

# **MEDICAL POLICY UPDATE**

### IN THIS ISSUE

Injectable Drugs Added to Site of Care	8
ADAMTS13, recombinant-krhn (Adzynma) Added to Site of Care	8
Secukinumab (Cosentyx) Added to Site of Care	9
Coverage Criteria Established for Nogapendekin alfa inbakicept-pmln (Anktiva)	9
Criteria Established for Imetelstat (Ryelto)	10
Updated criteria for Nerve Conduction Studies and Electromyography	10

# **Policy**

Policy Titles	Anticipated Issue Date	30 Day Notification Information
E-31 - Negative Pressure Wound Therapy Pumps/Vacuum Assisted Closure of Chronic Wounds	09/02/2024	This is an annual review. The policy position was reorganized. Coding was updated. Age restriction was removed.
E-47 - Non-Powered Negative Pressure Wound Therapy System	09/02/2024	This is an annual review. The policy position was reorganized. Coding was updated. Age restriction was removed.
E-68 - High Frequency Chest Wall Oscillation Devices	09/02/2024	This policy is scheduled for annual review. Administrative changes have been made with no criteria updates. The policy will publish on September 02, 2024.
	09/23/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Additional minor administrative changes were made to the policy. Policy will publish on September
I-4 - Hemophilia Treatment	09/23/2024	23, 2024.  This policy is scheduled for annual review. There is no indication for change in coverage. Policy will
I-76 - Ziconotide (Prialt®)	09/16/2024	publish September 16, 2024.

		This policy is scheduled for annual
		review. There is no indication for
		change in coverage. Policy will
I-78 - Intravitreal Implants	09/16/2024	publish September 16, 2024.
		This policy is scheduled for annual
		review. There is no indication for
		change in coverage. Policy will
I-86 - Bevacizumab (Avastin®)	09/09/2024	publish on September 9, 2024
		This policy is scheduled for annual
		review. There is no indication for
		change in coverage. Policy will
I-94 - Intravitreal Injections	09/09/2024	publish September 9, 2024.
		This policy is being revised to
		include the new FDA approved
		expanded indication for Keytruda
		and Imfinzi. Keytruda and Imfinzi
		are now indicated as a single agent
L 120 Programmed Death Becenter (PD 1)		in combination with carboplatin and
I-120 - Programmed Death Receptor (PD-1)/ Programmed Death-Ligand (PD-L1) Blocking		paclitaxel for the treatment of individuals with primary advanced
Antibodies	07/29/2024	or recurrent endometrial carcinoma.
Aitibodies	0112312024	This policy is up for annual review.
		There are no indications for a
		change in coverage at this time.
		Policy will publish on September
I-149 - Chelation Therapy for Off-Label Uses	09/23/2024	23, 2024.
		This policy was updated to add
		Cosentyx and Adzynma to the Site
		of Care program. Policy will publish
		on November 1, 2024 with the
I-151 - Site of Care	11/01/2024	standard 90 day notification.
		This policy is scheduled for annual
		review. There is no indication for
		change in coverage. Policy will
I-181 - Pralatrexate (Folotyn)	09/23/2024	publish September 23, 2024.
		This policy is being revised to
		capture the new FDA expanded
		indication for Skyrizi for the
		treatment of adult individuals with
		moderate to severe ulcerative
		colitis (UC). Additional
1 100 Interleukin 22 Antercanista /Illumica CO		administrative changes are also
I-199 - Interleukin-23 Antagonists (Ilumya SC	07/29/2024	being made to the policy. Policy will
and Skyrizi IV)	0112912024	publish on July 29, 2024.  This policy is scheduled for annual
		review. There is no indication for
		change in coverage. Policy update
		includes minor language revisions.
		Policy will publish September 23,
I-207 - Tagraxofusp-erzs (Elzonris)	09/23/2024	2024.
1 _ 1		This policy is up for annual review.
		There are no indications for a
		change in coverage at this time.
I-209 - Emapalumab-Izsg (Gamifant)	09/23/2024	Minor administrative changes were

