Covalent EGFR Inhibitor Analysis Reveals Importance of Reversible Interactions to Potency and Mechanisms of Drug Resistance

Phillip A. Schwartz, Petr Kuzmič, James Solowiej, Simon Bergqvist, Ben Bolanos, Chau Almaden, Asako Nagata, Kevin Ryan, Junli Feng, Deepak Dalvie, John Kath, MeiRong Xu, Revati Wani, Brion W. Murray

1. Substrate kinetics

This section describes the determination of kinetic properties of both the fluorescent peptide substrate “Y12”, and ATP as the co-substrate, in the Omnia assay. We first demonstrate by theoretical analysis that the specificity number \( k_{\text{sub}} = \frac{k_{\text{cat}}}{K_m} \) is well defined by the experimental data collected at low peptide substrate concentration \([S]_0 << K_m\) provided that the enzyme’s active-site concentration is known independently. We also derive a reduced “hit-and-run” mechanistic model \((E+S \rightarrow E+P)\) for substrate catalysis, which can be used to auto-generate a suitable system of first-order ordinary differential equations (ODEs) under these specialized experimental conditions. Finally, we present experimental data to show that the Michaelis constant for the peptide substrate, \(K_{m,\text{Pep}}\), cannot be determined because it is immeasurably high. However, we report a lower-limit estimate for \(K_{m,\text{Pep}}\) and demonstrate that it is indeed significantly higher than the maximum experimentally attainable peptide substrate concentration. These experimental findings justify our truncated theoretical model for the substrate catalysis branch of the overall inhibition mechanism.

### 1.1. Kinetic theory under the \([S]_0 << K_m\) conditions

**Algebraic analysis.** Under the experimental conditions where the substrate concentration remains very much lower than the Michaelis constant, \([S]_0 << K_m\), the classic Michaelis-Menten rate Eqn (1) reduces to Eqn (2). This is because if \([S]_0 << K_m\) then \([S]_0\) can be neglected in the summation term and therefore we obtain the approximate equality \([S]_0 + K_m \approx K_m\).

\[
\nu = \frac{[E]_0 k_{\text{cat}} [S]_0}{[S]_0 + K_m} \quad (1)
\]

\[
\nu \approx \frac{[E]_0 k_{\text{cat}} [S]_0}{K_m} \quad (2)
\]
The reaction rate is defined as the rate of change in substrate concentration, taken with negative sign, because the substrate is being consumed in the enzyme-catalyzed reaction, \( \nu \equiv -d[S]/dt \). The resulting differential equation (3) can be integrated analytically after separation of variables, to obtain the integral equation (4) as the theoretical model for reaction progress.

\[
\frac{d[S]}{dt} = -[E]_0 k_{sub} [S] \quad (3)
\]

\[
[S] = [S]_0 \exp(-[E]_0 k_{sub} t) \quad (4)
\]

\[
k_{sub} \equiv \frac{k_{cat}}{K_m} \quad (5)
\]

Thus, at substrate concentrations significantly lower than the Michaelis constant, the reaction progress curve conforms to the simple exponential model, and the apparent first-order rate constant is equal to \([E]_0 k_{cat}/K_m\). The enzyme concentration \([E]_0\) can be determined experimentally by active-site titration. Therefore, it is possible to determine the specificity number \(k_{sub} = k_{cat}/K_m\) simply by fitting the reaction progress to the first-order exponential equation and dividing the apparent first-order rate constant by \([E]_0\).

In conclusion, enzyme kinetic data observed under first-order conditions \(([S]_0 << K_m)\) do contain sufficient information (in the information-theoretic sense [1]) about \(k_{sub}\) as a potential fitting parameter.

**Differential Equation Modeling.** A number of software packages [2–4] allow the biochemist to specify the kinetic mechanism of an enzyme reaction using a symbolic notation, such as \(E + S \rightleftharpoons ES \rightarrow E + P\). The software then internally derives the corresponding mathematical model for the reaction progress, as a system of first-order ordinary differential equations (ODEs). The question arises what particular symbolic notation should be used to define the time-course of an enzyme reaction following the Michaelis-Menten kinetic mechanism under the first-order experimental conditions \(([S]_0 << K_m)\).

We do require that the reaction rate is strictly proportional to the enzyme concentration. We also require that the integrated kinetic equation describes a first-order exponential. This leads to the differential equation (3), which corresponds to the symbolic term \(E + S \rightarrow \cdots\). The remaining question is what should appear on the right-hand side. The answer lies in the requirement that the enzyme concentration should be treated as a constant in Michaelis-Menten kinetics. Therefore the free enzyme must appear on the right-hand side, because only then the rate terms for the consumption and generation of \(E\) cancel out, \(E + S \rightarrow E + P\), resulting in zero overall change in enzyme concentration over time. Thus we arrive at the “hit-and-run” symbolic model for enzyme catalysis, shown in Scheme S1.

In DynaFit notation [2], the “hit-and-run” model for substrate catalysis is represented as “\(E + S \rightarrow E + P : k_{sub}\)”. Scheme S1 is conceptually similar to the Theorell-Chance mechanism for bisubstrate enzymes [5, p. 594] \((E + A \rightarrow EA; EA + B \rightarrow EQ + P; EQ \rightarrow E + Q)\). No ternary molecular complex is postulated, even though at least one such complex is implied. In the “hit-and-run” mechanism, we also know that the Michaelis complex ES must be physically present. We are choosing to ignore it, so that we can produce a practically useful minimal model of the reaction progress.

**1.2 Substrate kinetics of the “Y12” peptide**

The “Y12” peptide (Ac-EEEEY(cSx)IV-NH\(_2\), Omnia Y Peptide 12, Invitrogen / Life Technologies) was assayed at ATP concentrations subsequently used in all inhibition assays ([ATP] = 800 \(\mu\)M) and at the enzyme concentration set to \([E]_0 = 50 \text{ nm}\). The peptide concentration \([S]_0\) was varied from 4 \(\mu\)M to 12 \(\mu\)M stepping by 2 \(\mu\)M, in triplicate. Fig. S1 shows a representative set of the reaction progress curves. Reaction progress curves are nonlinear throughout the entire assay and conform to the exponential fitting model, suggesting that the Michaelis constant is very much larger than the maximum substrate concentration.

![Scheme S1](image)

**Fig. S1:** Substrate kinetics of the “Y12” peptide vs. EGFR-L858R/T790M double mutant. Legend: peptide concentrations used in each kinetic experiment.

All reaction progress curves were nonlinear throughout the entire time course of the assay. The progress curves conformed ideally to the exponential fitting equation (6), where \(F\) is the fluorescence intensity recorded at the reaction time \(t\); \(F_0\) is the baseline fluorescence (an instrument offset); \(A\) is the exponential amplitude; and \(k\) is the empirical first order rate constant. Initial reaction rates were computed from the best-fit values of...
A and $k$ by using Eqn (7). The initial rates were subsequently fit to the Michaelis-Menten rate equation (8), in an attempt to determine the Michaelis constant $K_m$.

\[
F = F_0 + A \left[ 1 - \exp(-k \cdot t) \right] \quad (6)
\]

\[
v = A k \quad (7)
\]

\[
v = \frac{V_{\text{max}} \cdot [S]_0}{[S]_0 + K_m} \quad (8)
\]

![Image of substrate kinetics graph](image)

Fig. S 2: Substrate kinetics of the “Y12” peptide vs. EGFR-L858R/T790M double mutant: Circles - initial reaction rates computed from the best-fit values of exponential amplitude $A$ and exponential rate constants $k$, using Eqn (7). Solid curve - least square model curve corresponding to Eqn (8) while keeping the Michaelis constant at the lower limit of its 95% confidence interval, $K_m = 174 \mu M$. Dashed curves - inference (prediction) bands at the 95% confidence level.

