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Predicting resistance of clinical Abl mutations to targeted kinase inhibitors using alchemical free-energy calculations

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Abstract The therapeutic effect of targeted kinase inhibitors can be significantly reduced by intrinsic or 13 acquired resistance mutations that modulate the affinity of the drug for the kinase. In cancer, the majority of 14 missense mutations are rare, making it difficult to predict their impact on inhibitor affinity. This complicates 15 the practice of precision medicine, pairing of patients with clinical trials, and development of next-generation 16 inhibitors. Here, we examine the potential for alchemical free-energy calculations to predict how kinase 17 mutations modulate inhibitor affinities to Abl, a major target in chronic myelogenous leukemia (CML). We 18 find these calculations can achieve useful accuracy in predicting resistance for a set of eight FDA-approved 19 kinase inhibitors across 144 clinically-identified point mutations, achieving a root mean square error in 20 binding free energy changes of 1.1^{1.3}/_{0.9} kcal/mol (95% confidence interval) and correctly classifying mutations 21 as resistant or susceptible with 88_{82}^{93} % accuracy. Since these calculations are fast on modern GPUs, this 22 benchmark establishes the potential for physical modeling to collaboratively support the rapid assessment 23 and anticipation of the potential for patient mutations to affect drug potency in clinical applications. 24

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Targeted kinase inhibitors are a major therapeutic class in the treatment of cancer. A total of 38 selective 26 small molecule kinase inhibitors have now been approved by the FDA [1], including 34 approved to treat 27 cancer, and perhaps 50% of all current drugs in development target kinases [2]. Despite the success of 28 selective inhibitors, the emergence of drug resistance remains a challenge in the treatment of cancer [3-10] 29 and has motivated the development of second- and then third-generation inhibitors aimed at overcoming 30 recurrent resistance mutations [11-15]. 31 While a number of drug resistance mechanisms have been identified in cancer (e.g., induction of splice 32 variants [16], or alleviation of feedback [17]), inherent or acquired missense mutations in the kinase domain 33 of the target of therapy are a major form of resistance to tyrosine kinase inhibitors (TKI) [10, 18, 19]. Oncology 34 is entering a new era with major cancer centers now deep sequencing tumors to reveal genetic alterations 35 that may render subclonal populations susceptible or resistant to targeted inhibitors [20], but the use of 36 this information in precision medicine has lagged behind. It would be of enormous value in clinical practice 37 if an oncologist could reliably ascertain whether these mutations render the target of therapy resistant or 38 susceptible to available inhibitors; such tools would facilitate the enrollment of patients in mechanism-based 39

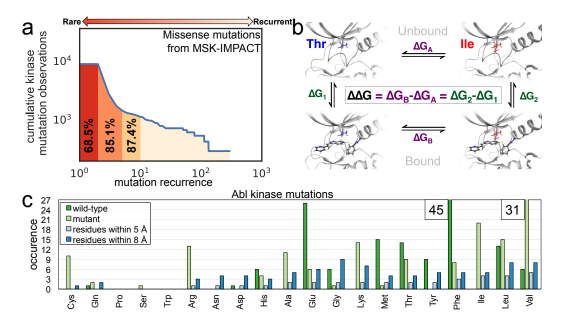


Figure 1. Relative alchemical free-energy calculations can be used to predict affinity changes of FDA-approved selective kinase inhibitors arising from clinically-identified mutations in their targets of therapy. (*a*) Missense mutation statistics derived from 10,336 patient samples subjected to MSK-IMPACT deep sequencing panel [20] show that 68.5% of missense kinase mutations in cancer patients have never been observed previously, while 87.4% have been observed no more than ten times. (*b*) To compute the impact of a clinical point mutation on inhibitor binding free energy, a thermodynamic cycle can be used to relate the free energy of the wild-type and mutant kinase in the absence (top) and presence (bottom) of the inhibitor. (*c*) Summary of mutations studied in this work. Frequency of the wild-type (dark green) and mutant (green) residues for the 144 clinically-identified Abl mutations used in this study (see *Table 1* for data sources). Also shown is the frequency of residues within 5 Å (light blue) and 8 Å (blue) of the binding pocket. The number of wild-type Phe residues (n=45) and mutant Val residues (n=31) exceeded the limits of the y-axis.