		made to the policy. Policy will
		publish on September 23, 2024.
		This policy is up for annual review. There are no indications for a change in coverage at this time. Minor administrative changes were made to the policy. Coding was updated to NCCN
I-214 - Luspatercept (Reblozyl)	09/23/2024	recommendations. This policy will publish on September 23, 2024.
I-218 - Crizanlizumab (Adakveo)	09/16/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 16, 2024.
I-253 - Betibeglogene autotemcel (Zynteglo)	09/09/2024	This policy is up for annual review. The policy was revised to remove the upper age limit and allow for additional TDT genotypes on a case by case basis. Denial statement was also updated to not medically necessary. The policy will publish on September 9, 2024.
I-254 - Spesolimab (Spevigo)	09/16/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 16, 2024.
I-259 - Entranacogene dezaparvovec (EntranaDez)	09/09/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 9, 2024.
I-270 - Epcoritamab-bysp (Epkinly)	09/09/2024	This policy is scheduled for annual review. Policy revision includes coding update. Policy will publish September 9, 2024.
I-271 - Valoctocogene roxaparvovec (Roctavian)	09/09/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 9, 2024.
I-273- ADAMTS13, recombinant-krhn (Adzynma)	11/01/2024	Policy is being updated with addition of Adzyma to Site of Care Program. Policy will publish November 1, 2024
I-280 - Secukinumab (Cosentyx)	11/01/2024	This policy is up for annual review. The policy was updated to include which specialist physician (dermatologist, rheumatologist) should be prescribing the medication based on indication, in order to align with pharmacy policy criteria. Place of service was also

		updated to outpatient-infusion.
		Policy will publish on November 1, 2024.
		This policy is up for annual review.
		There are no indications for a
		change in coverage at this time. Policy will publish on September 2,
I-281 - Exagamglogene autotemcel	09/02/2024	2024.
		This policy is up for annual review.
		There are no indications for a
I-282 - Lovotibeglogene autotemcel		change in coverage at this time. Policy will publish on September 2,
(Lyfgenia)	09/02/2024	2024.
,		This is a new policy for the recently
		FDA approved imetelstat (Ryelto)
		for the treatment of adults with low- to intermediate-1 risk
		myelodysplastic syndromes (MDS)
		with transfusion-dependent anemia
		requiring four or more red blood cell
		units over 8 weeks who have not
		responded to or have lost response to or are ineligible for
		erythropoiesis-stimulating agents.
I-288 - Imetelstat (Rytelo)	07/29/2024	Policy will publish on July 29, 2024.
		This is a new policy establishing
		criteria for new to market therapy (Tarlatamab-dlle) Imdelltra, a Tcell
		engager therapy indicated for adult
		individuals with extensive small cell
		lung cancer. Policy will publish July
I-290 - (Tarlatamab-dlle) Imdelltra	07/29/2024	29, 2024. This policy is up for annual review.
		There are no indications for a
		change in coverage at this time.
I-291 - Lovotibeglogene autotemcel		Policy will publish on September 2,
(Lyfgenia)	09/02/2024	2024.
		This policy was revised to include dx code C43.111 in the diagnosis
		code table for procedure code
MA I-120 - Programmed Death Receptor		J9299. This code was inadvertently
(PD-1)/ Programmed Death-Ligand (PD-L1)	07/00/0004	omitted from the table. Policy will
Blocking Antibodies	07/29/2024	publish on July 29, 2024.
		This policy is being revised to capture the diagnosis codes for the
		new FDA expanded indication for
		Skyrizi for the treatment of adult
MAL 400 Interleukin 32 Antonosisto ///		individuals with moderate to severe
MA I-199 - Interleukin-23 Antagonists (Ilumya SC and Skyrizi IV)	07/29/2024	ulcerative colitis (UC). Policy will publish on July 29, 2024.
OG GING ORYTIZETV)	0112012024	This policy is scheduled for annual
		review. There is no indication for
	00/00/00	change in coverage. Policy update
MA I-207 - Tagraxofusp-erzs (Elzonris)	09/23/2024	includes minor language revisions.

		Policy will publish September 23, 2024.
MA I-209 - Emapalumab-Izsg (Gamifant)	09/23/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Minor administrative changes were made to the policy. Policy will publish on September 23, 2024.
MA L 221 - Crizonlizumob (Adakuoo)	09/16/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September
MA I-221 - Crizanlizumab (Adakveo)  MA I-223 - Luspatercept (Reblozyl)	09/10/2024	This policy is up for annual review. Coding was updated to NCCN recommendations. This policy will publish on September 23, 2024.
MA I-264- Spesolimab (Spevigo)	09/16/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 16, 2024.
MA I-269 - Entranacogene dezaparvovec (EntranaDez)	09/10/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 9, 2024.
MA I-279 - Epcoritamab-bysp (Epkinly)	09/09/2024	This policy is scheduled for annual review. Policy update includes language and coding revisions. Policy will publish September 9, 2024.
MA I-280 - Valoctocogene roxaparvovec (Roctavian)	09/09/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 9, 2024.
MA I-289 - Secukinumab (Cosentyx)	11/01/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on November 1, 2024.
MA I-290 - Exagamglogene autotemcel	09/02/2024	This policy is up for annual review. There are no indications for a change in coverage at this time. Policy will publish on September 2, 2024.
MA I-298 - Imetelstat (Rytelo)	07/29/2024	This is a new policy for the recently FDA approved imetelstat (Ryelto) for the treatment of adults with low-to intermediate-1 risk myelodysplastic syndromes (MDS) with transfusion-dependent anemia requiring four or more red blood cell

