

Member Name: \_\_\_\_\_ Date of Birth: \_\_\_\_\_ Member ID#: \_\_\_\_\_

**Drug Information**

Pharmacy billing (NDC: \_\_\_\_\_) Start Date (or date of next dose): \_\_\_\_\_  
Dose: \_\_\_\_\_ Regimen: \_\_\_\_\_

**Billing Provider Information**

Provider NPI: \_\_\_\_\_ Provider Name: \_\_\_\_\_  
Provider Phone: \_\_\_\_\_ Provider Fax: \_\_\_\_\_

**Prescriber Information**

Prescriber NPI: \_\_\_\_\_ Prescriber Name: \_\_\_\_\_  
Prescriber Phone: \_\_\_\_\_ Prescriber Fax: \_\_\_\_\_ Specialty: \_\_\_\_\_

**Criteria**

**\*Page 1 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays.\***

**For Initial Authorization (Initial approval will be for the duration of 6 months):**

1. Please indicate the diagnosis and information:

**Unresectable or Metastatic Melanoma**

- A. Does member have BRAF V600E or V600K mutation? Yes \_\_\_\_\_ No \_\_\_\_\_
- B. Does member have wild-type BRAF melanoma? Yes \_\_\_\_\_ No \_\_\_\_\_
- C. Will trametinib be used as a single-agent? Yes \_\_\_\_\_ No \_\_\_\_\_
- D. Will trametinib be used in combination with dabrafenib (Tafinlar®)? Yes \_\_\_\_\_ No \_\_\_\_\_
- E. Will trametinib be used as first-line therapy? Yes \_\_\_\_\_ No \_\_\_\_\_
- F. Will trametinib be used as second-line or subsequent therapy? Yes \_\_\_\_\_ No \_\_\_\_\_
  - i. If using as second-line or subsequent therapy, please indicate member's ECOG performance status (0-5): \_\_\_\_\_
- G. Has member received prior BRAF inhibitor therapy (e.g., dabrafenib, vemurafenib)? Yes \_\_\_\_\_ No \_\_\_\_\_
  - i. If member has received prior BRAF inhibitor therapy, please indicate the following:
    - a. Was member intolerant to prior BRAF inhibitor therapy? Yes \_\_\_\_\_ No \_\_\_\_\_
    - b. Was there evidence of progression on prior BRAF inhibitor therapy? Yes \_\_\_\_\_ No \_\_\_\_\_

**Non-Small Cell Lung Cancer (NSCLC)**

- A. Is the diagnosis refractory or metastatic disease? Yes \_\_\_\_\_ No \_\_\_\_\_
- B. Does member have BRAF V600E or V600K mutation? Yes \_\_\_\_\_ No \_\_\_\_\_
- C. Does member have wild-type BRAF NSCLC? Yes \_\_\_\_\_ No \_\_\_\_\_
- D. Will trametinib be used in combination with dabrafenib (Tafinlar®)? Yes \_\_\_\_\_ No \_\_\_\_\_

**Anaplastic Thyroid Cancer (ATC)**

- A. Is the diagnosis locally advanced or metastatic disease? Yes \_\_\_\_\_ No \_\_\_\_\_
- B. Does member have BRAF V600E mutation? Yes \_\_\_\_\_ No \_\_\_\_\_
- C. Will trametinib be used in combination with dabrafenib (Tafinlar®)? Yes \_\_\_\_\_ No \_\_\_\_\_
- D. Are there any satisfactory locoregional treatment options for the member? Yes \_\_\_\_\_ No \_\_\_\_\_

**Serous Ovarian Cancer**

- A. Is diagnosis persistent or recurrent low-grade serous ovarian cancer? Yes \_\_\_\_\_ No \_\_\_\_\_
- B. Will trametinib be used as immediate treatment for serially rising CA-125 in members who previously received chemotherapy? Yes \_\_\_\_\_ No \_\_\_\_\_
- C. Will trametinib be used for disease progression on primary, maintenance, or recurrence therapy?  
Yes \_\_\_\_\_ No \_\_\_\_\_
- D. Will trametinib be used for stable or persistent disease (if member is not on maintenance therapy)?  
Yes \_\_\_\_\_ No \_\_\_\_\_
- E. Will trametinib be used for complete remission and relapse after completing chemotherapy?  
Yes \_\_\_\_\_ No \_\_\_\_\_

PLEASE PROVIDE THE INFORMATION REQUESTED AND RETURN TO:

University of Oklahoma College of Pharmacy  
Pharmacy Management Consultants  
Product Based Prior Authorization Unit  
Fax: 1-800-224-4014  
Phone: 1-800-522-0114 Option 4

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**Mekinist® (Trametinib) Prior Authorization Form**

Member Name: \_\_\_\_\_ Date of Birth: \_\_\_\_\_ Member ID#: \_\_\_\_\_

**Criteria****\*Page 2 of 2– Please complete and return all pages. Failure to complete all pages will result in processing delays.\*****For Initial Authorization, continued:**

1. Please indicate the diagnosis and information, continued:

 **If diagnosis is not listed on the previous page, please indicate diagnosis:** \_\_\_\_\_

Additional Information: \_\_\_\_\_

**For Continued Authorization:**

1. Date of last dose: \_\_\_\_\_

2. Does member have any evidence of progressive disease while on trametinib? Yes \_\_\_\_\_ No \_\_\_\_\_

3. Has the member experienced any adverse drug reactions related to trametinib therapy? Yes \_\_\_\_\_ No \_\_\_\_\_

*If yes, please specify adverse reactions:* \_\_\_\_\_

Additional Information: \_\_\_\_\_

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Please complete and return all pages. Failure to complete all pages will result in processing delays.

Prescriber Signature: \_\_\_\_\_ Date: \_\_\_\_\_

***I certify that the indicated treatment is medically necessary and all information is true and correct to the best of my knowledge. Please do not send in chart notes. Specific information will be requested if necessary.***PLEASE PROVIDE THE INFORMATION REQUESTED AND RETURN TO:University of Oklahoma College of Pharmacy  
Pharmacy Management Consultants  
Product Based Prior Authorization Unit  
Fax: 1-800-224-4014  
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