- ⁴⁰ basket trials [21, 22], help prioritize candidate compounds for clinical trials, and aid the development of
- ⁴¹ next-generation inhibitors.

⁴² The long tail of rare kinase mutations frustrates prediction of drug resistance

43 While some cancer missense mutations are highly recurrent and have been characterized clinically or

44 biochemically, a "long tail" of rare mutations collectively accounts for the majority of clinically observed

⁴⁵ missense mutations (*Figure 1*a), leaving clinicians and researchers without knowledge of whether these

⁴⁶ uncharacterized mutations might lead to resistance. While rules-based and machine learning schemes

- 47 are still being assessed in oncology contexts, work in predicting drug response to microbial resistance has
- 48 shown that rare mutations present a significant challenge to approaches that seek to predict resistance

⁴⁹ to therapy [23]. Clinical cancer mutations may impact drug response through a variety of mechanisms

⁵⁰ by altering kinase activity, ATP affinity, substrate specificities, and the ability to participate in regulatory

interactions, compounding the difficulties associated with limited datasets that machine learning approaches
 face. In parallel with computational approaches, high-throughput experimental techniques such as MITE-

⁵³ Seq [24] have been developed to assess the impact of point mutations on drug response. However, the

⁵⁴ complexity of defining selection schemes that reliably correlate with *in vivo* drug effectiveness and long

⁵⁵ turn-around times might limit their ability to rapidly and reliably impact clinical decision-making.

- ⁵⁶ Alchemical free-energy methods can predict inhibitor binding affinities
- ⁵⁷ Physics-based approaches could be complementary to machine-learning and experimental techniques
- ⁵⁸ in predicting changes in TKI affinity due to mutations with few or no prior clinical observations. Modern
- ⁵⁹ atomistic molecular mechanics forcefields such as OPLS3 [25], CHARMM [26], and AMBER FF14SB [27] have
- ⁶⁰ reached a sufficient level of maturity to enable the accurate and reliable prediction of receptor-ligand binding
- ⁶¹ free energy. Alchemical free-energy methods permit receptor-ligand binding energies to be computed

- rigorously, including all relevant entropic and enthalpic contributions [28]. Encouragingly, kinase:inhibitor
- ⁶³ binding affinities have been predicted using alchemical free-energy methods with mean unsigned errors of
- 1.0 kcal/mol for CDK2, JNK1, p38, and Tyk2 [29, 30]. Beyond kinases, alchemical approaches have predicted
- the binding affinity of BRD4 inhibitors with mean absolute errors of 0.6 kcal/mol [31]. Alchemical methods
- ⁶⁶ have also been observed to have good accuracy (0.6 kcal/mol mean unsigned error for Tyk2 tyrosine kinase)
- ⁶⁷ in the prediction of relative free energies for ligand transformations within a complex whose receptor
- ⁶⁸ geometry was generated using a homology model [32].
- ⁶⁹ Alchemical approaches can predict the impact of protein mutations on free energy
- 70 Alchemical free-energy calculations have also been used to predict the impact of mutations on protein-
- protein binding [33] and protein thermostabilities [34]. Recent work has found that protein mutations can
- ⁷² be predicted to be stabilizing or destabilizing with a classification accuracy of 71% across ten proteins and 62
- ⁷³ mutations [35]. The impact of Gly to D-Ala mutations on protein stability was predicted using an alchemical
- ⁷⁴ approach with a similar level of accuracy [36]. Recently, one study has hinted at the potential utility of
- ⁷⁵ alchemical free-energy calculations in oncology by predicting the impact of a single clinical mutation on the
- ⁷⁶ binding free energies of the TKIs dasatinib and RL45 [37].
- 77 Assessing the potential for physical modeling to predict resistance to FDA-approved TKIs