Fig. S2 shows the results of the least-squares fit of the initial reaction rates, computed from Eqn (7), to the Michaelis-Menten rate Eqn (8). The best least-squares fit values and the corresponding standard errors of model parameters $V_{\text{max}}$ and $K_m$ could not be determined. In fact, the least-squares model curve was essentially a perfect straight line through the origin, corresponding to infinite values of both $V_{\text{max}}$ and $K_m$.

However, we were able to estimate at least to lower limit of the confidence interval for $K_m$ utilizing the profile-t (likelihood profile) method of Bates & Watts [6, 7]. Using this method we determined, at the 95% confidence level, that the apparent $K_m$ value of the peptide substrate, at [ATP] = 800 μM, must be larger than 174 μM. This value is very much larger than the corresponding Michaelis constant at the lower limit of its 95% confidence interval, $K_m << K_m$.

Thus, we conclude that in all enzyme assays we employed in this study to characterize the irreversible inhibitors of EGFR, the peptide substrate concentration [S] = 13 μM is (effectively) negligibly small when compared to the corresponding Michaelis constant. This justifies the use of the truncated (“hit-and-run”) kinetic model described in Scheme S1, which gives rise to first-order exponential kinetics in the absence of inhibitors, as defined by Eqn (9). In Eqn (9), [S] is the substrate concentration at the arbitrary reaction time $t$; $[S]_0$ is the initial substrate concentration at the $t = 0$; $[E]_0$ is the enzyme concentration; and $k_{\text{sub}}$ is the second-order rate constant by definition equal to $k_{\text{cat}}/K_m$.

\[
[S] = [S]_0 \exp(-k_{\text{sub}} [E]_0 t) \quad (9)
\]

It should be noted that under the special experimental conditions where the substrate concentration is effectively infinitely lower than the corresponding Michaelis constant ([S]_0 << K_m), essentially all enzyme exists in the unbound (free) state, and thus the biochemical system behaves as of the enzyme-substrate complex were not formed at all.

### 1.3. Substrate kinetics of ATP

ATP was assayed at the “Y12” peptide (Ac-EEEEY(cSx)IV-NH2, Omnia Y Peptide 12, Invitrogen / Life Technologies) concentrations subsequently used in all inhibition assays ([Pep] = 13 μM) and at the enzyme concentration set to [E]_0 = 30 nM. ATP concentration [ATP]_0 was varied according to the dilution series shown in the inset to Fig. S3. All assays were performed in triplicate, on separate micro-titer plates. Fig. S3 shows one of three replicated sets of the reaction progress curves.

![Image of ATP kinetics graph](image)

Fig. S 3: Substrate kinetics of ATP (Replicate 1 of 3) vs. EGFR-L858R/T790M double mutant: Reaction progress curves are nonlinear throughout the entire assay and conform to the exponential fitting model. Legend: ATP concentrations used in each kinetic experiment.

As was the case for peptide assays described above, all reaction progress curves were nonlinear throughout the entire time course of the assay. The progress curves conformed ideally to the exponential fitting equation (6). Initial reaction rates were computed from the best-fit values of $A$ and $k$ by using Eqn (7). The initial rates were subsequently fit to the Michaelis-Menten rate equation (8), in order to determine the Michaelis constant $K_m$. Fig. S4 shows the results of the least-squares fit of the
initial reaction rates. The best least-squares fit values and the corresponding standard errors of model parameters were $V_{\text{max}} = (37.7 \pm 1.5)$ RFU/sec and $K_{m, \text{ATP}} = (49.1 \pm 5.2) \mu M$.

2. Initial rate kinetics of EGFR inhibitors

In this section we describe the determination of apparent inhibition constants from initial reaction rates. The main purpose was to compare the results with the corresponding values of inhibition constants obtained by the global fit of complete reaction progress curves, and in so doing verify the inhibition constants obtained by either of the two independent method. The initial rates were determined by the least-squares fit on the “early” portion of each inhibition progress curve to an empirical model, represented by the first-order exponential. The reaction rates so obtained were subsequently fit to the Morrison Eqn [8] for tight-binding inhibition.

2.1. Exponential fit of “early” reaction progress

All EGFR inhibitors listed in Table 2 of the main manuscript contain the $\alpha,\beta$-unsaturated carboxamide moiety and therefore act as irreversible inhibitors of EGFR. Furthermore, the inhibitory effect for most inhibitors in this study is seen very prominently already at very low inhibitor concentrations comparable with the concentration of the enzyme (“tight binding”) [8]. For these reason the complete reaction progress curves ($t_{\text{max}} = 30 \text{ min}$) have a complex shape that can only be described by a numerical model represented by a system of differential equations [9]. However, we observed that the initial portion ($t_{\text{max}} = 7 \text{ min}$) of every progress curve at any given inhibitor concentration can be successfully fit to the single exponential equation (6). Fig. S5 shows the results or the exponential fit for neratinib. The initial reaction rates form EGFR assay at various inhibition concentrations were computed as the product of exponential amplitudes $A$ and the corresponding first-order rate constants $k$, as shown in Eqn (7).

$$K_i = \frac{K_i^*}{1 + [\text{ATP}]_0 / K_{m, \text{ATP}}} \quad (10)$$

In all inhibition assays we utilized $[\text{ATP}] = 800 \mu M$. Thus, the presumed “true” inhibition constants reported in this study are always $(1 + 800/50) = 17$ times lower than the apparent inhibition constants determined in data fitting.
the first reversible step (effectively instantaneous on the time-scale of these kinetic experiments) we observe the formation of the non-covalent complex. In the second, much slower and irreversible step, we subsequently observe the formation of the covalent conjugate.

2.2. Determination of apparent inhibition constants

![Graph showing initial rate kinetics of neratinib](image)

**Fig. S 6: Initial rate kinetics of neratinib:** Symbols - initial reaction rates computed from the best-fit values of exponential amplitude $A$ and exponential rate constants $k_i$ using Eqn (7). Each data point is an average from three independent determinations. Error bars - standard deviations from replicated ($n = 3$) initial rate measurements. Solid curve - least square model curve corresponding to the Morrison Eqn (11). Dashed curves - inference (prediction) bands at the 95% confidence level.

The initial reaction rates, determined by exponential fit of the initial portion of each inhibition progress curve, were fit to the Morrison Eqn (11) to assess the strength of enzyme/inhibitor binding in the initial non-covalent complex. In Eqn (11), $V_0$ is the uninhibited reaction rate observed in the absence of the inhibitor; $[E]$ is the enzyme concentration; $[I]$ is the inhibitor concentration; and $K_i^*$ is the apparent inhibition constant. The adjustable parameters were $V_0$ and $K_i^*$. For all ATP competitive inhibitors, the apparent inhibition constant relates to the “true” inhibition constant as is shown in Eqn (12).


$$K_i^* = K_i \left(1 + \frac{[\text{ATP}]}{K_{m,\text{ATP}}}ight)$$

(11)  
(12)

The results of fit are shown in Fig. S6 for neratinib as a representative example. The best-fit value of the apparent inhibition constant was $K_i^* = (4.0 \pm 0.5)$ nM. This result corresponds to the “true” inhibition constant $K_i = (0.24 \pm 0.03)$ nM, according to Eqn (10). It is important to note that the concentration of the peptide substrate was shown to be very much lower than the corresponding Michaelis constant $K_{m,\text{Pep}}$. Thus, the “true” inhibition constants computed by using Eqn (10) above do represent the thermodynamic dissociation equilibrium constant of the instantaneously formed non-covalent complex.

