimzia or Taltz) using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria.

5. Rheumatoid Arthritis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvog/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.

B) If the patient has met criterion 5Ai (the standard Inflammatory Conditions – Rinvoq/LQ Prior Authorization Policy criteria), but criterion 5Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

6. <u>Psoriatic Arthritis – Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvog/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR

- Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
- Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy criteria), but criterion 6Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Stelara subcutaneous, Taltz, or Tremfya) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

7. <u>Ulcerative Colitis – Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one adalimumab product.

 Note: Examples of adalimumab products include Humira,
 Abrilada, adalimumab-adaz, adalimumab-adbm,
 adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,
 Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,
 Idacio, Yuflyma, and Yusimry. A trial of an infliximab
 product (e.g., Remicade, biosimilars; Zymfentra) or
 Simponi subcutaneous also counts.
- If the patient has met criterion 7Ai (the standard Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy criteria), but criterion 7Aii is not met: offer to review for a Preferred Product (Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 8. Ankylosing Spondylitis, Crohn's Disease, nr-axSpA, Rheumatoid Arthritis, Psoriatic Arthritis, or Ulcerative Colitis - Patient is Currently Receiving Rinvoq.
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following (a, b, c, d, e, f, g, or h):

- a) Patient has Ankylosing Spondylitis and has tried one of Enbrel or an adalimumab product; OR
 Note: Examples of adalimumab products include
 Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
- b) Patient has <u>Crohn's Disease</u> and has tried one adalimumab product; OR <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars; Zymfentra) or Cimzia also counts.
- c) Patient has <u>Juvenile Idiopathic Arthritis</u> and has tried ONE of Enbrel or an adalimumab product; OR <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.
- d) Patient has nr-axSpA and has tried Cimzia; OR
 Note: A trial of Enbrel, an adalimumab product, an infliximab product (Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts. Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry.
- e) Patient has Rheumatoid Arthritis and has tried one of Enbrel or an adalimumab product; OR

 Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.

- f) Patient has <u>Psoriatic Arthritis</u> and has tried one of Enbrel or an adalimumab product; OR

 <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
- g) Patient has <u>Ulcerative Colitis</u> and has tried one adalimumab product; OR

 <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars; Zymfentra) or Simponi subcutaneous also counts.
- h) Patient has been established on Rinvoq for at least 90 days <u>and</u> prescription claims history indicates <u>at least a 90-day supply of Rinvoq was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].</u>

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Rinvoq for at least 90 days AND the patient has been receiving Rinvoq via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Rinvoq).

- **B)** If the patient has met criterion 8Ai (the standard Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy criteria), but criterion 8Aii is not met: offer to review for one of the following Products using the respective standard Inflammatory Conditions Prior Authorization Policy criteria:
 - i. Ankylosing Spondylitis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, or Taltz.
 - ii. Crohn's Disease: Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with

- 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra.
- iii. Juvenile Idiopathic Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, or Simlandi.
- iv. nr-axSpA: Cimzia or Taltz.
- v. Rheumatoid Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, or Simlandi.
- vi. Psoriatic Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous (pen or syringe), Stelara subcutaneous, Taltz, or Tremfya.
- vii. Ulcerative Colitis: Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra.
- **9.** <u>All Other Conditions</u>. Approve <u>Rinvoq</u> (initial therapy for a duration as directed or <u>1 year</u> for a patient continuing therapy) if the patient meets the standard <u>Inflammatory Conditions</u> <u>Rinvoq/LQ Prior Authorization Policy</u> criteria.

Rinvoq LQ

- 1. <u>Juvenile Idiopathic Arthritis Initial Therapy</u>.
 - **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvog/LQ Prior Authorization Policy* criteria; AND
 - **ii.** Patient has tried one of Enbrel or an adalimumab product; OR
 - Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.
 - B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. <u>Psoriatic Arthritis – Initial Therapy</u>.

A) Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):

ii. Patient has tried one of Enbrel or an adalimumab

- i. Patient meets the standard *Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy* criteria; AND
- product; OR

 <u>Note</u>: Examples of adalimumab products include Humira,
 Abrilada, adalimumab-adaz, adalimumab-adbm,
 adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,
 Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,

adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.

B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions – Rinvoq/LQ Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Stelara subcutaneous, Taltz, or Tremfya) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

3. <u>Juvenile Idiopathic Arthritis or Psoriatic Arthritis – Patient is Currently Receiving Rinvoq/LQ</u>.

- **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following conditions (a, b, or c):
 - a) Patient has <u>Juvenile Idiopathic Arthritis</u> and has tried one of Enbrel or an adalimumab product; OR <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.
 - b) Patient has <u>Psoriatic Arthritis</u> and has tried one of Enbrel or an adalimumab product; OR

 <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g.,

- Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
- c) Patient has been established on Rinvoq/LQ for at least 90 days and prescription claims history indicates at least a 90-day supply of Rinvoq/LQ was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required].

 Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if

prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Rinvoq/LQ for at least 90 days AND the patient has been receiving Rinvoq/LQ via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Rinvoq/LQ).

- **B)** If the patient has met criterion 3Ai (the standard *Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy* criteria but criterion 3Aii is not met: offer to review for one of the following Products using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria:
 - i. Juvenile Idiopathic Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, or Simlandi.
 - ii. Psoriatic Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous (pen or syringe), Stelara subcutaneous, Taltz, or Tremfya.
- **4.** Other Conditions. Approve Rinvoq LQ (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Rinvoq/LQ Prior Authorization Policy criteria.

Xeljanz tablets, Xeljanz XR tablets

1. Ankylosing Spondylitis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR

<u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,

- Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
- B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Xeljanz/XR Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, or Taltz) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. Rheumatoid Arthritis - Initial Therapy.

A) Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):

ii. Patient has tried one of Enbrel or an adalimumab

Simponi (Aria or subcutaneous) also counts.

- i. Patient meets the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria; AND
- product; OR
 Note: Examples of adalimumab products include Humira,
 Abrilada, adalimumab-adaz, adalimumab-adbm,
 adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,
 Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,
 Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an
 infliximab product (e.g., Remicade, biosimilars), or
- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Xeljanz/XR Prior Authorization Policy criteria), but criterion 2Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

3. <u>Juvenile Idiopathic Arthritis – Initial Therapy</u>.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Xelianz/XR Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.

B) If the patient has met criterion 3Ai (the standard Inflammatory Conditions – Xeljanz/XR Prior Authorization Policy criteria), but criterion 3Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions – Prior Authorization Policy criteria.

4. Psoriatic Arthritis - Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR

 Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
- B) If the patient has met criterion 4Ai (the standard Inflammatory Conditions Xeljanz/XR Prior Authorization Policy criteria), but criterion 4Aii is not met: offer to review for a Step 1 Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous [pen or syringe], Stelara subcutaneous, Taltz, or Tremfya) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

5. Ulcerative Colitis – Initial Therapy.

- **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one adalimumab product.

 Note: Examples of adalimumab products include Humira,
 Abrilada, adalimumab-adaz, adalimumab-adbm,
 adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk,
 Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz,
 Idacio, Yuflyma, and Yusimry. A trial of an infliximab
 product (e.g., Remicade, biosimilars; Zymfentra) or
 Simponi subcutaneous also counts.
- **B)** If the patient has met criterion 5Ai (the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria), but criterion 5Aii is not met: offer to review

for a Preferred Product (<u>Humira [NDCs starting with 00074]</u>, adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra) using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria.

- 6. Ankylosing Spondylitis, Rheumatoid Arthritis, Juvenile Idiopathic Arthritis, Psoriatic Arthritis, or Ulcerative Colitis Patient is Currently Receiving Xeljanz/XR.
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following (a, b, c, d, e, or f):
 - a) Patient has Ankylosing Spondylitis and has tried one of Enbrel or an adalimumab product; OR

 Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
 - b) Patient has Rheumatoid Arthritis and has tried one of Enbrel or an adalimumab product; OR

 Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
 - c) Patient has <u>Juvenile Idiopathic Arthritis</u> and has tried one of Enbrel or an adalimumab product; OR <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.
 - **d)** Patient has <u>Psoriatic Arthritis</u> and has tried one of Enbrel or an adalimumab product; OR <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty,

- adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of Cimzia, an infliximab product (e.g., Remicade, biosimilars), or Simponi (Aria or subcutaneous) also counts.
- e) Patient has <u>Ulcerative Colitis</u> and has tried one adalimumab product; OR

 <u>Note</u>: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars; Zymfentra) or Simponi subcutaneous also counts.
- f) Patient has been established on Xeljanz/XR for at least 90 days and prescription claims history indicates at least a 90-day supply of Xeljanz/XR was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required]; OR Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Xelianz/XR for at least 90 days AND the patient has been receiving Xeljanz/XR via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Xeljanz/XR).
- **B)** If the patient has met criterion 6Ai (the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria but criterion 6Aii is not met: offer to review for one of the following Products using the respective standard *Inflammatory Conditions Prior Authorization Policy* criteria:
 - i. Ankylosing Spondylitis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, or Taltz.
 - ii. Rheumatoid Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, or Simlandi.
 - **iii. Juvenile Idiopathic Arthritis:** Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo,

- Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, or Simlandi.
- iv. Psoriatic Arthritis: Enbrel, Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Otezla, Skyrizi subcutaneous (pen or syringe), Stelara subcutaneous, Taltz, or Tremfya.
- v. Ulcerative Colitis: Humira (NDCs starting with 00074), adalimumab-adbm, Cyltezo, Hyrimoz (NDCs starting with 61314), adalimumab-adaz, adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, or Zymfentra.
- **7.** Other Conditions. Approve Xeljanz/XR (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Xeljanz/XR Prior Authorization Policy criteria.

