AvMed

PHARMACY PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

<u>Directions</u>: <u>The prescribing physician must sign and clearly print name (preprinted stamps not valid)</u> on this request. All other information may be filled in by office staff; <u>fax to 1-305-671-0200</u>. No additional phone calls will be necessary if all information (<u>including phone and fax #s</u>) on this form is correct. <u>If the information provided is not complete, correct, or legible, the authorization process can be delayed.</u>

Drug Requested: Zymfentra[™] (infliximab-dyyb) (**Pharmacy**)

| MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete. | | |
|--|--------------------------------------|--|
| Member Name: | | |
| Member AvMed #: | Date of Birth: | |
| Prescriber Name: | | |
| Prescriber Signature: | | |
| Office Contact Name: | | |
| Phone Number: | Fax Number: | |
| NPI #: | | |
| DRUG INFORMATION: Authoriz | zation may be delayed if incomplete. | |
| Drug Form/Strength: | | |
| Dosing Schedule: | Length of Therapy: | |
| Diagnosis: | ICD Code, if applicable: | |
| Weight (if applicable): | Date weight obtained: | |

<u>ATTENTION</u>: All patients must complete an intravenous induction regimen with an infliximab product before starting Zymfentra. IV induction (loading dose) for treatment of Crohn's disease & ulcerative colitis can only be billed under the <u>MEDICAL BENEFIT</u>. Provider please note: Renflexis[®] (infliximab-adba) is the Health Plan's **PREFFERED** infliximab product, O5104. NDC: 00006430501/02; 78206016201/99

Quantity Limits: 2 syringes/pens per 28 days

Adult Dosing:

- Zymfentra is indicated as maintenance treatment only, starting at Week 10 and thereafter.
 - All patients must complete an intravenous induction regimen with an infliximab product before starting Zymfentra. For induction dosing information, see the corresponding full prescribing information for the chosen infliximab product
- Zymfentra is for subcutaneous use only
- Maintenance dosage starting at Week 10 and thereafter: 120 mg subcutaneously once every two weeks.
- To switch patients who are responding to maintenance therapy with an infliximab product administered intravenously, administer the first subcutaneous dose of Zymfentra in place of the next scheduled intravenous infusion and every two weeks thereafter

NOTE: The Health Plan considers the use of concomitant therapy with more than one biologic immunomodulator (e.g., Dupixent, Entyvio, Humira, Rinvoq, Stelara) prescribed for the same or different indications to be experimental and investigational. Safety and efficacy of these combinations has **NOT** been established and will **NOT** be permitted.

| • Wi | ill th | ne member be discontinuing a previously | prescribed biologic if approved for requested Yes | d medication? OR □ No |
|-------|---|---|--|------------------------|
| | | please list the medication that will be disc al along with the corresponding effective | continued and the medication that will be inidate. | tiated upon |
| Mo | edic | ation to be discontinued: | Effective date: | |
| Mo | edic | ation to be initiated: | Effective date: | |
| | | | | |
| suppo | ort e | | at apply. All criteria must be met for approvading lab results, diagnostics, and/or chart no | |
| | | | | |
| | | ntenance Dose – 120 mg administ er than Week 10, and then every | ered by subcutaneous injection start 2 weeks thereafter | ting at no |
| Autl | hor | ization Criteria: To be reviewed f | or approval under the pharmacy be | enefit |
| | Me | ember is 18 years of age or older | | |
| | Me | ember has ONE of the following diagnose | es: | |
| | | Moderate-to-severe active Crohn's dise | ase | |
| | | Moderate-to-severe active ulcerative co | litis | |
| | ☐ Prescribed by or in consultation with a Gastroenterologist | | | |
| | Me | ember meets ONE of the following: | | |
| | | Member has tried and failed budesonide | or high dose steroids (40-60 mg prednisone) |) |
| | | Member has tried and failed at least ON months | \mathbf{E} of the following DMARD therapies for at | least three (3) |
| | | □ 5-aminosalicylates (balsalazide, olsa | lazine, sulfasalazine) | |
| | | ☐ oral mesalamine (Apriso, Asacol/HD |), Delzicol, Lialda, Pentasa) | |
| | | Member has already started on or is curr IV product and is requesting continuatio | ently undergoing an induction regimen with n of therapy with Zymfentra | an infliximab |

(Continued on next page)

| Authorization Criteria: To be reviewed for one-time approval under the medical benefit Medication will be used as induction therapy Medication being provided by: Location/site of drug administration: NPI or DEA # of administering location: Member to receive FDA approved loading dose of Renflexis® to be administered by intravenous infusion at a dose of 5 mg/kg at 0, 2 & 6 weeks | du | duction Dose for <u>PREFERED</u> Renflexis [®] only (If required) – One time approval for irration of 2 months. Member to receive up to three (3) IV infusion doses, at a dose 5 mg/kg at 0, 2 & 6 weeks |
|--|-------------|---|
| □ Medication being provided by: □ Location/site of drug administration: □ NPI or DEA # of administering location: □ Member to receive FDA approved loading dose of Renflexis® to be administered by intravenous infusion | <u>Auth</u> | orization Criteria: To be reviewed for one-time approval under the medical benefit |
| □ Location/site of drug administration: □ NPI or DEA # of administering location: □ Member to receive FDA approved loading dose of Renflexis® to be administered by intravenous infusion | | Medication will be used as induction therapy |
| □ NPI or DEA # of administering location: □ Member to receive FDA approved loading dose of Renflexis® to be administered by intravenous infusion | | Medication being provided by: |
| ☐ Member to receive FDA approved loading dose of Renflexis® to be administered by intravenous infusion | | □ Location/site of drug administration: |
| | | □ NPI or DEA # of administering location: |
| | | Member to receive FDA approved loading dose of Renflexis® to be administered by intravenous infusion at a dose of 5 mg/kg at 0, 2 & 6 weeks |
| Medication being provided by Specialty Pharmacy – Proprium Rx | Med | lication being provided by Specialty Pharmacy – Proprium Rx |

**Use of samples to initiate therapy does not meet step edit/preauthorization criteria. **

*Previous therapies will be verified through pharmacy paid claims or submitted chart notes. *