		units over 8 weeks who have not
		responded to or have lost response
		to or are ineligible for
		erythropoiesis-stimulating agents.
		Policy will publish on July 29, 2024
		This is a new policy for
		Betibeglogene autotemcel
		(Zynteglo). Standardized Medicare
		coverage criteria was established
MA I-299 - Betibeglogene autotemcel		for this policy. Policy will publish on
(Zynteglo)	09/09/2024	September 9, 2024.
		This is a new policy establishing
		criteria for new to market therapy
		(Tarlatamab-dlle) Imdelltra, a Tcell
		engager therapy indicated for adult
		individuals with extensive small cell
		lung cancer. Policy will publish July
MA I-300 - (Tarlatamab-dlle) Imdelltra	07/29/2024	29, 2024.
100 (1000)		This policy is scheduled for annual
		review. There is no indication for
		change in coverage. Policy will
MA I-75 - Bevacizumab (Avastin)	09/09/2024	publish on September 9, 2024
With 10 Bevacizarias (1tvastiri)	00/00/2024	This policy is scheduled for annual
		review. Administrative changes
		were made. Further changes
		include an update to criteria for
		non-ambulatory orthoses and an
		addition to list microprocessor
		controlled KAF/KAFO as
		experimental and investigational.
		This policy will publish on
O-24 - Ankle-Foot/Knee-Ankle-Foot Orthosis	09/02/2024	September 02, 2024.
		The policy is being revised to
		include surgical codes that require
		additional documentation to
		determine if the service is eligible
		for team surgery reimbursement.
		This follows the Medicare Physician
S-12 - Team Surgery	07/22/2024	Fee Schedule.
,		This is an annual review. The policy
		position was reorganized, and
		language was updated. Coding was
S-131 - Sacral Nerve Modulation/Stimulation	09/02/2024	updated.
		This is an annual review. There are
		no recommended revisions. This
S-186 - Magnetic Resonance Imaging (MRI)-		policy will publish on September 2,
Guided Focused	09/02/2024	2024.
34,404 1 004004	00/02/2027	This policy is scheduled for annual
		review. Criteria requiring the
		performing provider and facility to
		meet the recommendations for
C 202 Transactheter Dulmanam / Value		performing TPV implantation
S-203 - Transcatheter Pulmonary Valve	00/00/0004	outlined in SCAI/AATS/ACC/STS
Implantation	09/02/2024	Operator and Institutional

		Requirements for Transcatheter Valve Repair and Replacement has been added. Administrative changes have been made.
S-204 - Endoscopic Radiofrequency Ablation/Cryotherapy	09/02/2024	This is an annual review with no recommended changes. The policy will publish on September 2, 2024.
S-226 - Placental/Umbilical Cord Blood as a Source of Stem Cells	09/09/2024	This policy was revised to clarify that an individual must not have a suitable matched related donor, matched unrelated donor (MUD), mismatched unrelated donor (MMUD), or haploidentical donor readily available. Additional minor administrative changes were made to the policy. Policy will publish on September 9, 2024.
S-262 - Eustachian Tube Balloon Dilatation	09/02/2024	This is an annual review. There are no recommended changes. This policy will publish on September 2, 2024.
S-268 - Endobronchial Valve Surgery	09/02/2024	This policy is scheduled for annual review. The place of service has been changed from Inpatient/Outpatient to Inpatient only. The policy will publish on September 2, 2024.
S-270 - Endoscopic Stricturotomy	09/02/2024	This policy is scheduled for annual review. There are no changes to criteria. The policy will publish on September 2, 2024.
U-7 - Fetal Surgery for Prenatally Diagnosed Malformations	09/02/2024	This policy is scheduled for annual review. This policy is being archived. The policy will publish on September 2, 2024.
U-8 - Treatment of Twin-Twin Transfusion Syndrome with Amnioreduction and/or Fetoscopic Laser Therapy	09/02/2024	This policy is scheduled for annual review. This policy is being archived. The policy will publish on September 2, 2024.
V-44 - Medical Nutrition Management Services (MNT)	09/09/2024	This is an annual review. No recommended changes to coverage criteria. Operational guidelines will be revised to add prepay logic to both covered and non-covered dx codes. This policy will publish on September 9, 2024.
Y-23 - Chronic Pain Management	09/02/2024	This policy is scheduled for annual review. There are no changes to policy criteria. The policy will publish on September 2, 2024.

#### Injectable Drugs Added to Site of Care



Highmark Blue Cross Blue Shield has established new criteria for I-151, Site of Care. Cosentyx and Adzynma are being added to the Site of Care program.

This revised Medical Policy will apply to professional providers and facility claims. The effective date is November 1, 2024.

