

# Entyvio<sup>®</sup> (vedolizumab) (Intravenous)

Document Number: IC-0202

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# I. Length of Authorization

Coverage will be provided for 14 weeks initially and may be renewed every 6 months thereafter unless otherwise specified.

- Dose escalation requests for Crohn's Disease and Ulcerative Colitis: Coverage will be provided for 3 months and may be renewed every 6 months thereafter (*see Section V for therapy continuation details*).
- Therapy for the Management of Immune Checkpoint Inhibitor-Related Diarrhea/Colitis: Coverage will be provided for 3 doses total and may not be renewed.
- Therapy for Ulcerative Colitis in patients who will be receiving subcutaneous maintenance doses: Coverage will be provided for 2 intravenous doses and 4 subcutaneous doses [see Entyvio SQ policy – Document Number: IC-0733]

# II. Dosing Limits

A. Quantity Limit (max daily dose) [NDC Unit]:

#### Loading Dose:

• Entyvio 300 mg single use vial: 1 vial at weeks 0, 2, & 6 (3 vials total per 42 days)

#### Maintenance Dose:

- Entyvio 300 mg single use vial: 1 vial every 4 weeks (28 days)
- B. Max Units (per dose and over time) [HCPCS Unit]:

#### Crohn's Disease and Ulcerative Colitis:

- Loading Dose: 300 billable units (300 mg) at weeks 0, 2, & 6
- <u>Maintenance Dose</u>: 300 billable units (300 mg) every 4 weeks

# Management of Immune Checkpoint Inhibitor-Related Diarrhea/Colitis

• 300 billable units (300 mg) at weeks 0, 2, & 6

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#### III. Initial Approval Criteria<sup>1</sup>

Coverage is provided in the following conditions:

- Patient is at least 18 years of age; AND
- Patient is up to date with all vaccinations, in accordance with current immunization guidelines, prior to initiating therapy; **AND**

#### Universal Criteria<sup>1</sup>

- Patient does not have an active infection, including clinically important localized infections; **AND**
- Patient has been evaluated and screened for the presence of latent tuberculosis (TB) infection prior to initiating treatment and will receive ongoing monitoring for presence of TB during treatment; **AND**
- Patient is not on concurrent treatment with another integrin receptor antagonist, TNFinhibitor, IL-inhibitor, biologic response modifier or other non-biologic agent (i.e., apremilast, tofacitinib, baricitinib, upadacitinib, abrocitinib, deucravacitinib, etc.); **AND**

#### Crohn's Disease † 1,2,15,16

- Physician has assessed baseline disease severity utilizing an objective measure/tool; AND
- Documented moderate to severe active disease; AND
  - Documented failure, contraindication, or ineffective response at maximum tolerated doses to a minimum 3-month trial of corticosteroids or immunomodulators (e.g., azathioprine, 6-mercaptopurine, or methotrexate, etc.); OR
  - Documented failure, contraindication, or ineffective response at maximum tolerated doses to a minimum 3-month trial on previous therapy with a TNF modifier such as adalimumab, certolizumab, or infliximab

#### Ulcerative Colitis † 1,12,18-20

- Physician has assessed baseline disease severity utilizing an objective measure/tool; AND
- Documented moderate to severe active disease; AND
  - Documented failure or ineffective response to a minimum 3-month trial of conventional therapy [aminosalicylates, corticosteroids, or immunomodulators (e.g., azathioprine, 6-mercaptopurine, methotrexate, etc.)] at maximum tolerated doses, unless there is a contraindication or intolerance to use; OR
  - Documented failure, contraindication, or ineffective response at maximum tolerated doses to a minimum 3-month trial on previous therapy with a TNF modifier such as adalimumab, golimumab, or infliximab

#### Management of Immune Checkpoint Inhibitor-Related Diarrhea/Colitis ‡ <sup>13,14</sup>



- Patient has been receiving therapy with an immune checkpoint inhibitor (e.g., nivolumab, pembrolizumab, atezolizumab, avelumab, durvalumab, cemiplimab, ipilimumab, tremelimumab, dostarlimab, retifanlimab, etc.); **AND** 
  - Patient has mild (G1) diarrhea or colitis related to their immunotherapy with persistent or progressive symptoms and a positive lactoferrin/calprotectin; **OR**
  - $\circ~$  Patient has moderate (G2) to severe (G3-4) diarrhea or colitis related to their immunotherapy

au FDA Approved Indication(s);  $\pm$  Compendia Recommended Indication(s);  $\Phi$  Orphan Drug