Here, we ask whether physical modeling techniques may be useful in predicting whether clinically-identified 78 kinase mutations lead to drug resistance or drug sensitivity. We perform state-of-the-art relative alchemical 79 free-energy calculations using FEP+ [29], recently demonstrated to achieve sufficiently good accuracy to drive 80 the design of small-molecule inhibitors for a broad range of targets during lead optimization [28–30, 38]. We 81 compare this approach against a fast but approximate physical modeling method implemented in Prime [39] 82 (an MM-GBSA approach) in which an implicit solvent model is used to assess the change in minimized 83 interaction energy of the ligand with the mutant and wild-type kinase. We consider whether these methods 84 can predict a ten-fold reduction in inhibitor affinity (corresponding to a binding free energy change of 1.36 85 kcal/mol) to assess baseline utility. As a benchmark, we compile a set of reliable inhibitor $\Delta p | C_{e0}$ data for 144 86 clinically-identified mutants of the human kinase Abl, an important oncology target dysregulated in cancers 87 like chronic myelogenous leukemia (CML), for which six [1] FDA-approved TKIs are available. While $\Delta p | C_{so}$ 88 can approximate a dissociation constant ΔK_n , other processes contributing to changes in cell viability might 89 affect IC_{so} in ways that are not accounted for by a traditional binding experiment, motivating a quantitative 90 comparison between ΔplC_{50} and ΔK_{D} . The results of this benchmark demonstrate the potential for FEP+ to 91

 $_{92}$ predict the impact that mutations in Abl kinase have on drug binding, and a classification accuracy of 88_{82}^{93} %

⁹³ (for all statistical metrics reported in this paper, the 95% confidence intervals (CI) is shown in the form of

 (x_{lower}^{upper})), an RMSE of $1.07_{0.89}^{1.26}$ kcal/mol, and an MUE of $0.79_{0.67}^{0.92}$ kcal/mol was achieved.

95 Results

⁹⁶ Free energy calculations can recapitulate the impact of clinical mutations on TKI affinity

Alchemical free-energy calculations utilize a physics-based approach to estimate the free energy of transform-97 ing one chemical species into another, incorporating all enthalpic and entropic contributions in a physically 98 consistent manner [28, 40–42]. While relative alchemical free-energy calculations have typically been em-99 ployed in optimizing small molecules for increased potency or selectivity [29, 38, 42, 43], a complementary 100 alchemical approach can be used to compute the impact of point mutations on ligand binding affinities. 101 *Figure 1*b depicts the thermodynamic cycle that illustrates how we used relative free energy calculations to 102 compute the change in ligand binding free energy in response to the introduction of a point mutation in the 103 kinase. In the *bound* leg of the cycle, the wild-type protein: ligand complex is transformed into the mutant 104 protein:ligand complex. In the unbound leg of the cycle, the apo protein is transformed from wild-type into 105 mutant. To achieve reliable predictions with short relative free-energy calculations, a reliable receptor: ligand 106 complex structure is required with the assumption that the binding mode of wild-type and mutant are 107 similar. In this work, high-resolution co-crystal structures of wild-type Abl bound to an inhibitor were utilized 108 when available. To assess the potential for using docked inhibitor poses, we also examined two systems for 109

					(kcal/mol)		(kcal/mol)
ткі	$\mathbf{N}_{\mathrm{mut}}$	R	S	PDB	$ \Delta G_{\max} - \Delta G_{\min} $	Source	$\Delta G_{ m WT}$
axitinib	26	0	26	4wa9	2.05	[44]	-8.35
bosutinib	21	4	17	3ue4	2.79	[45]	-9.81
dasatinib	21	5	16	4xey	5.08	[45]	-11.94
imatinib	21	5	16	1орј	2.16	[45]	-9.19
nilotinib	21	4	17	3cs9	3.88	[45]	-10.74
ponatinib	21	0	21	3oxz	1.00	[45]	-11.70
subtotal	131	18	113				
erlotinib	7	1	6	Dock to 3ue4	1.73	[46]	-9.77
gefitinib	6	0	6	Dock to 3ue4	1.79	[46]	-8.84
total	144	19	125				