The apparent inhibition constant determined for neratinib by the initial rate method, $K_i^* = 4.0$ nM, agrees very well with the dissociation equilibrium constant computed from the best-fit values of the association and dissociation rate constant (see Results in the main article), defined as $K_i^* = k_{\text{off}} / k_{\text{on}}$. In three independent kinetic determinations, the $k_{\text{off}} / k_{\text{on}}$ ratio was found to be 3.4 nM, 3.6 nM, and 3.3 nM, respectively. Within the bounds of the formal standard errors, these values agree very well with $K_i^* = 4.0$ nM determined here from initial rates.

The inhibition constants determined from initial rates, for all irreversible inhibitors of EGFR listed in Table 2 of the main manuscript, are summarized in Table S1.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$K_i^*_{nm}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI-1033</td>
<td>1.6 ± 0.7</td>
</tr>
<tr>
<td>dacomitinib</td>
<td>18 ± 1</td>
</tr>
<tr>
<td>afatinib</td>
<td>4.4 ± 0.4</td>
</tr>
<tr>
<td>neratinib</td>
<td>4.0 ± 0.5</td>
</tr>
<tr>
<td>1</td>
<td>3.2 ± 0.5</td>
</tr>
<tr>
<td>CL-387785</td>
<td>180 ± 11</td>
</tr>
<tr>
<td>2</td>
<td>63 ± 5</td>
</tr>
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<td>3</td>
<td>84 ± 3</td>
</tr>
<tr>
<td>4</td>
<td>2200 ± 100</td>
</tr>
<tr>
<td>5</td>
<td>430 ± 30</td>
</tr>
<tr>
<td>WZ-4002</td>
<td>340 ± 20</td>
</tr>
</tbody>
</table>

Table S 1: Apparent inhibition constants determined from the fit of initial reaction rates to the Morrison Eqn (11) and subsequent conversion of $K_i$ to $K_i^*$ using Eqn (10). See Fig. S6 for a representative example.

3. Global analysis of full reaction progress curves

3.1. Rapid-equilibrium two-step mechanism

In the immediately preceding section, based on the analysis of initial reaction rates, we have established that EGFR is inhibited by $\alpha,\beta$-unsaturated compounds listed in Table 2 of the main manuscript in two clearly distinct steps. The first step, essentially instantaneous under our particular experimental conditions, is the reversible formation of the non-covalent enzyme-inhibitor complex (E-I in Scheme S2). The second step is the gradual, irreversible formation of the covalent complex (E-I in Scheme S2). The dashed-line box shown in Scheme S2 represents that the formation of the initial non-covalent complex is essentially instantaneous on the time scale of our kinetic experiments.

![Scheme S2](image)
3.2. Global fit to an ODE model

DynaFit input script file. The DynaFit [2] script shown in Fig. S7 was used to analyze the time course of EGFR double mutant being inhibited by neratinib. Essentially identical scripts were utilized for all 11 compounds listed in Table 2 of the main manuscript. Each of the 11 compounds was analyzed in three independent experiments (three separate plate-reader plates in the Omnia assay format). All 33 DynaFit scripts and the corresponding data files from 33 independent experiments are available upon request.

The notation “algorithm = differential-evolution” signifies that the global [10] least-square fit of combined progress curves was performed by using the Differential Evolution (DE) algorithm [11]. Briefly, the DE algorithm is an Evolutionary Strategy scheme starting from a population of initial estimates spanning a sufficiently wide range of allowable values. In this case, DynaFit (under its default settings) randomly generated 300 initial parameter estimates spanning 12 orders of magnitude for each of the four rate constants listed in the “[constants]” section of the script. More specifically, the numerical value of each individual rate constant was allowed to vary from $10^{-6}$ to $10^6$. The “[mechanism]” section is a DynaFit representation of the reaction mechanism displayed in Schemes S1 and S2, combined.

All concentrations in the listing shown in Fig. S7 are in micromolar units. The notation “$E = 0.02 ? (0.005 .. 0.035)$” signifies that the nominal enzyme concentration ([E] = 20 nM) should be treated as one of the adjustable model parameters spanning from 5 to 35 nM. The notation “conc I = 0.03906 ?’ (and similarly for other data sets in the global super-set of experimental data shown in Listing S1.1) signifies that all nonzero inhibitor concentrations should also be treated as adjustable model parameter. The notation “Constraints Concentrations = 0.2” signifies that the inhibitor concentrations are allowed to vary by at most 20% of their respective nominal value (in this example [I] = 39.06 nM). The notation “P = 5000 ? (1000 .. 10000)” signifies that the difference molar response coefficient (i.e., the change in fluorescence intensity in Relative Fluorescence Units per micromolar amount of product formed) should be constrained within one order of magnitude. These particular constraints were set on the basis of preliminary experiments. The notation “offset = auto ? (-2000 .. +1000)” signifies that the offset on the fluorescence intensity axis should be optimized, separately for each recorded progress curves, within the indicated bounds.

In accordance with the postulated mechanism, the software internally generated as the least-squares fitting model the system of Ordinary Differential Equations (ODE system) (8) - (13).

This ODE system was integrated numerically using the LSODE algorithm [12].

\[
\begin{align*}
\frac{d[E]}{dt} &= -k_{on}[E][I] + k_{off}[E.I] \quad (13) \\
\frac{d[S]}{dt} &= -k_{cat}[E][S] \quad (14) \\
\frac{d[P]}{dt} &= +k_{cat}[E][S] \quad (15) \\
\frac{d[I]}{dt} &= -k_{on}[E][I] + k_{off}[E.I] \quad (16) \\
\frac{d[E.I]}{dt} &= +k_{cat}[E][I] - k_{cat}[E.I] - k_{inact}[E.I] \quad (17) \\
\frac{d[E-I]}{dt} &= +k_{inact}[E.I] \quad (18)
\end{align*}
\]

Results of global fit. The results of fit for three independent kinetic experiments with neratinib are shown in Table S2. The overlay of the best-fit model on the experimental data for one of the three replicates is shown in graphical form in Fig. S8.

A fundamentally important question arises regarding the confidence intervals for microscopic rate constants appearing in the combined Schemes S1 and S2. Namely, the question is which of the four microscopic rate constants (if any) can be reliably determined from our particular type experimental data. The answer is unambiguously obtained upon examining the replicated values of $k_{sub}$, $k_{on}$, $k_{off}$, and $k_{inact}$ listed in Table S2.
In this context it is very important to note that the “plus or minus” values listed in the right-most column are not the formal standard errors arising from nonlinear regression of any particular global data set. Instead the averages and standard deviations for microscopic rate constants listed in Table S2 arose in three entirely independent kinetic experiments, each “experiment” consisting of six to nine reaction progress curves obtained at various inhibitor concentrations pooled together and subjected to global [10] fit.

Thus, the rate constant values in the right-most two columns of Table S2 are averages and standard deviations from the three independent replicates. The controversial [3, 13, 14] formal standard errors of rate constants from nonlinear regression [15, p. 815, Eqn 5.6.4] were entirely ignored in this report.

To assess which particular microscopic rate constant is sufficiently well defined by our experimental data, we reasoned that any particular microscopic rate constant that is well defined will also be well reproduced upon going from one independent set of kinetic experiments to the next replicated set. The conclusions are as follows.

