

## **Pharmacy Prior Authorization Form** Fax completed form to: 877.974.4411 toll free, or 616.942.8206 ☐ Commercial (Traditional) ☐ Commercial (Individual/Optimized) This form applies to: Medicaid ☐ Urgent (life threatening) ☐ Non-Urgent (standard review) This request is: Urgent means the standard review time may seriously jeopardize the life or health of the patient or the patient's ability to regain maximum function. Fluorouracil 0.5% cream Member Last Name: First Name: DOB: \_\_\_\_\_ Gender: Primary Care Physician: Prov. Phone: \_\_\_\_\_ Prov. Fax: \_\_\_\_\_ Requesting Provider: Provider Address: Provider NPI: \_\_\_\_\_ Contact Name: \_\_\_\_\_ Provider Signature: **Product Information** ☐ New request ☐ Continuation request Start date (or date of next dose): \_\_\_\_\_\_ Date of last dose (if applicable): \_\_\_\_\_ ☐ Fluorouracil 0.5% cream Drug product: Dosing frequency: **Precertification Requirements** Before this drug is covered, the patient must meet all of the following requirements (initial approval 3 months): 1. Must have a diagnosis of actinic keratosis; AND 2. Must have had an inadequate response or intolerance to office-based treatments (liquid nitrogen cryotherapy, surgical curettage) or have been considered and ruled out due to nature/number of lesions or limited resources to provide such treatments; AND 3. Must have had an inadequate response to a full treatment or intolerance/contraindication to a trial of imiguimod. For continuation, patient must have met the following requirements (continued approval 3 months): 1. Must have documentation that there is a recurrence of active lesions and treatment with another course of therapy is required. 2. If there is no demonstrable clinically significant improvement in condition continued treatment will not be approved. Note: Authorization for indications, dosing, or a route of administration not approved by the Food and Drug Administration (FDA) or recognized in CMSaccepted compendia (e.g. DrugDex, AHFS, U.S. Pharmacopeia, and also Clinical Pharmacology for oncology indications only) require supporting evidence for coverage. Please provide two published peer-reviewed literature articles supporting the appropriateness of the drug, the dosing of the drug, or the route of administration to be used for the identified indication. New request **Priority Health Precertification Documentation** A. What condition is this drug being requested for? ☐ Actinic keratosis ☐ Other – the patient's condition is:



	Rationale for use:
В.	Has the patient had an inadequate response to office-based treatments (liquid nitrogen cryotherapy, surgical curettage)?  Yes, describe: No, reason:
C.	Has the patient had a trial with a full treatment course of imiquimod?  Yes, Dates:  No, rationale:
	No, rationale.
Request to continue a previously authorized approval Priority Health Precertification Documentation	
Α.	What condition is this drug being requested for?  Actinic keratosis  Other – the patient's condition is:  Rationale for use:
В.	Did the patient have a clinically significant improvement in condition after initiation of fluorouracil?  Yes No
C.	Has there been a recurrence of active lesions prompting another course of therapy?  ☐ Yes ☐ No

## **Additional information**

**Note:** Fluorouracil 0.5% cream is limited to 30 grams every 30 days. Approval will be granted in 3 month intervals (new request required every 3 months for continuation).