Place of Service: Outpatient-Infusion

Please refer to Medical Policy I-151, Site of Care, for additional information.

#### ADAMTS13, recombinant-krhn (Adzynma) Added to Site of Care



Highmark Blue Cross Blue Shield has revised criteria for I-273 ADAMTS13, recombinant-krhn (Adzynma). The policy was updated with the addition of Adzynma to the Site of Care Program.

This revised Medical Policy will apply to professional providers and facility claims. The effective date is November 1, 2024.

Place of Service: Outpatient- Infusion

Please refer to Medical Policy I-273 ADAMTS13, recombinant-krhn (Adzynma), and Medical Policy I-151, Site of Care, for additional information.

#### Secukinumab (Cosentyx) Added to Site of Care



Highmark Blue Cross Blue Shield has revised criteria for I-280 Secukinumab (Cosentyx). The policy was updated to include which specialist physician (dermatologist, rheumatologist) should be prescribing the medication based on indication. The following indications now require a specialist physician to prescribe:

- Ankylosing Spondylitis
- Psoriatic Arthritis
- Non-radiographic Axial Spondyloarthritis

Secukinumab (Cosentyx) is also being added to the Site of Care program.

This revised Medical Policy will apply to professional providers and facility claims. The effective date is November 1, 2024.

Place of Service: Outpatient-Infusion

Please refer to Medical Policy I-280, Secukinumab (Cosentyx), for additional information.

# Coverage Criteria Established for Nogapendekin alfa inbakicept-pmln (Anktiva)



Highmark Blue Cross Blue Shield has established new criteria for I-287, Nogapendekin alfa inbakicept-pmln (Anktiva). This is a new policy creating criteria for Anktiva, an intravesical therapy indicated with Bacillus Calmette-Guérin (BCG) for the treatment of adult patients with BCGunresponsive nonmuscle invasive bladder cancer (NMIBC) with carcinoma in situ (CIS) with or without papillary tumors.

This revised Medical Policy will apply to professional providers and facility claims. The effective date is July 8, 2024.

Place of Service: Outpatient

Please refer to Medical Policy for I-287, Nogapendekin alfa inbakicept-pmln (Anktiva), for additional information.

#### **Criteria Established for Imetelstat (Ryelto)**



Highmark Blue Cross Blue Shield has established new criteria for Imetelstat (Ryelto) for the treatment of myelodysplastic syndrome (MDS) with transfusion dependent anemia. The following criteria will be established:

- Individual is 18 years of age or older; and
- Individual has a documented diagnosis of low- to intermediate- risk MDS as defined by **ONE** of the following:
  - Revised International Prognostic Scoring System (IPSS-R); Very low, low, intermediate (defined as a score of 0 to 4.5); or
  - IPSS: Low/Intermediate-1 (Score 0 to 1); or
  - WHO-Based Prognostic Scoring System (WPSS): very low, low, intermediate (Score 0 to 2); and
- Prescribed by or in consultation with a hematologist, oncologist, or other specialist with expertise in the diagnosis and management of myelodysplastic syndromes; and
- Prescriber has ruled out and/or addressed other causes of anemia [e.g., abnormal bleeding (gastrointestinal, uterine, etc.), hemolysis, nutritional deficiency, renal disease]; **and**
- Individual has required four (4) or more red blood cell units over an eight (8) week period; and
- Individual has had no response to or is ineligible for an erythropoiesis-stimulating agents (ESA)

This new Medical Policy will apply to professional providers and facility claims. The effective date is July 29, 2024.

Place of Service: Outpatient

Please refer to Medical Policy I-288, Imetelstat (Rytelo), for additional information.

#### **Updated criteria for Nerve Conduction Studies and Electromyography**



Highmark Blue Cross Blue Shield has established new criteria for M-28, Nerve Conduction Studies and Electromyography. The following criteria has been added to the policy:

Testing for the purpose of monitoring disease intensity or treatment efficacy in polyneuropathy of diabetes or end stage renal disease (ESRD) is not covered.

This revised Medical Policy will apply to professional providers and facility claims. The effective date is October 28, 2024.

Place of Service: Inpatient/Outpatient

Please refer to Medical Policy M-28, Nerve Conduction Studies and Electomyography, for additional information.



### **Comments on These Medical Policies?**

We want to know what you think about our new medical policy changes. Send us an email with any questions or comments that you may have on the new medical policies in this edition of Medical Policy Update.

Write to us at medicalpolicy@highmark.com



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Medical Policy Update is a monthly newsletter for the health care providers who participate in our networks and submit claims to Highmark using the appropriate HIPAA transactions or claim forms as required by Highmark. This publication focuses only on medical policy and claims administration updates, including coding guidelines and procedure code revisions, and is the sole source for this information. For all other news, information, and updates, be sure to read *Provider News*, available on the Provider Resource Center.

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