The microscopic rate constant $k_{\text{sub}}$ (i.e., the specificity number, $k_{\text{sub}} = k_{\text{cat}}/K_m$) is very well reproduced across independent replicates, with the coefficient of variation CV approximately equal to 20%. The inactivation rate constant, $k_{\text{inact}}$, is also very well reproduced across all independent replicates (CV $\approx$ 20%, $n = 3$).

In contrast, the bimolecular association rate constant $k_{\text{on}}^*$ is not defined by the kinetic data shown in Fig. 8. The dissociation rate constant $k_{\text{off}}$ (i.e., the “off rate” constant $k_{\text{off}}$) is not defined by the data, either. However, the rate constant ratio $k_{\text{off}}/k_{\text{on}}^*$ (i.e., the apparent inhibition constant $K_i^*$) is very well reproduced across all three replicates (CV $\approx$ 20%, $n = 3$).

In other words, to characterize any given irreversible inhibitor on the basis of our particular type of experimental data, we can only determine $k_{\text{inact}}$ and $K_i^* = k_{\text{off}}/k_{\text{on}}^*$. We cannot determine the individual values of $k_{\text{off}}$ or $k_{\text{on}}^*$. Thus, by implication, neither can we determine at all the value of $K_i^* = (k_{\text{off}} + k_{\text{inact}})/k_{\text{on}}^*$. As will be shown in a forthcoming report, only the lower limit for both $k_{\text{on}}^*$ and $k_{\text{off}}$ can be determined under rapid-equilibrium experimental conditions, such as those utilized in our study.

The results shown in Table S2 demonstrate that the association rate constant $k_{\text{on}}^*$ is not defined by our type of transient kinetic data. The nominal average value of $k_{\text{on}}^*$ is $3.7 \times 10^{10}$ M$^{-1}$s$^{-1}$, which would appear to violate the diffusion limit of approximately $10^9$ M$^{-1}$s$^{-1}$. However, the nominal standard error, $\pm 4.6 \times 10^{10}$ M$^{-1}$s$^{-1}$, is greater than the average value itself. Under such circumstances the “best-fit” value of any rate constant is meaningless, which why we identified $k_{\text{on}}^*$ as undefined.

A detailed analysis revealed that any value of $k_{\text{on}}^*$ that is greater than $100 \mu M^{-1}s^{-1}$ will fit all our data sets equally well, for all compounds. Therefore, in the next round of kinetic anal-

<table>
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<th>Parameter</th>
<th>Low</th>
<th>High</th>
<th>Replicate 1</th>
<th>Replicate 2</th>
<th>Replicate 3</th>
<th>Mean ± Std.Dev.</th>
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<td>$10^6$</td>
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</tr>
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<td>offset#3, RFU</td>
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<td>1000</td>
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<td>1</td>
<td>-296</td>
<td>-126 ± 153</td>
</tr>
<tr>
<td>offset#4, RFU</td>
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<td>1000</td>
<td>-298</td>
<td>-390</td>
<td>-405</td>
<td>-364 ± 58</td>
</tr>
<tr>
<td>offset#5, RFU</td>
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<td>1000</td>
<td>-145</td>
<td>-177</td>
<td>-361</td>
<td>-228 ± 116</td>
</tr>
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<td>offset#6, RFU</td>
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<td>1000</td>
<td>-460</td>
<td>-387</td>
<td>-606</td>
<td>-485 ± 112</td>
</tr>
<tr>
<td>offset#7, RFU</td>
<td>-2000</td>
<td>1000</td>
<td>-454</td>
<td>-632</td>
<td>-608</td>
<td>-565 ± 97</td>
</tr>
<tr>
<td>offset#8, RFU</td>
<td>-2000</td>
<td>1000</td>
<td>-558</td>
<td>-617</td>
<td>-407</td>
<td>-528 ± 108</td>
</tr>
</tbody>
</table>

Table S2: Results of Differential Evolution (DE) global least-squares fit of three independent kinetic experiments with neratinib vs. EGFR-L858R/T790M. The “Low” / “High” columns show the constraints imposed on each given parameters. Each replicate measurement consisted of eight progress curves obtained at the nominal enzyme concentration $[E] = 20$ nM. The nominal values of seven nonzero inhibitor concentrations are the geometric means of the “Low” and “High” values (see also inset in Fig. S8). The eighth kinetic trace was the control experiment with [I] = 0. The value of $K_i^*$ was computed from the best fit values of microscopic rate constants as $k_{\text{off}}/k_{\text{on}}^*$. 
ysis, the rate constant $k_{inact}^*$ was held fixed at 100 $\mu$m-1s-1 and only the rate constants $k_{sub}$, $k_{off}$ and $k_{inact}$ were optimized in the regression. For very “tight binding” [8] compounds, but not for others, the nominal enzyme concentration was also optimized in the least squares regression.

Table S3 summarizes the results. As was the case in values listed in Table S2, all standard errors shown in Table S3 are the standard deviations from replicated determinations ($n = 3$), not formal standard errors from nonlinear regression. To compute the $K_i$ values listed in Table 2 of the main manuscript, we have first computed the apparent $K_i^*$ value as the ratio of microscopic rate constants $K_i^* = k_{off}/k_{sub}$. Subsequently, assuming that all inhibitors are strictly competitive with ATP, the “true” $K_i$ values were computed using Eqn (10).

The standard deviations from replicates listed in Table S3 are sufficiently small (coefficient of variation lower than 20%) for all three adjustable rate constants, $k_{sub}$, $k_{off}$, and $k_{inact}$.

Strictly speaking, one would expect that the $k_{sub}$ values would be identical across experiments conducted with various inhibitors. The minor variations we do see are probably due to the fact that the enzyme concentration was not exactly identical in each series of experiments ($[E]_0$ and $k_{sub}$ are closely correlated in the regression model).

These results convincingly prove that all three microscopic rate constants treated as adjustable parameters in our differential equations, including $k_{sub}$, are well determined by our experimental data.

**Comparison of $K_i^*$ values from different methods.** Fig. S9 shows in graphical form the comparison between the apparent $K_i^*$ values computed by two fully independent methods: (i) from the initial reaction rates obtained by the exponential fit; and (ii) from the global Differential Evolution fit of complete reaction progress curves. The results show that the two methods are in excellent agreement (coefficient of determination $R^2 > 0.99$). The slope of the regression line (0.85) suggests that the apparent $K_i^*$ values computed by progress curve analysis are systematically 15% lower than the apparent $K_i^*$ values obtained by the analysis of initial reaction rates.

*These results convincingly prove that, with our newly described data-analytic method, we can reliably determine sub-nanomolar $K_i$ values even though the enzyme concentration is higher than 10 nM.*

4. Biochemical vs. cellular potency correlations

In this section we present a detailed graphical comparison between biochemical potency, as measured by various kinetic parameters ($k_{inact}$, $K_i$, and $k_{inact}/K_i$)

The results displayed in Fig. S10 are drawn from the numerical values listed in Table 2 of the main manuscript. Cellular potency, as measured by inhibition of autophosphorylation in tumor cells, is influenced both by the chemical reactivity of the “warhead” moiety, as measured by $k_{inact}$, and the dissociation equilibrium constant of the initially formed non-covalent enzyme/inhibitor complex, as measured by $K_i^*$ or $K_i$. This is evidenced by the fact that both $k_{inact}$ and $K_i^*$ show significant correlation with the cellular IC$_{S0}$.

