

PATIENT INFORMATION

PATIENT NAME: _____ PHONE: _____

DATE OF BIRTH: _____ SEX: M F HEIGHT: _____ WEIGHT: _____ LBS KG

ALLERGIES: _____ PREFERRED CLINIC: _____

REFERRAL STATUS: NEW REFERRAL ORDER CHANGE ORDER RENEWAL

DIAGNOSIS & CLINICAL DOCUMENTATION

- J45.50 Severe persistent asthma, uncomplicated
- J45.51 Severe persistent asthma with (acute) exacerbation
- J82.83 Eosinophilic Asthma
- J45.51 Polyarteritis with lung involvement [EGPA/Churg-Strauss]
- ICD-10 CODE: _____ DESCRIPTION: _____

REQUIRED DOCUMENTATION

- Insurance Information
- List of Medications
- Tried & Failed Therapies
- Most Recent History & Physical
- Lab results and/or Pulmonary Function Tests to support diagnosis
- Blood Eosinophil Level/CBC

MEDICATION ORDER

- LOADING:** Fasenra® (benralizumab) 30mg via subcutaneous injection every 4 weeks initial 3 doses
- MAINTENANCE:** Fasenra® (benralizumab) 30mg via subcutaneous injection every 8 weeks
- REFILL X 12 MONTHS UNLESS OTHERWISE NOTED HERE:** _____

Provider Attestation of need for HCP administration

- _____
INITIALS Patient has experienced severe hypersensitivity reactions to Fasenra within the past 6 months and requires administration and direct monitoring by a healthcare professional.
- _____
INITIALS Patient or caregiver are not competent or are physically unable to administer the Fasenra formulation labeled for self-administration.
- _____
INITIALS Patient has history of uncontrolled disease and in the clinical opinion of the ordering provider, it is not advisable to try the self-administered formulation of Fasenra.
- _____
INITIALS The circumstances and location for self-administration are not adequate for the potential treatment of anaphylaxis.
- _____
INITIALS Patient's weight is such that in the clinical opinion of the ordering provider, it is not advisable to try the self-administered formulation of Fasenra

Patient to be observed for 30 minutes following the first injection and then for 15 minutes following each subsequent injection. Administer per protocol. In the event of an adverse reaction occurring in the infusion clinic, utilize the Immersiv Health adverse reaction protocol.

PRESCRIBER INFORMATION

PROVIDER NAME: _____ NPI #: _____

EMAIL: _____ PHONE: _____ FAX: _____

ADDRESS (INCLUDE CITY, STATE, ZIP): _____

SUPERVISING PHYSICIAN: _____ CONTACT NAME: _____
(IF APPLICABLE)

SIGNATURE: _____ DATE: _____
(NO STAMPS)

SUBSTITUTION PERMITTED

DISPENSE AS WRITTEN