Xeljanz oral solution

- 1. Juvenile Idiopathic Arthritis Initial Therapy.
 - **A)** Approve for 6 months if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria; AND
 - ii. Patient has tried one of Enbrel or an adalimumab product; OR

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.

- B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions Xeljanz/XR Prior Authorization Policy criteria), but criterion 1Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- 2. <u>Juvenile Idiopathic Arthritis Patient is Currently</u> Receiving Xeljanz.
 - **A)** Approve for 1 year if the patient meets BOTH of the following (i and ii):
 - i. Patient meets the standard *Inflammatory Conditions Xeljanz/XR Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following (a or b):
 - **a)** Patient has <u>Juvenile Idiopathic Arthritis</u> and has tried one of Enbrel or an adalimumab product; OR

- Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of an infliximab product (e.g., Remicade, biosimilars) or Simponi Aria also counts.
- b) Patient has been established on Xeljanz for at least 90 days and prescription claims history indicates at least a 90-day supply of Xeljanz was dispensed within the past 130 days [verification in prescription claims history required], or if claims history is not available, according to the prescriber [verification by prescriber required]; OR

Note: In cases when 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Xeljanz for at least 90 days AND the patient has been receiving Xeljanz via paid claims (e.g., patient has not been receiving samples or coupons or other types of waivers in order to obtain access to Xeljanz).

- B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions Xeljanz/XR Prior Authorization Policy criteria but criterion 2Aii is not met: offer to review for a Preferred Product (Enbrel, Humira [NDCs starting with 00074], adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-adaz, adalimumab-ryvk, or Simlandi) using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.
- **3.** Other Conditions. Approve Xeljanz oral solution (initial therapy for a duration as directed or 1 year for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions Xeljanz/XR Prior Authorization Policy criteria.

Phosphodiesterase Type 4 Inhibitor

Otezla

<u>All Conditions</u>. Approve <u>Otezla</u> (initial therapy for a duration as directed or <u>1 year</u> for a patient continuing therapy) if the patient meets the standard <u>Inflammatory Conditions</u> – <u>Otezla Prior Authorization Policy</u> criteria.

Sphingosine 1-Phosphate Receptor Modulator

Velsipity

- 1. <u>Ulcerative Colitis Initial Therapy</u>.
 - **A)** Approve for 6 months if the patient meets ALL of the following (i, ii, <u>and</u> iii):
 - i. Patient meets the standard *Inflammatory Conditions Velsipity Prior Authorization Policy* criteria; AND

ii. Patient has tried TWO of an adalimumab product, Skyrizi subcutaneous, Stelara subcutaneous, Omvoh subcutaneous, Rinvoq, Simponi subcutaneous, Xeljanz/XR, or Zymfentra [documentation required]; AND

Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-aacf, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of an infliximab product (e.g., Remicade, biosimilars), Entyvio intravenous or subcutaneous, Omvoh intravenous, Skyrizi intravenous, or Stelara intravenous also counts **[documentation required]**.

iii. Patient has tried Zeposia [documentation required].

B) If the patient has met criterion 1Ai (the standard Inflammatory Conditions – Velsipity Prior Authorization Policy criteria), but criterion 1Aii or criterion 1Aiii are not met, offer to review for a Step 1 or Step 2 Product (Humira [NDCs starting with 00074], adalimumab-adaz, adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, Omvoh subcutaneous, Rinvoq, Simponi SC, Xeljanz/XR, or Zymfentra), or Zeposia using the respective standard Inflammatory Conditions Prior Authorization Policy criteria.

2. <u>Ulcerative Colitis – Patient is Currently Receiving</u> Velsipity.

- **A)** Approve for 1 year if the patient meets BOTH of the following (i <u>and</u> ii):
 - i. Patient meets the standard *Inflammatory Conditions Velsipity Prior Authorization Policy* criteria; AND
 - ii. Patient meets ONE of the following conditions (a or b):
 - a) Patient meets BOTH of the following [(1) and (2)]:(1) Patient has tried TWO of an adalimumab
 - product, Skyrizi subcutaneous, Stelara subcutaneous, Omvoh subcutaneous, Rinvoq, Simponi subcutaneous, Xeljanz/XR, or Zymfentra [documentation required]; AND Note: Examples of adalimumab products include Humira, Abrilada, adalimumab-adaz, adalimumab-adbm, adalimumab-fkjp, adalimumab-aaty, adalimumab-ryvk, Simlandi, Amjevita, Cyltezo, Hadlima, Hulio, Hyrimoz, Idacio, Yuflyma, and Yusimry. A trial of multiple adalimumab products