Table 1.	Public ΔpIC_{50} datasets for 144 Abl kinase mutations and eight tyrosine kinase inhibitors (TKIs) with
correspo	nding wild-type co-crystal structures used in this study.

 \mathbf{N}_{mut} : Total number of mutants for which ΔpIC_{50} data was available.

Number of **R**esistant, **S**usceptical mutants using 10-fold affinity change threshold.

PDB: Source PDB ID, or *Dock to 3ue4*, which used 3ue4 as the receptor for Glide-SP docking inhibitors without co-crystal structure.

 $\Delta G_{\rm WT}$: Binding free energy of inhibitor to wild-type Abl, as estimated from IC₅₀ data.

which co-crystal structures were not available (Abl:erlotinib and Abl:gefitinib) and used docking to generate
 initial coordinates.

¹¹² Compiled ΔpIC_{50} data provides a benchmark for predicting mutational resistance

To construct a benchmark evaluation dataset, we compiled a total of $144 \Delta pIC_{so}$ measurements of AbI:TKI 113 affinities, summarized in Table 1, taking care to ensure all measurements for an individual TKI were reported 114 in the same study from experiments run under identical conditions. 131 $\Delta p I C_{so}$ measurements were available 115 across the six TKIs with available co-crystal structures with wild-type Abl—26 for axitinib and 21 for bosutinib, 116 dasatinib, imatinib, nilotinib, and ponatinib. 13 $\Delta p I C_{so}$ measurements were available for the two TKIs for 117 which docking was necessary to generate Abl:TKI structures—7 for erlotinib and 6 for gefitinib. For added 118 diversity, this set includes TKIs for which Abl is not the primary target—axitinib, erlotinib, and gefitinib, All 119 mutations in this benchmark dataset have been clinically-observed (Table S1). Due to the change in bond 120 topology required by mutations involving proline, which is not currently supported by the FEP+ technology for 121 protein residue mutations, the three mutations H396P (axitinib, gefitinib, erlotinib) were excluded from our 122 assessment. As single point mutations were highly represented in the IMPACT study analyzed in Figure 1a, 123 we excluded double mutations from this work. However, the impact of mutations from multiple sites can 124 potentially be modeled by sequentially mutating each site and this will be addressed in future work. 125 Experimental ΔplC_{s_0} measurements for wild-type and mutant Abl were converted to $\Delta \Delta G$ in order 126 to make direct comparisons between physics-based models and experiment. However, computation 127 of experimental uncertainties were required to understand the degree to which differences between 128 predictions and experimental data were significant. Since experimental error estimates for measured IC₅₀s 129 were not available for the data in Table 1, we compared that data to other sources that have published 130 IC₅₀s for the same mutations in the presence of the same TKIs (*Figure 2a*,b,c). Cross-comparison of 97 131 experimentally measured $\Delta\Delta$ Gs derived from cell viability assay IC₅₀ data led to an estimate of experimental 132 variability of 0.320.36 kcal/mol root-mean square error (RMSE) that described the expected repeatability of the 133 measurements. Because multiple factors influence the IC₅₀ aside from direct effects on the binding affinity— 134 the focus of this study—we also compared $\Delta\Delta$ Gs derived from Δ plC₅₀s with those derived from binding 135 affinity measurements (ΔK_d) for which data for a limited set of 27 mutations was available (*Figure 2*d); 136 the larger computed RMSE of 0.81^{1.04}_{0.59} kcal/mol represents an estimate of the lower bound of the RMSE to 137 the IC_{so}-derived $\Delta\Delta$ Gs that we might hope to achieve with FEP+ or Prime, which were performed using 138