---

**[task]**

data = progress | task = fit | algorithm = differential-evolution

**[mechanism]**

E + S $\rightarrow$ E + P : ksub
E + I $\leftrightarrow$ E.I : kon* koff
E.I $\rightarrow$ E-I : kinact

**[constants]**

| $k_{sub} = 1$ | $kon* = 1$ | $koff = 1$ | $kinact = 1$ |

**[concentrations]**

| $E = 0.02$ | (0.005 .. 0.035) | $S = 13$ |

**[responses]**

| $P = 5000$ | (1000 .. 10000) |

**[data]**

directory ./project/kinase/EGFR/inhib/neratinib/Replicate-1/data

sheet sheet.txt

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column 8 | offset = auto | (-2000 .. +1000) | conc I = 0.01465 |
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column 13 | offset = auto | (-2000 .. +1000) | conc I = 0 |

**[settings]**

{Constraints} | Concentrations = 0.2

---

Fig. S 7: DynaFit [2] script used to analyze the time course of EGFR double mutant being inhibited by neratinib.
However, the influence of the initial, non-covalent binding interactions is far stronger than the chemical reactivity of the “warhead”. This is made evident by the fact that the coefficient of determination for log(\(k_{\text{inact}}\)) vs. log(IC\(_{50}\)) is merely \(R^2 = 0.60\), suggesting only weak to moderate correlation. In contrast, the correlation coefficient for log(\(K_i\)) is 0.89, suggesting strong correlation.

Interestingly, the strongest correlation (\(R^2 = 0.95\)) between the biochemical and cellular potency is seen for the ratio \(K_i/k_{\text{inact}}\) vs. log(IC\(_{50}\)). In fact if the eleven inhibitors in Table 1 were ordered in terms decreasing \(K_i/k_{\text{inact}}\), the resulting ordering would correctly predict the cellular potency (i.e., decreasing cellular IC\(_{50}\)) for ten out of eleven inhibitors listed in Table 1. Only compound 3 would appear moderately out of order.

In this context it should be noted that the inverse ratio, \(k_{\text{inact}}/K_i\), can be viewed as the lower limit estimate on the microscopic bimolecular association rate constant \(k_{\text{on}}\). The microscopic rate constant \(k_{\text{on}}\) characterizes the “coming together” of the enzyme and the inhibitor, to form the initial non-covalent complex.

In addition to the 11 compounds listed in Table 2 of the main manuscript, we analyzed the correlation between biochemical potency (\(K_i\) against EGFR-L858R/T790M double mutant) and cellular potency (autophosphorylation in H1975 tumor cells) for 154 cell-permeable, covalent inhibitors spanning six inhibitor scaffolds, see Fig. S11. The coefficient of determination corresponding to the plot in Fig. S11 is \(R^2 = 0.72\), which again suggests that initial non-covalent binding contributes very importantly to overall cellular potency.

5. Kinetics of different oxidation states

Table S4 lists the apparent \(k_{\text{cat}}\) and \(K_m\) values of different oxidation states of EGFR mutants. All kinetic parameters listed in Table S3 were obtained from global fit of reaction progress curves (EGFR-L858R/T790M) to the system of differential equations (13)–(18). The plus-or-minus values are standard deviations from averaging three replicated, entirely independent experiments. For further details see text.

<table>
<thead>
<tr>
<th>compound</th>
<th>(k_{\text{cat}}, \mu M^{-1} s^{-1})</th>
<th>(k_{\text{off}}, s^{-1})</th>
<th>(k_{\text{inact}}, s^{-1})</th>
<th>(K_i^*, \text{NM})</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI-1033</td>
<td>0.028 ± 0.005</td>
<td>0.19 ± 0.04</td>
<td>0.011 ± 0.0002</td>
<td>1.9 ± 0.4</td>
</tr>
<tr>
<td>dacomitinib</td>
<td>0.023 ± 0.002</td>
<td>1.1 ± 0.1</td>
<td>0.0018 ± 0.0001</td>
<td>10.7 ± 0.9</td>
</tr>
<tr>
<td>afatinib</td>
<td>0.017 ± 0.004</td>
<td>0.3 ± 0.1</td>
<td>0.0024 ± 0.0003</td>
<td>2.8 ± 0.6</td>
</tr>
<tr>
<td>neratinib</td>
<td>0.016 ± 0.002</td>
<td>0.2 ± 0.1</td>
<td>0.0011 ± 0.0002</td>
<td>2.4 ± 0.5</td>
</tr>
<tr>
<td>1</td>
<td>0.023 ± 0.004</td>
<td>0.2 ± 0.1</td>
<td>0.008 ± 0.004</td>
<td>2.4 ± 1.2</td>
</tr>
<tr>
<td>CL-387785</td>
<td>0.017 ± 0.001</td>
<td>18 ± 4</td>
<td>0.0020 ± 0.0003</td>
<td>180 ± 40</td>
</tr>
<tr>
<td>2</td>
<td>0.014 ± 0.004</td>
<td>4 ± 0.5</td>
<td>0.0035 ± 0.0006</td>
<td>40 ± 5</td>
</tr>
<tr>
<td>3</td>
<td>0.026 ± 0.003</td>
<td>7 ± 2</td>
<td>0.0018 ± 0.0001</td>
<td>70 ± 20</td>
</tr>
<tr>
<td>4</td>
<td>0.021 ± 0.001</td>
<td>180 ± 30</td>
<td>0.00015 ± 0.00002</td>
<td>1800 ± 300</td>
</tr>
<tr>
<td>5</td>
<td>0.020 ± 0.004</td>
<td>50 ± 5</td>
<td>0.0011 ± 0.0001</td>
<td>500 ± 40</td>
</tr>
<tr>
<td>WZ-4002</td>
<td>0.024 ± 0.001</td>
<td>23 ± 5</td>
<td>0.0049 ± 0.0015</td>
<td>230 ± 50</td>
</tr>
</tbody>
</table>

Table S3: Results of global fit reaction progress curves (EGFR-L858R/T790M) to the system of differential equations (13)–(18). The plus-or-minus values are standard deviations from averaging three replicated, entirely independent experiments. For further details see text.
proteins were immobilized on a CM5 sensor chip by standard
5% glycerol, 1% DMSO, 0.5 mM TCEP, 0.005% P20. EGFR
versible or reversible inhibition enzyme kinetics.

The peptide substrate in these determinations was al-
substrate.

6. Inhibitor Pharmacology: Oxidized EGFR-Cys797

Table S5 shows the potency values for inhibitors of oxidized
EGFR-L858R single mutant in biochemical and cellular assays.
Cellular potency was quantified by determining the inhibition
of EGFR autophosphorylation in H3255 tumor cells. Table
S6 shows similar results for the EGFR-L858R/T790M double
mutant. In this case cellular potency is quantified as inhibi-
tion of EGFR autophosphorylation in H1975 tumor cells. The
results show that EGFR-Cys797 oxidation (-SH, unoxidized, -
SO2H sulfinylated, -SSG, glutathiolated) can profoundly affect
inhibitor affinity depending on the inhibitor characteristics and
EGFR context.

7. Surface plasmon resonance (SPR) binding studies

To evaluate the energetic contribution of the Cys797-Michael
Acceptor interaction to overall potency, reversible EGFR in-
hibitor affinities to EGFR were determined by surface plas-
mon resonance and compared to the inhibitor affinities (K_i) of
the corresponding covalent inhibitors determined by either irre-
versible or reversible inhibition enzyme kinetics.