counts as **ONE** product. A trial of either or both Xeljanz products (Xeljanz and Xeljanz XR) collectively counts as **ONE** product. A trial of an infliximab product (e.g., Remicade, biosimilars), Entyvio intravenous or subcutaneous, Omvoh intravenous, Skyrizi intravenous, or Stelara intravenous also counts [documentation required]. Patient has tried Zeposia [documentation (2) required]; OR **b)** Patient has been established on Velsipity for at least 90 days and prescription claims history indicates at least a 90-day supply of Velsipity was dispensed within the past 130 days [verification in **prescription claims history required**], or if claims history is not available, according to the prescriber [verification by prescriber required] Note: In cases where 130 days of the patient's prescription claim history file is unavailable to be verified, an exception to this requirement is allowed if the prescriber has verified that the patient has been receiving Velsipity for at least 90 days AND the patient has been receiving Velsipity via paid claims (e.g., patient has <u>not</u> been receiving samples or coupons or other types of waivers in order to obtain access to Velsipity). B) If the patient has met criterion 2Ai (the standard Inflammatory Conditions - Velsipity Prior Authorization Policy criteria), but criterion 2Aii is not met, offer to review for a Step 1 or Step 2 Product (Humira [NDCs starting with 00074], adalimumab-adaz, adalimumab-adbm, Cyltezo, Hyrimoz [NDCs starting with 61314], adalimumab-ryvk, Simlandi, Skyrizi subcutaneous (on-body injector), Stelara subcutaneous, Omvoh subcutaneous, Rinvog, Simponi SC, Xeljanz/XR, or Zymfentra) or Zeposia using the respective standard Inflammatory Conditions Prior Authorization Policy criteria. **3.** Other Conditions. Approve Velsipity (initial therapy for a duration as directed or <u>1 year</u> for a patient continuing therapy) if the patient meets the standard Inflammatory Conditions -Velsipity Prior Authorization Policy criteria. Zeposia **All Conditions.** Approve Zeposia if the patient meets the standard Multiple Sclerosis and Ulcerative Colitis - Zeposia Preferred Specialty Management Policy criteria. **Tyrosine Kinase 2 Inhibitor** Sotyktu **All Conditions.** Approve Sotyktu (initial therapy for a duration as directed or <u>1 year</u> for a patient continuing therapy) if the patient

meets the standard *Inflammatory Conditions – Sotyktu Prior Authorization Policy* criteria.

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HISTORY

70 Pages - Cigna National Formulary Coverage - Policy:Inflammatory Conditions Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies

Revision Feffective 01/01/2024 Adalimumab Products: Amjevita was removed and adalimumab-adbm was added to the Preferred Adalimumab Products. It was clarified that the Preferred Hyrimoz Product is specific for NDCs starting with 61314. A Note was added stating that Adalimumab-adaz and Adalimumab-adbm are Non-Preferred for some plans. Refer to respective Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies Policies for more information. Bimzelx: Bimzelx was added as a Non-Preferred Product for Plaque Psoriasis. A patient is directed to try two Preferred Products prior to Bimzelx (documentation required). Cosentyx Subcutaneous: Throughout the policy, it was clarified that this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	11/22/2023
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are Non-Preferred for some plans. Refer to respective Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies Policies for more information. Bimzelx: Bimzelx was added as a Non-Preferred Product for Plaque Psoriasis. A patient is directed to try two Preferred Products prior to Bimzelx (documentation required). Cosentyx Subcutaneous: Throughout the policy, it was clarified that this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
Policy for National Preferred, High Performance, and Basic Formularies Policies for more information. Bimzelx: Bimzelx was added as a Non-Preferred Product for Plaque Psoriasis. A patient is directed to try two Preferred Products prior to Bimzelx (documentation required). Cosentyx Subcutaneous: Throughout the policy, it was clarified that this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
Policies for more information. Bimzelx: Bimzelx was added as a Non-Preferred Product for Plaque Psoriasis. A patient is directed to try two Preferred Products prior to Bimzelx (documentation required). Cosentyx Subcutaneous: Throughout the policy, it was clarified that this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
Bimzelx: Bimzelx was added as a Non-Preferred Product for Plaque Psoriasis. A patient is directed to try two Preferred Products prior to Bimzelx (documentation required). Cosentyx Subcutaneous: Throughout the policy, it was clarified that this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
Psoriasis. A patient is directed to try two Preferred Products prior to Bimzelx (documentation required). Cosentyx Subcutaneous: Throughout the policy, it was clarified that this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
Bimzelx (documentation required). Cosentyx Subcutaneous: Throughout the policy, it was clarified that this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
this is the subcutaneous formulation. For a patient with Ankylosing Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
Spondylitis, Non-Radiographic Spondyloarthritis, and Psoriatic Arthritis currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
currently taking Cosentyx, an exception was added if, according to the prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
prescriber, the patient has been established on Cosentyx intravenous for at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
at least 90 days. For a patient < 18 years of age with Psoriatic Arthritis, a patient is directed to a trial of Enbrel or Stelara subcutaneous	
(documentation required). A Note was added that a previous trial of another tumor necrosis factor inhibitor counts towards a trial of Enbrel.	
Previously, a patient < 18 years of age was not required to try a	
Preferred Product prior to Cosentyx.	
Orencia Subcutaneous: A patient < 18 years of age is directed to a	
trial of Enbrel or Stelara SC (documentation required). A Note was added	
that a previous trial of another tumor necrosis factor inhibitor counts towards a trial of Enbrel. Previously, Orencia subcutaneous was not	
indicated in a patient < 18 years of age.	
Selected Effective 01/15/2024	01/03/2024
Revision Step 3c to Step Step 3c to Step	
2. The requirement for previous therapy was changed from three Preferred Products to one Preferred Product. Documentation supporting	
the previous trial is not required.	
Bimzelx: For Plaque Psoriasis, Bimzelx was moved from Step 3a to Step	
3c. A trial of two Step 1 Products is required (previously was two Step 1	
or Step 2 Products). Documentation of the previous trials is required.	
Cimzia: For Plaque Psoriasis, Cimzia was moved from Step 3a to Step 3c. A trial of two Step 1 Products is required (previously was two Step 1	
or Step 2 Products). Documentation of the previous trials is required.	
Cosentyx Subcutaneous: For Plaque Psoriasis, Cosnetyx subcutaneous	
was moved from Step 3a to Step 3c. A trial of two Step 1 Products is	
required (previously was two Step 1 or Step 2 Products). Documentation	
of the previous trials is required. Ilumya: For Plaque Psoriasis, Ilumya was moved from Step 3a to Step	
3c. A trial of two Step 1 Products is required (previously was two Step 1	
or Step 2 Products). Documentation of the previous trials is required.	
Siliq: For Plaque Psoriasis, Siliq was moved from Step 3a to Step 3c. A	
trial of two Step 1 Products is required (previously was two Step 1 or Step 2 Products). Documentation of the previous trials is required.	
Selected Effective 03/01/2024	01/24/2024
Revision Omvoh Subcutaneous: For Ulcerative Colitis, Omvoh subcutaneous	, ,===.
was added to Step 2. A trial of one Step 1 Product is required. An	
infliximab product, Simponi subcutaneous, and Stelara intravenous	
counts towards a trial of a Preferred Product. An exception was added for a patient who had already received induction with Omvoh intravenous	
who is not required to try a Preferred Product.	

Selected Revision	Entyvio Subcutaneous: For Ulcerative Colitis, Entyvio subcutaneous was added to Step 3a. A trial of two Step 1 or Step 2 Products is required (documentation required). An infliximab product, Omvoh intravenous, and Stelara intravenous counts towards a trial of a Preferred Product. An exception was added for a patient currently receiving Entyvio intravenous who is not required to try the Preferred Products. Velsipity: For Ulcerative Colitis, Velsipity was added to Step 4. A trial of two Step 1 or Step 2 Products plus Zeposia is required (documentation required). An infliximab product, Entyvio intravenous or subcutaneous, Omvoh intravenous, and Stelara intravenous counts towards a trial of a Preferred Product. Effective 04/01/2024 It was clarified that the Preferred Humira formulation includes NDCs starting with 00074.	02/28/2024
Selected	Effectivce 05/15/2024	05/08/2024
Revision	Actemra Subcutaneous: For Rheumatoid Arthritis and Polyarticular Juvenile Idiopathic Arthritis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Actemra subcutaneous. Rinvoq: For Rheumatoid Arthritis, Ankylosing Spondylitis, Psoriatic Arthritis, Crohn's Disease, and Ulcerative Colitis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Rinvoq. For Crohn's Disease and Ulcerative Colitis, Zymfentra was added as a Preferred Product. Xeljanz tablets/Xeljanz XR tablets: For Rheumatoid Arthritis, Juvenile Idiopathic Arthritis (Xeljanz tablets only), Ankylosing Spondylitis, Psoriatic Arthritis, and Ulcerative Colitis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Xeljanz tablets/Xeljanz XR tablets. For Ulcerative Colitis, Zymfentra was added as a Preferred Product. Xeljanz solution: For Juvenile Idiopathic Arthritis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Xeljanz. Cimzia: For Rheumatoid Arthritis, Ankylosing Spondylitis, Psoriatic Arthritis, Plaque Psoriasis, and Crohn's Disease, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Cimzia. For Crohn's Disease, Zymfentra was added as a Preferred Product. Kevzara: For Rheumatoid Arthritis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Kevzara. Kineret: For Rheumatoid Arthritis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Kineret. Olumiant: For Rheumatoid Arthritis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Orencia Subcutaneous: For Rheumatoid Arthritis, Ankylosing Spondylitis, Psoriatic Arthr	