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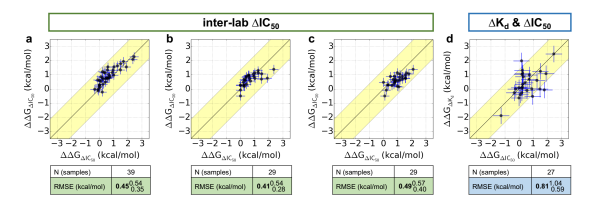


Figure 2. Cross-comparison of the experimentally measured effects that mutations in Abl kinase have on ligand binding, performed by different labs. $\Delta\Delta G$ was computed from publicly available ΔplC_{50} or ΔpK_d measurements and these values of $\Delta\Delta G$ were then plotted and the RMSE between them reported. (a) ΔplC_{50} measurements (X-axis) from [45] compared with ΔplC_{50} measurements (Y-axis) from [47]. (b) ΔplC_{50} measurements (X-axis) from [45] compared with ΔplC_{50} measurements (Y-axis) from [48]. (c) ΔplC_{50} measurements (X-axis) from [47] compared with ΔplC_{50} measurements (Y-axis) from [48]. (c) ΔplC_{50} measurements (X-axis) from [47] compared with ΔplC_{50} measurements (Y-axis) from [48]. (d) ΔplC_{50} measurements (X-axis) from [45] compared with ΔpK_d measurements (Y-axis) from [46] using non-phosphorylated Abl kinase. Scatter plot error bars in a,b,and c are ±standard error (SE) taken from the combined 97 inter-lab $\Delta\Delta G$ s derived from the ΔplC_{50} measurements, which was $0.32^{0.36}_{0.28}$. the RMSE was $0.45^{0.51}_{0.39}$ kcal/mol. Scatter plot error bars in d are the ±standard error (SE) of $\Delta\Delta G$ s derived from ΔpL_{50} and ΔpK_d from a set of 27 mutations, which is $0.58^{0.74}_{0.59}$. the RMSE was $0.81^{1.04}_{0.59}$ kcal/mol.

- ¹³⁹ non-phosphorylated models, when comparing sample statistics directly. In comparing 31 mutations for
- which phosphorylated and non-phosphorylated ΔK_d s were available, we found a strong correlation between
- the $\Delta\Delta$ Gs derived from those data (r=0.94, Supplementary *Figure S1*); the statistics of that comparison are
- similar to those of the inter-lab variability comparison.

¹⁴³ Most clinical mutations do not significantly reduce TKI potency

The majority of mutations do not lead to resistance by our 10-fold affinity loss threshold: 86.3% of the

co-crystal set (n=113) and 86.8% of the total set (n=125). Resistance mutations, which are likely to result in a

failure of therapy, constitute 13.7% of the co-crystal set (n=18) and 13.2% of the total set of mutations (n=19).

¹⁴⁷ The ΔpIC_{50} s for all 144 mutations are summarized in *Table S2—Table S7* in the Supplementary Information.

Two mutations exceeded the dynamic range of the assays (IC $_{50}$ >10,000 nM); as these two mutations clearly

raise resistance, we excluded them from quantitative analysis (RMSE and MUE) but included them in truth

table analyses and classification metrics (accuracy, specificity, sensitivity).

¹⁵¹ How accurately does physical modeling predict affinity changes for clinical Abl mutants?