SPR studies were carried out on a Biacore 3000 instrument
at 25°C in 150mM NaCl, 25mM HEPES, pH 7.2, 5mM MgCl2,
5% glycerol, 1% DMSO, 0.5 mM TCEP, 0.005% P20. EGFR
proteins were immobilized on a CM5 sensor chip by standard
amine coupling, in 10 mM sodium acetate pH 5.0 (1 M EGFR).
Injections were made using the Kinject mode at 50 µL/min with
a 200-1200s compound dissociation time. Compound injec-
tions were referenced to a blank surface and by a buffer blank.
Data analysis and fitting to a simple 1:1 kinetic model was per-
formed using the Scrubber2 software (BioLogic Software, Pty.,
Australia).

The results are summarized in Table S7 for wild-type enzyme
and in Table S8 for the L858R/T790M double mutant.

8. Mass spectrometric characterization of EGFR

8.1. Mass spectrometry methods

Intact mass analysis of modified EGFR protein was measured
by electrospray ionization on an Agilent (San Jose, CA) 6210
time-of-flight mass spectrometer (TOF-MS) coupled to an Ag-
ilent 1200 LC. EGFR protein samples, 500 ng/10 µL, were in-
jected onto an Acquity UPLC BEH300 C4 2.1 x 100 mm col-
umn (Waters, Bedford, MA.). At a flow rate of 0.5 mL/min, a
six-minute mobile phase gradient was constructed using 0.1%
formic acid (Solvent A) and acetonitrile (Solvent B). In the
first two minutes of the gradient, acetonitrile (Solvent B) was
ramped up to 40%, during which the column elution was sent to
waste. Subsequently, the LC stream was directed into the MS,
and solvent B was increased to 60% for the next three minutes
for elution of EGFR protein. In the final minute of the gradient,
the column was exposed to 90% acetonitrile and then allowed
re-equilibrated to 2% Solvent A. The protein was eluted off the
column into the MS-TOF, which was set to detect a mass range
from 600-2000 m/z. The MS fragmentor was set at 200 V, and
the skimmer at 140 V. Once eluted, the protein spectra were ex-
tacted from the eluted chromatography peaks in Agilent Qual-
itative Analysis software, and subsequently deconvoluted with
the Maximum Entropy algorithm. The deconvolution was run
over a mass range of 10-100 kDa, with a 1 Dalton mass step,
and 20 signal/noise threshold.

For identification of modified residues, “bottom-up” mass
spectrometry was employed, using nanoLC-MS. For “in-gel”
digestion, EGFR protein was first denatured in 8M urea (30
min, room temp) and then alkylated with 10 mM iodoacetamide
(2 hour, room temp) and then digested for 16 hours at 37°C.
Subsequently, 10 µL of sample (0.5 µg/µL) was loaded into wells on a non-reducing
SDS-PAGE gel, which was Coomassie blue stained. EGFR gel
bands were excised, cut and destained twice in 60/40 acetonit-
trile/water for 1 hour at room temp. For the subsequent prote-
olysis, the gel slices were first washed in 5% formic acid, be-
fore addition of 50 ng of pepsin (Protea, Morgantown, WV.
) and incubated for 16 hours at 37°C. The resultant EGFR di-
gest was analyzed on a LTQ mass spectrometer (Thermo Fisher
Scientific) equipped with a Michrom Captive Spray ionization
source, and coupled to a Proxeon nanoLC. Sample was injected at
5µL onto a Reprosil ProteCol Trap C18-AQ (SGE Analytical
Chemical) equipped with a Michrom Captive Spray ionization
source, and coupled to a Proxeon nanoLC. Sample was injected at
5µL onto a Reprosil ProteCol Trap C18-AQ (SGE Analytical
Chemical), for initial desalting with 20 µL of 0.1% formic acid
solution. Analytical separation was performed using a NanoES-
C18 column (Michrom BioResources, Auburn, CA.), 0.2x150
mm, running at a flow rate of 1.8 µL/min. The LC gradient con-
sisted of a 35-minute gradient from 2-40% acetonitrile with
0.1% (v/v) formic acid to elute peptides into the mass spectrometer. Mass spectrometry data was collected in “triple-play” mode, consisting of a full scan at 400-2000 m/z for selection of the three most predominant eluting peptides, followed by a high resolution ultra-zoom scan, and MS-MS of isolated peaks of interest. MS-MS spectra were collected in an isolation window of 3 m/z and a collision activation energy (CAD) of 35 eV. Data was then processed in Agilent Spectrum Mill rev. 4.0, for peptide identification. The EGFR mutation protein sequences were appended to the Spectrum Mill in-house database, and peptides were searched directly against the prescribed EGFR sequence. Also, peptides were searched with variable methionine oxidation, cysteine oxidation and cysteine glutathiolation. Precursor mass tolerance was set at 1.0 Da, and 0.7 Da for the fragment ion. Peptides were validated by fixed thresholds of forward-reverse scores < 8, with spectrum intensities (SPI) < 70. The search parameters were set for up to three missed cleavage sites.

8.2. Mass spectrometry results

Localization of EGFR oxidation was performed by LC-MS-MS analysis of the EGFR pepsin digest. Pepsin was the preferred protease because it provided greater than 89% sequence coverage for both the single activating mutation (L858R) and double mutant EGFR (T790M, L858R), and most importantly allowed for observation all six cysteine residues. LC-MS-MS analysis of the oxidized single and double mutant EGFR protein identified Cys797 as the site of oxidation. Shown in Fig. S12 is both the intact mass MS spectrum of the oxidized double mutant EGFR (T790M, L858R), and the subsequent MS-MS spectra for the pepsin generated peptides. Verification of Cys797 oxidation to its sulfinic acid form was verified from MS-MS fragmentation of the proteolytic peptide from residues 782-805, from which Spectrum Mill data analysis confidently assigned modification of Cys797 (Score = 18.40, Fwd Rev Score = 18.40). It should be noted that both the cysteine-sulfinic (SO₂⁻) and -sulfonic (SO₃⁻) acid forms were observed in MS-MS analysis, which is not surprising given there was indication of a minimal portion (~<10%) of SO₃ oxidation in the intact mass analysis. EGFR-Cys797 was identified as the only EGFR residue with an altered oxidation state for both the EGFR-L858R and EGFR-L858R/T790M H₂O₂ oxidized samples (Fig. S12). As with H₂O₂ oxidation, glutathiolation of EGFR-Cys797 was the only modification site for both observed for both EGFR-L858R and EGFR-L858R/T790M. Shown in Fig. S13 is the MS-MS spectrum for the glutathiolated peptide spanning residues 788-800 of double mutant of EGFR, from which Spectrum Mill data analysis confidently assigned modification of Cys797 (Score = 11.70, Fwd Rev Score = 11.70).

9. Molecular modeling

The active site orientation of the inhibitor Michael Acceptor to the reactive cysteine is defined by the inhibitor scaffold. Molecular modeling (Figure S14) predicts the binding mode of compound I (magenta) and WZ4002 (yellow) is very different for the EGFR L858R mutant. The distance is calculated between C(ligand)-S(Cys797) (yellow residue).

We postulate that sulfinylation of EGFR-Cys797 may enhance inhibitor affinity by creating a positive interaction between the

### Table S 4: Kinetic analysis of EGFR and oxidized-EGFR (-SH, unoxidized, -SO₂H sulfinylated, -SSG, glutathiolated) catalysis for L858R and L858R/T770M variants using the Ac-E-E-E-Y-Y-NH₂ peptide. The S-glutathiolation did not reduce the fraction of functional active sites for either protein but sulfinylation did by 50%. EGFR protein was highly pure but were normalized for the fraction of functional active sites to provide the most accurate kinetic constants.