list of Preferred adalimumab products that may have been tried prior to Cosentyx subcutaneous. **Sotyktu:** For Plague Psoriasis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Sotvktu. Bimzelx: For Plaque Psoriasis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Bimzelx. Ilumya: For Plaque Psoriasis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Ilumya. Siliq: For Plague Psoriasis, Simlandi and adalimumab-ryvk were added to the list of Preferred adalimumab products that may have been tried prior to Silia. Omvoh Subcutaneous: For Ulcerative Colitis, Simlandi, adalimumabryvk, and Zymfentra were added to the list of Preferred products that may have been tried prior to Omvoh Subcutaneous. Entyvio Subcutaneous: For Ulcerative Colitis, Simlandi, adalimumabryvk, and Zymfentra were added to the list of Preferred products that may have been tried prior to Entyvio subcutaneous. Zeposia: For Ulcerative Colitis, Simlandi, adalimumab-ryvk, and Zymfentra were added to the list of Preferred products that may have been tried prior to Zeposia. **Velsipity:** For Ulcerative Colitis, Simlandi, adalimumab-ryvk, and Zymfentra were added to the list of Preferred products that may have been tried prior to Velsipity. Selected Rinvoq: For Juvenile Idiopathic Arthritis, Rinvoq was added to Step 2. A 05/15/2024 Revision trial of a Step 1 Product is required. A trial of an infliximab product or Simponi Aria also counts. Rinvoq LQ: For Juvenile Idiopathic Arthritis, Rinvoq LQ was added to Step 2. A trial of a Step 1 Product is required. A trial of an infliximab product or Simponi Aria also counts. For Psoriatic Arthritis, Rinvoq LQ was added to Step 2. A trial of Enbrel or an adalimumab product is required. A trial of Cimzia, an infliximab product, or Simponi Aria or subcutaneous also counts. **Entyvio Subcutaneous:** For Crohn's Disease, Entyvio subcutaneous was added to Step 3a. A trial of two Step 1 or Step 2 Products is required (documentation required). An infliximab product, Skyrizi intravenous, and Stelara intravenous also counts. An exception was added for a patient currently receiving Entyvio intravenous who is not required to try the Preferred Products. **Sotyktu:** For Plaque Psoriasis, Sotyktu was moved from Step 2 to Step 1. As a result of this change, the requirement for one previous Preferred therapy was removed. Bimzelx: For Plaque Psoriasis, Bimzelx was moved from Step 3c to Step 3a. A trial of two Step 1 or Step 2 Products is required (previously was two Step 1 Products). Sotyktu was added as a Step 1 Product that may have been tried prior to Bimzelx. Documentation of previous trials remains required. Cimzia: For Plaque Psoriasis, Cimzia was moved from Step 3c to Step 3a. A trial of two Step 1 or Step 2 Products is required (previously was two Step 1 Products). Sotyktu was added as a Step 1 Product that may have been tried prior to Cimzia. For Psoriatic Arthritis, Rinvoq LQ was added as a Step 2 Product that may have been tried prior to Cimzia. Documentation of previous trials remains required. **Cosentyx Subcutaneous:** For Plaque Psoriasis, Cosentyx subcutaneous was moved from Step 3c to Step 3a. A trial of two Step 1 or Step 2 Products is required (previously was two Step 1 Products). Sotyktu was added as a Step 1 Product that may have been tried prior to Cosentyx