From prior experience with relative alchemical free-energy calculations for ligand design, good initial receptor-152 ligand geometry was critical to obtaining accurate and reliable free energy predictions [29], so we first focused 153 on the 131 mutations in Abl kinase across six TKIs for which wild-type Abl:TKI co-crystal structures were 154 available. *Figure 3* summarizes the performance of predicted binding free-energy changes ($\Delta\Delta G$) for all 155 131 mutants in this set for both a fast MM-GBSA physics-based method that only captures interaction 156 energies for a single structure (Prime) and rigorous alchemical free-energy calculations (FEP+). Scatter plots 157 compare experimental and predicted free-energy changes ($\Delta\Delta G$) and characterize the ability of these two 158 techniques to predict experimental measurements. Statistical uncertainty in the predictions and experiment-159 to-experiment variability in the experimental values are shown as ellipse height and widths respectively. 160 The value for experimental variability was 0.32 kcal/mol, which was the standard error computed from the 161 cross-comparison in *Figure 2*. For FEP+, the uncertainty was taken to be the standard error of the average 162 from three independent runs for a particular mutation, while Prime results are deterministic and are not 163 contaminated by statistical uncertainty (see Methods). 164

To better assess whether discrepancies between experimental and computed $\Delta\Delta G$ s simply arise for known forcefield limitations or might indicate more significant effects, we incorporated an additional error model in which the forcefield error was taken to be a random error $\sigma_{\rm FF} pprox$ 0.9 kcal/mol, a value established

¹⁶⁸ form previous benchmarks on small molecules absent conformational sampling or protonation state

issues [25]. Thin error bars in *Figure 2* represent the overall estimated error due to both this forcefield error

and experimental variability or statistical uncertainty.

To assess overall quantitative accuracy, we computed both root-mean-squared error (RMSE)—which is rather sensitive to outliers, and mean unsigned error (MUE). For Prime, the MUE was $1.16_{0.96}^{1.37}$ kcal/mol and the RMSE was $1.72_{1.41}^{2.00}$ kcal/mol. FEP+, the alchemical free-energy approach, achieved a significantly higher level of quantitative accuracy with an MUE of $0.82_{0.69}^{0.95}$ kcal/mol and an RMSE of $1.11_{0.91}^{1.30}$ kcal/mol. Notably, alchemical free energy calculations come substantially closer than MMGBSA approach to the minimum achievable RMSE of $0.81_{0.59}^{1.04}$ kcal/mol (due to experimental error; *Figure 2*) for this dataset.

How accurately can physical modeling classify mutations as susceptible or resistant?

While quantitative accuracy (MUE, RMSE) is a principle metric of model performance, an application of potential interest is the ability to classify mutations as causing resistance to a specific TKI. To characterize the

accuracy with which Prime and FEP+ classified mutations in a manner that might be therapeutically relevant,

we classified mutations by their experimental impact on the binding affinity as *susceptible* (affinity for mutant

is diminished by no more than 10-fold, $\Delta\Delta G \le 1.36$ kcal/mol) or as *resistant* (affinity for mutant is diminished by least 10-fold, $\Delta\Delta G > 1.36$ kcal/mol). Summary statistics of experimental and computational predictions of

these classes are shown in *Figure 2* (bottom) as truth tables (also known as *confusion matrices*).

The simple minimum-energy scoring method Prime correctly classified 9 of the 18 resistance mutations 185 in the dataset while merely 85 of the 113 susceptible mutations were correctly classified (28 false positives). 186 In comparison, the alchemical free-energy method FEP+, which includes entropic and enthalpic contributions 187 as well as explicit representation of solvent, correctly classified 9 of the 18 resistance mutations while a 188 vast majority, 105, of the susceptible mutations were correctly classified (merely 8 false positives). Prime 189 achieved a classification accuracy of 0.72^{0.79}_{0.64}, while FEP+ achieved an accuracy that is significantly higher (both 190 in a statistical sense and in overall magnitude), achieving an accuracy of 0.87^{0.92}_{0.81}. Sensitivity (also called *true* 191 positive rate) and specificity (true negative rate) are also informative statistics in assessing the performance of 192 a binary classification scheme. For Prime, the sensitivity was $0.50^{0.73}_{0.25}$, while the specificity was $0.75^{0.83}_{0.67}$. To 193 put this in perspective, a CML patient bearing a resistance mutation in the kinase domain of Abl has an 194 equal chance of Prime correctly predicting this mutation would be resistant to one of the TKIs considered 195 here, while if the mutation was susceptible, the chance of correct prediction would be ~75%. By contrast, 196 the classification specificity of FEP+ was substantially better. For FEP+, the sensitivity was 0.50^{0.74}_{0.29} while the 197 specificity was $0.93_{0.88}^{0.97}$. There is a very high probability that FEP+ will correctly predict that one of the eight 198 TKIs studied here will remain effective for a patient bearing a susceptible mutation. 199