<table>
<thead>
<tr>
<th>Cys797 oxidation</th>
<th>L858R</th>
<th>L858R/T770M</th>
<th>w.t.</th>
</tr>
</thead>
<tbody>
<tr>
<td>kₕcat, s⁻¹</td>
<td>6.8±0.2</td>
<td>7.6±0.2</td>
<td>6.9±0.4</td>
</tr>
<tr>
<td>Kₘ, µM</td>
<td>1120±30</td>
<td>280±20</td>
<td>260±10</td>
</tr>
<tr>
<td>kₕcat/Kₘ, µM⁻¹s⁻¹</td>
<td>0.0061±0.00002</td>
<td>0.027±0.0002</td>
<td>0.027±0.0002</td>
</tr>
</tbody>
</table>

### Table S 5: Potency values for inhibitors of oxidized EGFR-L858R single mutant in biochemical and cellular assays. ND = not determined.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Biochemical Kᵢₕ, nm</th>
<th>Cellular IC₅₀, nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI-1033</td>
<td>4.2 ± 0.6</td>
<td>4.7 ± 0.3</td>
</tr>
<tr>
<td>dacomitinib</td>
<td>21 ± 2</td>
<td>1.0 ± 0.1</td>
</tr>
<tr>
<td>afatinib</td>
<td>6.7 ± 0.9</td>
<td>7.0 ± 1.1</td>
</tr>
<tr>
<td>erlotinib</td>
<td>13 ± 1</td>
<td>1.7 ± 0.2</td>
</tr>
<tr>
<td>gefitinib</td>
<td>29 ± 3</td>
<td>1.4 ± 0.1</td>
</tr>
<tr>
<td>lapatinib</td>
<td>7.4 ± 0.8</td>
<td>5.0 ± 1</td>
</tr>
<tr>
<td>WZ4002</td>
<td>46 ± 4</td>
<td>1.2 ± 0.1</td>
</tr>
<tr>
<td>WZ4003</td>
<td>5100 ± 300</td>
<td>14900 ± 500</td>
</tr>
<tr>
<td>8</td>
<td>3700 ± 40</td>
<td>11000 ± 100</td>
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<td>7700 ± 300</td>
<td>86 ± 21</td>
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<td>8</td>
<td>5100 ± 300</td>
<td>&gt;10000</td>
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<tr>
<td>9</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td>8</td>
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</tr>
<tr>
<td>9</td>
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<td>ND</td>
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<td>8</td>
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</tr>
<tr>
<td>9</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>
oxidized cysteine residue S=O group and an inhibitor basic substituent. Molecular modeling studies (Fig. S15) predict a binding mode of gefitinib (magenta, PDB ID 2ITZ) and dacomitinib (yellow, docked) in the unoxidized form of EGFR-L858R with a short N(ligand)-S(Cys797) distance (4.58 - 5.23 Å). With sulfinylated EGFR-Cys797, the cysteine S=O and the inhibitor NH group are expected to be at a good interaction distance.

Histograms of hydrogen bond length distributions of NH(ligand) and S=O(Cys797) in Cambridge Structure Database (Fig. S16) suggest the mean N–O distance of 2.94 Å (Top Panel) and the mean N–S distance of 4.07 Å (Bottom Panel).

### 10. Detailed experimental methods

#### 10.1. Expression and purification of EGFR proteins

The cDNA encoding the core domain (residues 696-1022) of human EGFR double mutant (L858R/T790M) was synthesized and cloned into pKRIC-N3 using the BamH1 and HindIII sites. Similarly, the cDNA encoding the single mutant (L858R) core domain (residues 695-1022) was cloned into pKRIC-N5 and the cDNA encoding the WT core domain with the juxtamembrane region (residue 668-1022) was cloned into pKRIC-N3.

<table>
<thead>
<tr>
<th>Inhibitor</th>
<th>biochemical $K_i$, nm</th>
<th>cellular IC$_{50}$, nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI1033</td>
<td>16 ± 2 21 ± 2 0.2 ± 0.1</td>
<td>2.3 ± 0.5</td>
</tr>
<tr>
<td>dacomitinib</td>
<td>49 ± 6 2.4 ± 0.6 0.9 ± 0.1</td>
<td>10 ± 1</td>
</tr>
<tr>
<td>1</td>
<td>36 ± 1 55 ± 8 0.3 ± 0.1</td>
<td>6.6 ± 0.2</td>
</tr>
<tr>
<td>6</td>
<td>82 ± 2 2.2 ± 0.4 29 ± 2</td>
<td>7.3 ± 1.1</td>
</tr>
<tr>
<td>erlotinib</td>
<td>80 ± 7 47 ± 2 54 ± 1</td>
<td>5200 ± 300</td>
</tr>
<tr>
<td>gefitinib</td>
<td>420 ± 50 5.5 ± 0.3 22 ± 3</td>
<td>4300 ± 2300</td>
</tr>
<tr>
<td>lapatinib</td>
<td>210 ± 10 210 ± 20 120 ± 3</td>
<td>7400 ± 100</td>
</tr>
<tr>
<td>WZ4002</td>
<td>2200 ± 200 1400 ± 200 16 ± 4</td>
<td>78 ± 25</td>
</tr>
<tr>
<td>WZ4003</td>
<td>1700 ± 100 870 ± 10 340 ± 10</td>
<td>&gt;10,000</td>
</tr>
<tr>
<td>8</td>
<td>ND ND ND</td>
<td>≈660</td>
</tr>
<tr>
<td>7</td>
<td>ND ND ND</td>
<td>1800 ± 300</td>
</tr>
</tbody>
</table>

Table S6: Potency values for inhibitors of oxidized EGFR-L858R/T790M double mutant in biochemical and cellular assays. ND = not determined.

<table>
<thead>
<tr>
<th>unreactive analog</th>
<th>$k_{on}$, $\mu$m $^{-1}$s$^{-1}$</th>
<th>$k_{off}$, s$^{-1}$</th>
<th>$K_d$, nm</th>
<th>$K_i$, nm</th>
<th>reactive analog</th>
<th>$K_i$, nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>7</td>
<td>0.9</td>
<td>0.00046</td>
<td>0.51</td>
<td>0.8</td>
<td>CI-1033</td>
<td>0.093</td>
</tr>
<tr>
<td>8</td>
<td>0.84</td>
<td>0.0015</td>
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<td>ND</td>
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Table S7: Biacore results vs. biochemical inhibition potency: wild-type EGFR. ND = not determined. For further explanation see text.
Both the double mutant (DM, L858R/T790M) and WT constructs contain an N-terminal his tagged GST followed by a TEV (Tobacco Etch Virus) cleavage site and the single mutant (L858R) construct contains an N-terminal his tag followed by a TEV cleavage site. pKRiC-N3 and pKRiC-N5 are modified insect cell expression cloning vector pFastbac I. Therefore, they are only used in the Bac-to-Bac method for recombinant baculovirus production. Sf9 cells were used to generate recombinant baculovirus and to scale up expression for WT construct and sf21 cells were used to scale up expression for L858R/T790M and L858R constructs.