subcutaneous. For Psoriatic Arthritis, Rinvoq LQ was added as a Step 2 Product that may have been tried prior to Cosentyx subcutaneous. Documentation of previous trials remains required. **Ilumya:** For Plaque Psoriasis, Ilumya was moved from Step 3c to Step 3a. A trial of two Step 1 or Step 2 Products is required (previously was two Step 1 Products). Sotyktu was added as a Step 1 Product that may have been tried prior to Ilumya. Documentation of previous trials remains required. Siliq: For Plaque Psoriasis, Siliq was moved from Step 3c to Step 3a. A trial of two Step 1 or Step 2 Products is required (previously was two Step 1 Products). Sotyktu was added as a Step 1 Product that may have been tried prior to Siliq. Documentation of previous trials remains required. Orencia Subcutaneous: For Juvenile Idiopathic Arthritis, Rinvoq/LQ was added as a Step 2 Product that may have been tried prior to Orencia subcutaneous. For Psoriatic Arthritis, Rinvoq LQ was added as a Step 2 Product that may have been tried prior to Orencia subcutaneous. Documentation of previous trials remains required. Simponi Subcutaneous: For Psoriatic Arthritis, Rinvoq LQ was added as a Step 2 Product that may have been tried prior to Simponi Subcutaneous. Documentation of previous trials remains required. Selected **Tyenne Subcutaneous:** For Rheumatoid Arthritis and Juvenile 07/17/2024 Revision Idiopathic Arthritis, Tyenne subcutaneous was added as a Step 2 tocilizumab subcutaneous product. A patient is directed to one Step 1 Product prior to Tyenne. **Kevzara:** For Juvenile Idiopathic Arthritis, Kevzara was added to Step 3a. A patient is directed to two Step 1 or 2 products. Documentation of previous trials is required. For Rheumatoid Arthritis and Juvenile Idiopathic Arthritis, Tyenne subcutaneous was added as a Step 2 tocilizumab subcutaneous product that may have been tried prior to Kevzara. Orencia Subcutaneous: For Rheumatoid Arthritis and Juvenile Idiopathic Arthritis, Tyenne subcutaneous was added as a Step 2 tocilizumab subcutaneous product that may have been tried prior to Orencia Subcutaneous. For Juvenile Idiopathic Arthritis, a trial of Kevzara also counts. Skyrizi Subcutaneous: For Ulcerative Colitis, Skyrizi subcutaneous (onbody injector) was added to Step 1. Omvoh Subcutaneous: For Ulcerative Colitis, Skyrizi subcutaneous (on-body injector) was added to the list of Preferred products that may have been tried prior to Omvoh subcutaneous. A trial of Skyrizi intravenous also counts. **Rinvoq:** For Ulcerative Colitis, Skyrizi subcutaneous (on-body injector) was added to the list of Preferred products. **Simponi Subcutaneous:** For Ulcerative Colitis, Skyrizi subcutaneous (on-body injector) was added to the list of Preferred products. For Rheumatoid Arthritis, Tyenne subcutaneous was added as a Step 2 tocilizumab subcutaneous product that may have been tried prior to Simponi subcutaneous. Xeljanz/Xeljanz XR: For Ulcerative Colitis, Skyrizi subcutaneous (onbody injector) was added to the list of Preferred products. **Entyvio Subcutaneous:** For Ulcerative Colitis, Skyrizi subcutaneous (on-body injector) was added to the list of Preferred products that may have been tried prior to Entyvio subcutaneous. A trial of Skyrizi intravenous also counts. **Zeposia:** For Ulcerative Colitis, Skyrizi subcutaneous (on-body injector) was added to the list of Preferred products.

	Velsipity: For Ulcerative Colitis, Skyrizi subcutaneous (on-body injector) was added to the list of Preferred products that may have been tried prior to Velsipity. A trial of Skyrizi intravenous also counts. Cimzia: For Rheumatoid Arthritis, Tyenne subcutaneous was added as a Step 2 tocilizumab subcutaneous product that may have been tried prior to Cimzia. Kineret: For Rheumatoid Arthritis, Tyenne subcutaneous was added as a Step 2 tocilizumab subcutaneous product that may have been tried prior to Kineret. Olumiant: For Rheumatoid Arthritis, Tyenne subcutaneous was added as a Step 2 tocilizumab subcutaneous product that may have been tried prior to Olumiant.	
Selected Revision	A newly created Step 2b was added to the policy, which directs to a trial of one Step 1 Product. For existing Step 3a Products, it was clarified that these Products are directed a trial of two Step 1 or Step 2a Products (previously these were listed as Step 1 or Step 2 Products) with no changes to the criteria. For Plaque Psoriasis, Tremfya was clarified to be the subcutaneous formulation. Tremfya: For plaque psoriasis, it was clarified that the subcutaneous formulation is in Step 1. Bimzelx: Bimzelx was moved into Step 2a and requests are directed to a trial of one Step 1 Product (previously was in Step 2 and was directed to two Step 1 Products with documentation requirements). A previous trial of Tremfya was clarified to be the subcutaneous formulation.	10/02/2024

APPENDIX A

Table 1. Approved TNFis for Targeted Indications.*

	Rheumatology					Dermatology	Gastroen	terology	
	RA	JIA	AS	nr- axSpA	PsA	PsO	CD	UC	
Tumor Necrosis Factor Inhibitors									
Cimzia	\checkmark		\checkmark	\checkmark	\checkmark	\checkmark	√		
Enbrel	\checkmark	\checkmark	\checkmark		\checkmark	$\sqrt{}$			
Adalimumab Products (Humira, biosimilars)	√	√	√		√	V	√	√	
Infliximab Intravenous Products	√		√		√	√	√	√	
Zymfentra	1				1		√^	√^	
Simponi Subcutaneous	√		√		√			√	
Simponi Aria	√	√	√		√				

TNFis – Tumor necrosis factor inhibitors; * Refer to the selected standard *Inflammatory Conditions Prior Authorization Policies* for the specific patient population approved for each indication; RA – Rheumatoid arthritis; JIA – Juvenile idiopathic arthritis; AS – Ankylosing spondylitis; nr-axSpA – Non-radiographic spondyloarthritis; PsA – Psoriatic arthritis; PsO – Plaque psoriasis; CD – Crohn's disease; UC – Ulcerative colitis; ^ Maintenance dosing only.