²⁰⁰ How sensitive are classification results to choice of cutoff?

Previous work by O'Hare et al. utilized TKI-specific thresholds for dasatinib, imatinib, and nilotinib [49], which 201 were ~2 kcal/mol. Supplementary Figure S2 shows that when our classification threshold was increased 202 to a 20-fold change in binding (1.77 kcal/mol), FEP+ correctly classified 8 of the 13 resistant mutations 203 and with a threshold of 100-fold change in binding (2.72 kcal/mol). FEP+ correctly classified the only two 204 resistant mutations (T315I/dasatinib and T315I/nilotinib). With the extant multilavered and multinodal 205 decision-making algorithms used by experienced oncologists to manage their patients' treatment, or by 206 medicinal chemists to propose candidate compounds for clinical trials, the resistant or susceptible cutoffs 207 could be selected with more nuance than the simple 10-fold affinity threshold we consider here. With a larger 208 affinity change cutoff, for example, the accuracy with which physical models predict resistance mutations 209 increases beyond 90% (Supplementary Figure S2). For the alchemical approach, the two-class accuracy was 210 $0.92_{0.87}^{0.96}$ when an affinity change cutoff of 20-fold was used while using an affinity change cutoff of 100-fold 211 further improved the two-class accuracy to $0.98^{1.00}_{0.96}$ 212

²¹³ Bayesian analysis can estimate the true error

²¹⁴ The statistical metrics—MUE, RMSE, accuracy, specificity, and sensitivity—discussed above are based on

analysis of the apparent performance of the observed modeling results compared with the observed

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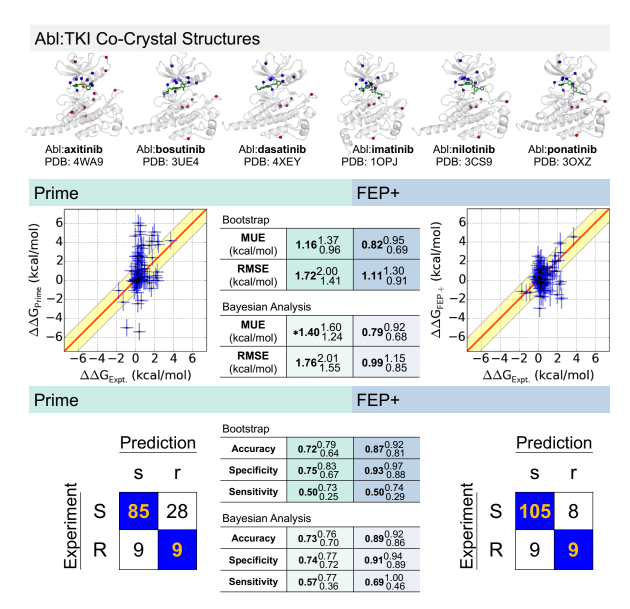