Specifically, sf9 or sf21 cells were infected with baculovirus based on cell density. Forty eight hours (for L858R and WT) or seventy two (for L858R/T790M) hours post infection, cells were harvested by centrifugation and the cell pellet was frozen and stored at -80°C. To purify the protein, a cell pellet was resuspended in a lysis buffer containing 100 mM HEPES, pH 8.0; 150 mM NaCl; 5 mM MgCl₂; 10% glycerol; 20 mM imidazole; 4 mM TCEP; and protease inhibitor tablets (Roche). Cells were lysed by stirring the suspension at 4°C for 45 min and the cell lysate was centrifuged at 10,000g-20,000g for 60 min at 4°C. The supernatant was collected by passing through four layers of cheesecloth. The tagged protein was then purified by batch binding with Invitrogen Probond resin for two hours at 4°C with slow rotation using a Roto-Shaker Genie, followed by washing the bound resin with the lysis buffer and then eluting with the lysis buffer containing 300 mM imidazole. The untagged and non-phosphorylated protein was obtained by overnight dialysis at 4°C in the presence of TEV protease and lambda phosphatase in a dialysis buffer containing 25 mM HEPES, pH 8.0; 500 mM NaCl; 5 mM MgCl₂; 10% Glycerol; 2 mM TCEP; and 1 mM MnCl₂, followed by nickel reverse chromatography. The untagged protein was further purified by Superdex75 size exclusion chromatography using an SEC buffer containing 25 mM HEPES, pH 8.0; 50-100 mM NaCl; 5 mM MgCl₂; 10% glycerol; 4 mM TCEP. The peak fractions were pooled based on the activity and purity (SDS-PAGE analysis). Pools were concentrated to a desired concentration (2-3 mg/mL).

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<th>$k_{off}$, s$^{-1}$</th>
<th>$K_d$, nm</th>
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Table S 8: Biacore results vs. biochemical inhibition potency: EGFR-L858R/T790M double mutant. ND = not determined. For further explanation see text.

Fig. S 12: Intact mass measure of the oxidized double mutant EGFR (L858R, T790M), indicating 90%, by peak area, in the SO₂-oxidized form. Subsequent pepsin digest of this oxidized protein, generated a peptide (residues 782-805) from which MS-MS identification of SO₂ on cysteine-797 is evident. Similar results were obtained for EGFR (L858R).
tion was measured using both Abs280 and colorimetric assay using BSA as standard (Biorad coomassie plus protein assay reagent). The protein purity was checked by SDS-PAGE with all proteins having >95% purity. The identity of protein was confirmed by intact mass spectroscopy. For WT protein purification, the lysis and nickel elution buffers had 200 mM NaCl and the dialysis and size exclusion chromatography buffers had 300 mM NaCl.

10.2. Determination of active enzyme concentration

Enzyme concentration for the non-oxidized EGFR mutants were determined by conducting the fluorometric covalent inhibitor assay in the presence of varying amounts of dacomitinib at concentrations ranging from one-tenth to two-fold of the nominal enzyme concentration. Assay conditions were similar to those used to test covalent inhibitors (see section Fluorometric enzyme assays), however EGFR (50 nm) and inhibitor were incubated for 20 minutes in the absence of ATP. The assay was initiated by the addition of 1.1 mM ATP.

The initial velocity was plotted against the ratio of [inhibitor]/[enzyme]. A straight line was fit through the data points which represented reactions that were not fully inhibited, and the intercept on the horizontal axis yielded the fraction of enzyme with active sites capable of turnover. L858R/T790M had 96% of the added protein capable of inhibition by this active conformation-binding inhibitor dacomitinib. L858R and WT had a smaller fraction of the active sites available to this inhibitor (53% and 23% respectively).

L858R and WT EGFR proteins were inhibited by the non-active conformation binding inhibitor lapatinib at varied concentrations. The initial rates were fit to the Morrison equation (11), where the active enzyme concentration $[E]_0$ was treated as one of the optimized parameters along with the apparent inhibition constant $K_i^*$. The best-fit value of $[E]_0$ was greater than 95% of the nominal enzyme concentration.

The same method was used to estimate the concentration of active sites for EGFR mutants oxidized at Cys797 (which are incapable of reactivity with covalent inhibitors). S-Glutathiolated EGFR-L858R/T790M has 100% of the active sites functional, while the sulfinylated form was 50% of the active sites functional. EGFR-L858R protein has 50% of the active sites functional after S-glutathiolation and 25% of active sites available after sulfinylation.

10.3. UV/Vis spectrophotometric enzyme assays

A spectrophotometric coupled enzymatic assay was used to monitor the kinase-catalyzed production of ADP from ATP that accompanies phosphoryl transfer to a phosphoacceptor peptide substrate Ac-EEEEYIIV-NH2. Activity was monitored by coupling NADH oxidation to the regeneration of ATP from product ADP through the action of pyruvate kinase (PK) and lactate
dehydrogenase (LDH). NADH conversion to NAD+ was monitored by the decrease in absorbance at 340 and 254\textsuperscript{deg}C.

Typical reaction solutions contained 2 mM phosphoenolpyruvate, 0.27 mM NADH, 12 mM free MgCl\(_2\), 1 mM DTT, 2 mM peptide (the solubility limit), 10 units/mL LDH, 150 mM NaCl, and 0.01% Tween-20 in 50 mM HEPES pH 7.5. ATP concentration was dependent upon the assay performed (Table S8.1). Reactions were performed either at 200 µL volumes in a quartz cuvette on a Beckman DU800 (kinetic parameter determination) or at 100 µL volumes in a clear-bottom, half-area 96-well plate on a Tecan Safire in absorbance mode (inhibitor affinity determination).

Kinetic parameters for the phosphoryl transfer reaction, the turnover number, \(k_{\text{cat},\text{ATP}}\), and the Michaelis constant for ATP, \(K_{\text{m,ATP}}\), respectively, were determined using the UV/Vis spectrophotometric coupled assay by varying [ATP] around that which produced half maximal velocity for each EGFR mutant (oxidized and non-oxidized). Reactions were initiated by the addition of 30 to 120 nm EGFR-L858R/T790M or EGFR-L858R (final concentration) for both the oxidized and non-oxidized species.

The initial portion of each reaction progress curve was fit to the straight-line model, to determine the initial rate as the slope. To determine the kinetic parameters \(k_{\text{cat},\text{ATP}}\) and \(K_{\text{m,ATP}}\), the initial rates so obtained were fit to Eqn (1), where \([E]_0\) is the enzyme concentration independently determined by active-site titration.

Inhibitor affinity (\(K_i\)) determinations were performed by fitting initial rate data to the Morrison Eqn (11). Apparent inhibition constants determined by nonlinear regression were converted to “true” inhibition constants by using Eqn (10), assuming strictly ATP-competitive inhibition mode. A liquid handling robot (BiomekFX) was used to make 1:2 inhibitor dilutions from DMSO stocks (2% final DMSO concentration), add inhibitor to the reaction mixture, and initiate the reaction by the addition of ATP substrate. Enzyme and ATP concentrations for each EGFR mutant (oxidized and non-oxidized) are found in Table S9.

The apparent \(K_{\text{m,ATP}}\) under fluorometric assay conditions for each non-oxidized EGFR mutant was determined by varying [ATP] around that which produced half maximal velocity. The Michaelis-Menten equation (14) was fitted to the data. Apparent \(K_{\text{m,ATP}}\) for EGFR-L858R, EGFR-L858R/T790M, EGFR-WT were determined to be 114, 50, and 34 M respectively.

Detailed kinetic analysis of covalent inhibitors acting on EGFR mutants not oxidized at EGFR-Cys\(_{797}\) was also performed by using the fluorometric assay. Determinations were made from a series of progress curves as a function of inhibitor concentration in DMSO (2% final DMSO concentration) to reactions containing 5 mM or 0.8 mM ATP for EGFR-L858R and EGFR-L858R/T790M assays, respectively.

11. Structures of nonreactive inhibitors

Figure 17 displays the chemical structures of nonreactive EGFR inhibitors utilized in this study. Corresponding reactive analogues are shown in parentheses.
References


