Afatinib

ErbB-1, -2 and -4 tyrosine kinase inhibitor

Afatinib is an irreversible kinase inhibitor and binds to the kinase domains of EGFR (ErbB1), HER2 (ErbB2), and HER4 (ErbB4) to inhibit tyrosine kinase autophosphorylation. This results in a downregulation of ErbB signaling and subsequent inhibition of proliferation of cell lines expressing wild-type EGFR, selected EGFR exon 19 deletion mutations, or exon 21 L858R mutations. It also inhibited in vitro proliferation of cell lines overexpressing HER2. Overall, tumor growth was inhibited by Afatinib very effectively with low nanomolar IC50 values ranging from approximately 6nM to below 500 nM.

Ordering Information

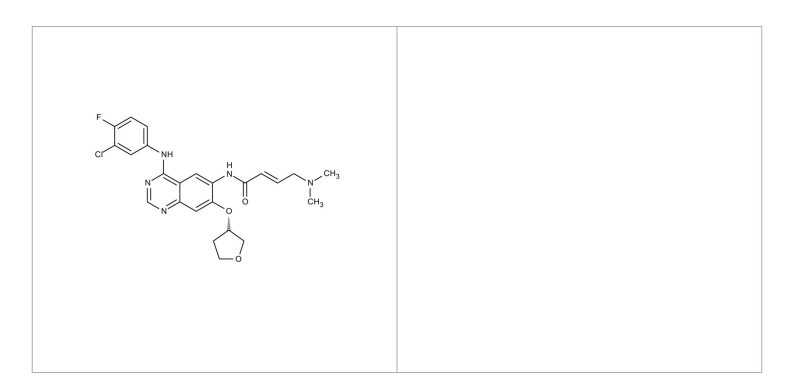
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ENZ-CHM158-0005

5mg

Manuals, SDS & CofA

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Handling & Storage

Use/Stability As indicated on product label or CoA when stored as recommended.

Short Term Storage +4°C

Long Term Storage -20°C

Shipping Ambient Temperature

Regulatory Status RUO - Research Use Only

Product Details

Alternative Name Tovok, BIBW-2992

Appearance Yellow powder.

CAS 850140-72-6

Couple Target EGFR, ErbB

Couple Type Inhibitor

Formula $C_{24}H_{25}CIFN_5O_3$

Identity Determined by EM-MS, NMR

MW 485.9

Purity ≥95% (HPLC)

Solubility Soluble in DMSO (97 mg/ml) or ethanol (15 mg/ml).

Last modified: May 29, 2024