# IV. Renewal Criteria<sup>1</sup>

Coverage may be renewed based upon the following criteria:

- Patient continues to meet universal and indication-specific criteria as identified in section III; **AND**
- Absence of unacceptable toxicity from the drug. Examples of unacceptable toxicity include: anaphylaxis or other serious allergic, severe infusion-related or hypersensitivity reactions, severe infections, progressive multifocal leukoencephalopathy (PML), jaundice or other evidence of significant liver injury, etc.; **AND**

# Crohn's Disease 11,16

• Disease response as indicated by improvement in signs and symptoms compared to baseline such as endoscopic activity, number of liquid stools, presence and severity of abdominal pain, presence of abdominal mass, body weight compared to IBW, hematocrit, presence of extra intestinal complications, use of anti-diarrheal drugs, tapering or discontinuation of corticosteroid therapy, and/or an improvement on a disease activity scoring tool [e.g., an improvement on the Crohn's Disease Activity Index (CDAI) score or the Harvey-Bradshaw Index score].

# Ulcerative Colitis 9-11,20

• Disease response as indicated by improvement in signs and symptoms compared to baseline such as stool frequency, rectal bleeding, and/or endoscopic activity, tapering or discontinuation of corticosteroid therapy, and/or an improvement on a disease activity scoring tool [e.g., an improvement on the Ulcerative Colitis Endoscopic Index of Severity (UCEIS) score or the Mayo Score].

# Management of Immune Checkpoint Inhibitor-Related Diarrhea/Colitis ‡ <sup>13,14</sup>

• May not be renewed.

# V. Dosage/Administration <sup>1,13,17</sup>

Indication	Dose	
Crohn's Disease	Induction dose:	



Indication	Dose	
	• Administer 300 mg intravenously at weeks 0, 2, & 6	
	Maintenance dose:	
	• Administer 300 mg intravenously every 8 weeks thereafter	
	<b>***NOTE:</b> Requests for higher dosing must be reviewed according to the	
	dose escalation information below	
	Induction dose:	
	• Patients who will be receiving <u>intravenous</u> maintenance doses: Administer 300 mg intravenously at weeks 0, 2, & 6 <i>(see maintenance dosing below)</i>	
Ulcerative Colitis	• Patients who will be receiving <u>subcutaneous</u> maintenance doses: Administer 300 mg intravenously at weeks 0 and 2 <i>(see Entyvio SQ policy [Document Number: IC-0733] for maintenance dosing starting at week 6).</i>	
	Maintenance dose:	
	Administer 300 mg intravenously every 8 weeks thereafter	
	<b>***NOTE:</b> Requests for higher <u>intravenous</u> dosing must be reviewed according to the dose escalation information below	
Management of Immune Checkpoint Inhibitor- Related Diarrhea/Colitis	Administer 300 mg intravenously at weeks 0, 2, & 6	

- Dose escalation (up to the maximum dose and frequency specified below) may occur upon clinical review on a case-by-case basis provided that the patient has:
  - Shown an initial response to therapy; AND
  - $\circ$   $\;$  Received the three loading doses at the dose  $\underline{AND}$  interval specified above;  $\underline{AND}$
  - Received a minimum of one maintenance dose at the dose <u>AND</u> interval specified above; AND
  - Responded to therapy (by treatment week 14\*) with subsequent loss of response; AND
  - Dose escalation must not exceed the following limits:
    - 300 mg every 4 weeks
      - Coverage will be provided for 3 months with continued approval (as specified in Sections I & IV) contingent upon demonstration of clinical improvement and vedolizumab levels (if available)\*\*
        - Patients who do not regain response should discontinue therapy
        - Patients who are responding to therapy may continue with their current dosing\*\*

#### \*<u>Note</u>:

- Request for dose escalation prior to week 14 will be evaluated considering the patient's clinical picture regarding severity of inflammation, factors which may result in subtherapeutic response to standard dosing (e.g., obesity, hypoalbuminemia, prior TNF-I exposure), timing of response and breakthrough/loss of response, AND one of the following:
  - vedolizumab trough (if available)\*\* at week 14 is <14 micrograms/mL; OR</li>
  - CRP elevation or calprotectin >150



\*\*vedolizumab trough levels must be obtained (if this is a covered test under the benefit).

