



Coding Resource

It is important to note that the codes identified below are examples only. Each provider is responsible for ensuring all coding is accurate and documented in the medical record based on the condition of the patient. The use of the following codes does not guarantee reimbursement.

National Drug Code (NDC)

10-digit NDC

Dosage	Code
40 mg TABLETS – 30 ct BOTTLE	0310-1349-30
80 mg TABLETS — 30 ct BOTTLE	0310-1350-30

11-digit NDC

Dosage	Code
40 mg TABLETS – 30 ct BOTTLE	00310-1349-30
80 mg TABLETS — 30 ct BOTTLE	00310-1350-30

Current Procedural Terminology (CPT)¹

Code	Description
81235	EGFR (epidermal growth factor receptor) (eg, non-small cell lung cancer) gene analysis, common variants (eg, exon 19 LREA deletion, L858R, T790M, G719A, G719S, L861Q)

Healthcare Common Procedure Coding System (HCPCS)²

Code	Description
G0452	Molecular pathology procedure; physician interpretation and report

Diagnosis Codes³

ICD-10-CM	Description
C34.00	Malignant neoplasm of unspecified main bronchus
C34.01	Malignant neoplasm of right main bronchus
C34.02	Malignant neoplasm of left main bronchus
C34.10	Malignant neoplasm of upper lobe, unspecified bronchus or lung
C34.11	Malignant neoplasm of upper lobe, right bronchus or lung
C34.12	Malignant neoplasm of upper lobe, left bronchus or lung
C34.20	Malignant neoplasm of middle lobe, bronchus or lung
C34.30	Malignant neoplasm of lower lobe, unspecified bronchus or lung
C34.31	Malignant neoplasm of lower lobe, right bronchus or lung

ICD-10-CM	Description
C34.32	Malignant neoplasm of lower lobe, left bronchus or lung
C34.80	Malignant neoplasm of overlapping sites of unspecified bronchus and lung
C34.81	Malignant neoplasm of overlapping sites of right bronchus and lung
C34.82	Malignant neoplasm of overlapping sites of left bronchus and lung
C34.90	Malignant neoplasm of unspecified part of unspecified bronchus or lung
C34.91	Malignant neoplasm of unspecified part of right bronchus or lung
C34.92	Malignant neoplasm of unspecified part of left bronchus or lung

Please read Important Safety Information on page 2 and accompanying complete Prescribing Information, including Patient Information.

For more information, call AstraZeneca Access 360™ at 1-844-ASK-A360, Monday through Friday, 8 AM to 8 PM ET.



1-844-ASK-A360 (1-844-275-2360)



1-844-FAX-A360 (1-844-329-2360)



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INDICATIONS

- TAGRISSO is indicated for the first-line treatment of patients with metastatic non-small cell lung cancer (NSCLC)
 whose tumors have epidermal growth factor receptor (EGFR) exon 19 deletions or exon 21 L858R mutations, as
 detected by an FDA-approved test
- TAGRISSO is indicated for the treatment of patients with metastatic EGFR T790M mutation-positive NSCLC, as detected by an FDA-approved test, whose disease has progressed on or after EGFR tyrosine kinase inhibitor (TKI) therapy

Important Safety Information About TAGRISSO

- There are no contraindications for TAGRISSO
- Interstitial lung disease (ILD)/pneumonitis occurred in 3.9% of the 1142 TAGRISSO-treated patients; 0.4% of
 cases were fatal. Withhold TAGRISSO and promptly investigate for ILD in patients who present with worsening
 of respiratory symptoms which may be indicative of ILD (eg, dyspnea, cough and fever). Permanently discontinue
 TAGRISSO if ILD is confirmed
- Heart rate-corrected QT (QTc) interval prolongation occurred in TAGRISSO-treated patients. Of the 1142 TAGRISSO-treated patients in clinical trials, 0.9% were found to have a QTc > 500 msec, and 3.6% of patients had an increase from baseline QTc > 60 msec. No QTc-related arrhythmias were reported. Conduct periodic monitoring with ECGs and electrolytes in patients with congenital long QTc syndrome, congestive heart failure, electrolyte abnormalities, or those who are taking medications known to prolong the QTc interval. Permanently discontinue TAGRISSO in patients who develop QTc interval prolongation with signs/symptoms of life-threatening arrhythmia
- Cardiomyopathy occurred in 2.6% of the 1142 TAGRISSO-treated patients; 0.1% of cardiomyopathy cases were fatal. A decline in left ventricular ejection fraction (LVEF) ≥10% from baseline and to <50% LVEF occurred in 3.9% of 908 patients who had baseline and at least one follow-up LVEF assessment. Conduct cardiac monitoring, including assessment of LVEF at baseline and during treatment, in patients with cardiac risk factors. Assess LVEF in patients who develop relevant cardiac signs or symptoms during treatment. For symptomatic congestive heart failure, permanently discontinue TAGRISSO
- Keratitis was reported in 0.7% of 1142 patients treated with TAGRISSO in clinical trials. Promptly refer patients with signs and symptoms suggestive of keratitis (such as eye inflammation, lacrimation, light sensitivity, blurred vision, eye pain and/or red eye) to an ophthalmologist
- Verify pregnancy status of females of reproductive potential prior to initiating TAGRISSO. Advise pregnant women
 of the potential risk to a fetus. Advise females of reproductive potential to use effective contraception during
 treatment with TAGRISSO and for 6 weeks after the final dose. Advise males with female partners of reproductive
 potential to use effective contraception for 4 months after the final dose
- Most common adverse reactions (≥20%) were diarrhea, rash, dry skin, nail toxicity, stomatitis, fatigue and decreased appetite

Please see accompanying complete <u>Prescribing Information</u> including Patient Information.

You are encouraged to report negative side effects of AstraZeneca prescription drugs by calling 1-800-236-9933. If you prefer to report these to the FDA, either visit www.FDA.gov/medwatch or call 1-800-FDA-1088.

References: 1. American Medical Association. *CPT*® 2017 Professional Edition. Chicago, IL: American Medical Association; 2017. 2. Centers for Medicare & Medicaid Services. HCPCS Release & Code Sets. https://www.cms.gov/Medicare/Coding/HCPCSReleaseCodeSets/Alpha-Numeric-HCPCS.html. Accessed December 1, 2017. 3. American Medical Association. *ICD-10-CM* 2017: The Complete Official Codebook. Chicago, IL: American Medical Association; 2017.