Table 2. Approved IL-17, IL-23, and IL-12/23 Blockers for Targeted Indications.*

Table 2. Approved											
	R	heumatolog	У	Dermatolo	Gastroenterology						
	Ankylosin g Spondyliti	nr-axSpA	Psoriatic Arthritis	gy Plaque Psoriasis	Crohn's Disease	Ulcerative Colitis					
	S										
Interleukin-17 Blockers											
Bimzelx	√	√	\checkmark	\checkmark							
Cosentyx Subcutaneous	√	√	√	√							
Cosentyx	√	√	√								
Intravenous	V	V	V								
Siliq				√							
Taltz	√	\checkmark	√	√							
Interleukin-23 Bl	ockers										
Ilumya				\checkmark	\checkmark						
Omvoh Intravenous						√#					
Omvoh Subcutaneous						√^					
Skyrizi Intravenous					√ #	√#					
Skyrizi Subcutaneous			√	√	√^	√^					
Tremfya Intravenous						√#					
Tremfya Subcutaneous			√	√		√^					
Interleukin-12/2	3 Blockers			<u> </u>		1					
Stelara Subcutaneous			√	√	√^	√^					
Stelara Intravenous					√#	√#					

IL – Interleukin; * Refer to the selected standard *Prior Authorization Policies* for the specific patient population approved for each indication; nr-axSpA – Non-radiographic spondyloarthritis; ^ Maintenance dosing only; # Induction dosing only.

Table 3. Approved Oral tsDMARDs for Targeted Indications.*

		Rh	eumatolog	Dermatol ogy	Gastroer	nterology		
	RA	JIA	AS	nr- axSpA	PsA	PsO	CD	UC
Janus K	inases Inh	ibitors						
Olumia nt	√							
Rinvoq	\checkmark	\checkmark	√	\checkmark	\checkmark		√	\checkmark
Rinvoq LQ		√		√			-	-
Xeljanz tablets	√	√#	√		√			√
Xeljanz oral solutio n		√ #		1				-
Xeljanz XR	√		√		√			√

⁷⁰ Pages - Cigna National Formulary Coverage - Policy:Inflammatory Conditions Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies

Phospho	Phosphodiesterase Type 4 Inhibitor									
Otezla		-	-		\checkmark	\checkmark		-		
Sphingo	Sphingosine 1-Phosphate Receptor Modulator									
Velsipit								\checkmark		
У										
Zeposi								\checkmark		
a										
Tyrosine	Tyrosine Kinase 2 Inhibitor									
Sotykt										
u										

tsDMARDs – Targeted synthetic disease-modifying antirheumatic drugs; *Refer to the selected standard *Prior Authorization Policies* for the specific patient population approved for each indication; RA – Rheumatoid arthritis; JIA – Juvenile idiopathic arthritis; AS – Ankylosing spondylitis; nr-axSpA – Nonradiographic axial spondyloarthritis; PsA – Psoriatic arthritis; PsO – Plaque psoriasis; CD – Crohn's disease; UC – Ulcerative colitis; *Indicated in polyarticular JIA.

Table 4. Other Approved Biologics for Targeted Indications.*

Table 4. Other Approved Bio		Rheumatology					
	Rheumatoid Arthritis	Juvenile Idiopathic Arthritis	Psoriatic Arthritis	Crohn's Disease	Ulcerativ e Colitis		
Integrin Receptor Antagoni	st						
Entyvio Intravenous				\checkmark	\checkmark		
Entyvio Subcutaneous				ô	ô		
Interleukin-6 Blockers							
Tocilizumab Intravenous Products (Actemra, biosimilar)	√	√^					
Tocilizumab Subcutaneous Products (Actemra, biosimilar)	√	√^					
Kevzara	√	√					
Interleukin-1 Blocker							
Kineret	\checkmark			-			
T-Cell Costimulation Modula	itor						
Orencia Intravenous	√	$\sqrt{*}$	\checkmark				
Orencia Subcutaneous	\checkmark	√#					
CD20-Directed Cytolytic An	tibody						
Rituximab Intravenous Products	√						

^{*}Refer to the selected standard *Prior Authorization Policies* for the specific patient population approved for each indication; ^ Indicated in polyarticular and systemic JIA; * Indicated in polyarticular JIA; * Maintenance dosing only.

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