- Patients whose trough is 14-20 micrograms/mL may continue with 300 mg every 4 weeks.
- Patients with a trough >20 micrograms/mL must increase the interval between administrations from 4 weeks to 6 weeks. Response should be assessed after receipt of 3 doses at this every 6-week interval. Those patients demonstrating loss of response may then decrease the interval back to 300 mg every 4 weeks.
- Patients whose trough is <14 micrograms/mL are candidates to decrease the interval between administrations from 8 weeks to 4 weeks

# VI. Billing Code/Availability Information

HCPCS Code:

- J3380 Injection, vedolizumab, 1 mg; 1 billable unit = 1 mg NDC:
- Entyvio 300 mg single use vial: 67464-0300-xx

# VII. References

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# **Appendix 1 – Covered Diagnosis Codes**

ICD-10	ICD-10 Description	
K50.00	Crohn's disease of small intestine without complications	
K50.011	Crohn's disease of small intestine with rectal bleeding	
K50.012	Crohn's disease of small intestine with intestinal obstruction	
K50.013	Crohn's disease of small intestine with fistula	
K50.014	Crohn's disease of small intestine with abscess	
K50.018	Crohn's disease of small intestine with other complication	
K50.019	Crohn's disease of small intestine with unspecified complications	
K50.10	Crohn's disease of large intestine without complications	
K50.111	Crohn's disease of large intestine with rectal bleeding	
K50.112	Crohn's disease of large intestine with intestinal obstruction	
K50.113	Crohn's disease of large intestine with fistula	
K50.114	Crohn's disease of large intestine with abscess	
K50.118	Crohn's disease of large intestine with other complication	
K50.119	Crohn's disease of large intestine with unspecified complications	
K50.80	Crohn's disease of both small and large intestine without complications	
K50.811	Crohn's disease of both small and large intestine with rectal bleeding	
K50.812	Crohn's disease of both small and large intestine with intestinal obstruction	
K50.813	Crohn's disease of both small and large intestine with fistula	
K50.814	Crohn's disease of both small and large intestine with abscess	
K50.818	Crohn's disease of both small and large intestine with other complication	
K50.819	Crohn's disease of both small and large intestine with unspecified complications	
K50.90	Crohn's disease, unspecified, without complications	
K50.911	Crohn's disease, unspecified, with rectal bleeding	
K50.912	Crohn's disease, unspecified, with intestinal obstruction	
K50.913	Crohn's disease, unspecified, with fistula	
K50.914	Crohn's disease, unspecified, with abscess	
K50.918	Crohn's disease, unspecified, with other complication	
K50.919	Crohn's disease, unspecified, with unspecified complications	
K51.00	Ulcerative (chronic) pancolitis without complications	
K51.011	Ulcerative (chronic) pancolitis with rectal bleeding	
K51.012	Ulcerative (chronic) pancolitis with intestinal obstruction	
K51.013	Ulcerative (chronic) pancolitis with fistula	
K51.014	Ulcerative (chronic) pancolitis with abscess	

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ICD-10	ICD-10 Description	
K51.018	Ulcerative (chronic) pancolitis with other complication	
K51.019	Ulcerative (chronic) pancolitis with unspecified complications	
K51.20	Ulcerative (chronic) proctitis without complications	
K51.211	Ulcerative (chronic) proctitis with rectal bleeding	
K51.212	Ulcerative (chronic) proctitis with intestinal obstruction	
K51.213	Ulcerative (chronic) proctitis with fistula	
K51.214	Ulcerative (chronic) proctitis with abscess	
K51.218	Ulcerative (chronic) proctitis with other complication	
K51.219	Ulcerative (chronic) proctitis with unspecified complications	
K51.30	Ulcerative (chronic) rectosigmoiditis without complications	
K51.311	Ulcerative (chronic) rectosigmoiditis with rectal bleeding	
K51.312	Ulcerative (chronic) rectosigmoiditis with intestinal obstruction	
K51.313	Ulcerative (chronic) rectosigmoiditis with fistula	
K51.314	Ulcerative (chronic) rectosigmoiditis with abscess	
K51.318	Ulcerative (chronic) rectosigmoiditis with other complication	
K51.319	Ulcerative (chronic) rectosigmoiditis with unspecified complications	
K51.50	Left sided colitis without complications	
K51.511	Left sided colitis with rectal bleeding	
K51.512	Left sided colitis with intestinal obstruction	
K51.513	Left sided colitis with fistula	
K51.514	Left sided colitis with abscess	
K51.518	Left sided colitis with other complication	
K51.519	Left sided colitis with unspecified complications	
K51.80	Other ulcerative colitis without complications	
K51.811	Other ulcerative colitis with rectal bleeding	
K51.812	Other ulcerative colitis with intestinal obstruction	
K51.813	Other ulcerative colitis with fistula	
K51.814	Other ulcerative colitis with abscess	
K51.818	Other ulcerative colitis with other complication	
K51.819	Other ulcerative colitis with unspecified complications	
K51.90	Ulcerative colitis, unspecified, without complications	
K51.911	Ulcerative colitis, unspecified with rectal bleeding	
K51.912	Ulcerative colitis, unspecified with intestinal obstruction	
K51.913	Ulcerative colitis, unspecified with fistula	
K51.914	Ulcerative colitis, unspecified with abscess	

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ICD-10	ICD-10 Description	
K51.918	Ulcerative colitis, unspecified with other complication	
K51.919	Ulcerative colitis, unspecified with unspecified complications	
K52.1	Toxic gastroenteritis and colitis	
R19.7	Diarrhea, unspecified	

# Appendix 2 – Centers for Medicare and Medicaid Services (CMS)

Medicare coverage for outpatient (Part B) drugs is outlined in the Medicare Benefit Policy Manual (Pub. 100-2), Chapter 15, §50 Drugs and Biologicals. In addition, National Coverage Determination (NCD), Local Coverage Determinations (LCDs), and Local Coverage Article (LCAs) may exist and compliance with these policies is required where applicable. They can be found at: <a href="https://www.cms.gov/medicare-coverage-database/search.aspx">https://www.cms.gov/medicare-coverage-database/search.aspx</a>. Additional indications may be covered at the discretion of the health plan.

	Medicare Part B Administrative Contractor (MAC) Jurisdictions				
Jurisdiction	Applicable State/US Territory	Contractor			
E (1)	CA, HI, NV, AS, GU, CNMI	Noridian Healthcare Solutions, LLC			
F (2 & 3)	AK, WA, OR, ID, ND, SD, MT, WY, UT, AZ	Noridian Healthcare Solutions, LLC			
5	KS, NE, IA, MO	Wisconsin Physicians Service Insurance Corp (WPS)			
6	MN, WI, IL	National Government Services, Inc. (NGS)			
H (4 & 7)	LA, AR, MS, TX, OK, CO, NM	Novitas Solutions, Inc.			
8	MI, IN	Wisconsin Physicians Service Insurance Corp (WPS)			
N (9)	FL, PR, VI	First Coast Service Options, Inc.			
J (10)	TN, GA, AL	Palmetto GBA, LLC			
M (11)	NC, SC, WV, VA (excluding below)	Palmetto GBA, LLC			
L (12)	DE, MD, PA, NJ, DC (includes Arlington & Fairfax counties and the city of Alexandria in VA)	Novitas Solutions, Inc.			
K (13 & 14)	NY, CT, MA, RI, VT, ME, NH	National Government Services, Inc. (NGS)			
15	KY, OH	CGS Administrators, LLC			

#### Medicare Part B Covered Diagnosis Codes (applicable to existing NCD/LCD/LCA): N/A



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# Nondiscrimination & Language Access Policy



Discrimination is Against the Law. Aspirus Health Plan, Inc. complies with applicable Federal civil rights laws and does not discriminate on the basis of race, color, national origin, age, disability, or sex, (including sex characteristics, including intersex traits; pregnancy or related conditions; sexual orientation, gender identity and sex stereotypes), consistent with the scope of sex discrimination described at 45 CFR § 92.101(a)(2). Aspirus Health Plan, Inc. does not exclude people or treat them less favorably because of race, color, national origin, age, disability, or sex.

Aspirus Health Plan, Inc.:

Provides people with disabilities reasonable modifications and free appropriate auxiliary aids and services to communicate effectively with us, such as:

- Qualified sign language interpreters.

- Written information in other formats (large print, audio, accessible electronic formats, other formats).

Provides free language assistance services to people whose primary language is not English, which may include:

- Qualified interpreters.
- Information written in other languages.

If you need reasonable modifications, appropriate auxiliary aids and services, or language assistance services, contact the Nondiscrimination Grievance Coordinator at the address, phone number, fax number, or email address below.

If you believe that Aspirus Health Plan, Inc. has failed to provide these services or discriminated in another way on the basis of race, color, national origin, age, disability, or sex, you can file a grievance with:

Nondiscrimination Grievance Coordinator Aspirus Health Plan, Inc. PO Box 1890 Southampton, PA 18966-9998 Phone: 1-866-631-5404 (TTY: 711) Fax: 763-847-4010 Email: customerservice@aspirushealthplan.com

You can file a grievance in person or by mail, fax, or email. If you need help filing a grievance, the Nondiscrimination Grievance Coordinator is available to help you.

*You* can also file a civil rights complaint with the U.S. Department of Health and Human Services, Office for Civil Rights, electronically through the Office for Civil Rights Complaint Portal, available at https://ocrportal.hhs.gov/ocr/portal/lobby.jsf, or by mail or phone at:

U.S. Department of Health and Human Services 200 Independence Avenue, SW Room 509F, HHH Building Washington, D.C. 20201 1.800.368.1019, 800.537.7697 (TDD)

Complaint forms are available at http://www.hhs.gov/ocr/office/file/index.html. This notice is available at Aspirus Health Plan, Inc.'s website: https://aspirushealthplan.com/webdocs/70021-AHP-NonDiscrim\_Lang-Assist-Notice.pdf.

#### Language Assistance Services

Albanian: KUJDES: Nëse flitni shqip, për ju ka në dispozicion shërbime të asistencës gjuhësore, pa pagesë. Telefononi në 1-800-332-6501 (TTY: 711). (711: (تق هاتف الصم والبك) 1-800-332-6501 تنبيه : إذا كنت تتحدث اللغة العربية، فإن خدمات المساعدة اللغوية متاحة لك مجاناً التصل بن اعلى رقم الهاتف ال-800-332-6501 (TTY: 711). French: ATTENTION: Si vous parlez français, des services d'aide linguistique vous sont proposés gratuitement. Appelez le 1-800-332-6501 (ATS: 711). German: ACHTUNG: Wenn Sie Deutsch sprechen, stehen Ihnen kostenlos sprachliche Hilfsdienstleistungen zurVerfügung. Rufnummer: 1-800-332-6501 (TTY: 711).

Hindi: \_यान द\_: य\_द आप िहंदी बोलते ह\_ तो आपके िलए मु\_त म\_ भाषा सहायता सेवाएं उपल\_ध ह\_11-800-332-6501 (TTY: 711) पर कॉल कर\_। Hmong: LUS CEEV: Yog tias koj hais lus Hmoob, cov kev pab txog lus, muaj kev pab dawb rau koj. Hu rau 1-800-332-6501 (TTY: 711).

Korean: 주의: 한국어를 사용하시는 경우, 언어 지원 서비스를 무료로 이용하실 수 있습니다.1-800-332-6501 (TTY: 711)번으로 전화해 주십시오.

Polish: UWAGA: Jeżeli mówisz po polsku, możesz skorzystać z bezpłatnej pomocy językowej. Zadzwoń pod numer1-800-332-6501 (TTY: 711). Russian: ВНИМАНИЕ: Если вы говорите на русском языке, то вам доступны бесплатные услуги перевода.Звоните 1-800-332-6501 (телетайп: 711).

Spanish: ATENCIÓN: si habla español, tiene a su disposición servicios gratuitos de asistencia lingüística. Llame all-800-332-6501 (TTY: 711). Tagalog: PAUNAWA: Kung nagsasalita ka ng Tagalog, maaari kang gumamit ng mga serbisyo ng tulong sa wika nangwalang bayad. Tumawag sa 1-800-332-6501 (TTY: 711).

Traditional Chinese: 注意: 如果您使用繁體中文, 您可以免費獲得語言援助服務。請 致電 1-800-332-6501 (TTY: 711)

Vietnamese: CHÚ Ý: Nếu bạn nói Tiếng Việt, có các dịch vụ hỗ trợ ngôn ngữ miễn phí dành cho bạn. Gọi số 1-800-332-6501 (TTY: 711). Pennsylvania Dutch: Wann du Deitsch (Pennsylvania German / Dutch) schwetzscht, kannscht du mitaus Koschte ebbergricke, ass dihr helft mit die englisch Schprooch. Ruf selli Nummer uff: Call 1-800-332-6501 (TTY: 711).

Lao: ໂປດຊາບ: ຖ້າວ່າ ທ່ານເວົ້າພາສາ ລາວ, ການບໍລິການຊ່ວຍເຫຼືອດ້ານພາສາ,ໂດຍບໍ່ເສັງຄ່າ, ແມ່ນມີພ້ອມໃຫ້ທ່ານ. ໂທຣ 1-800-332-6501 (TTY: 711).