Figure 3. Comparison of experimentally-measured binding free-energy changes ($\Delta\Delta$ G) for 131 clinically observed mutations and 6 selective kinase inhibitors for which co-crystal structures of wild-type kinase with inhibitor are available. Top panel: Abl:TKI co-crystal structures used in this study with locations of clinical mutants for each inhibitor highlighted (colored from blue to red for residues nearest to farthest from ligand) in relation to TKI (green sticks) on the corresponding AbI:TKI wild-type crystal structure. *Middle panel:* Scatter plots show Prime and FEP+ computed ΔΔG compared to experiment, with ellipse widths and heights $(\pm \sigma)$ for experiment and FEP+ respectively. The red diagonal line indicates when prediction equals experiment, while the yellow shaded region indicates area in which predicted $\Delta\Delta G$ is within 1.36 kcal/mol of experiment (corresponding to a ten-fold error in predicted affinity change). $\Delta\Delta G < 0$ denotes the mutation increases the susceptibility of the kinase to the inhibitor, while $\Delta\Delta G > 0$ denotes the mutation increases the resistance of the kinase to the inhibitor. The two mutations that were beyond the concentration limit of the assay (T315I/dasatinib, L248R/imatinib) were not plotted; 129 points were plotted. Truth tables of classification accuracy, sensitivity and specificity using two-classes. Bottom panel: Truth tables and classification results include T315I/dasatinib and L248R/imatinib; 131 points were used. For MUE, RMSE, and truth table performance statistics, sub/superscripts denote 95 % CIs. Variability in the experimental data is shown as ellipse widths and uncertainty in our calculations is shown as ellipse heights. Experimental variability was computed as the standard error between IC_{s0} -derived $\Delta\Delta G$ measurements made by different labs, 0.32 kcal/mol. The statistical uncertainty in the Prime calculations was zero because the method is deterministic ($\sigma_{cal} = 0$), while the uncertainty in the FEP+ calculations was reported as the standard error, σ_{cal} , of the mean of the predicted $\Delta\Delta Gs$ from three independent runs. To better highlight true outliers unlikely to simply result from expected forcefield error, we presume forcefield error ($\sigma_{\rm FF} \approx 0.9$ kcal/mol [25]) also behaves as a random error, and represent the total estimated statistical and forcefield error ($\sqrt{\sigma_{\text{FF}}^2 + \sigma_{\text{exp/cal}}^2}$) as vertical error bars. The horizontal error bars for the experiment (σ_{exp}) was computed as the standard error between ΔplC_{50} and ΔK_d measurements, 0.58 kcal/mol. For Prime, *MUE highlights that the Bayesian model yields a value for MUE that is noticeably larger than MUE for observed data due to the non-Gaussian error distribution of Prime.

experimental data via sample statistics. However, this analysis considers a limited number of mutants. 216 and both measurements and computed values are contaminated with experimental or statistical error. 21 To obtain an estimate of the *intrinsic performance* of our physical modeling approaches, accounting for 218 known properties of the experimental variability and statistical uncertainties, we used a hierarchical Bayesian 219 model (detailed in the Methods) to infer posterior predictive distributions from which expectations and 95% 220 predictive intervals could be obtained. The results of this analysis are presented in Figure 3 (central tables). 221 FEP+ is significantly better than Prime at predicting the impact of mutations on TKI binding affinities, as the 222 apparent performance (using the original observations) as well as the intrinsic performance (where Bayesian 223 analysis was used to correct for statistical uncertainty or experimental variation) were well-separated outside 224 their 95% confidence intervals in nearly all metrics. Applying the Bayesian model, the MUE and RMSE for 225 FEP+ was $0.79_{0.68}^{0.92}$ kcal/mol and $0.99_{0.85}^{1.15}$ kcal/mol respectively (N=129). For the classification metrics accuracy, specificity, and sensitivity, the model yields $0.89_{0.86}^{0.92}$, $0.91_{0.89}^{0.94}$, and $0.69_{0.46}^{1.00}$ respectively (N=131). The intrinsic 226 227 RMSE and MUE of Prime was 1.76^{2.01} kcal/mol and 1.40^{1.60} kcal/mol (N=129) respectively, and the classification 228 accuracy, specificity, and sensitivity was $0.73_{0.70}^{0.76}$, $0.74_{0.72}^{0.77}$, and $0.57_{0.36}^{0.77}$ respectively (N=131). The intrinsic MUE of 229 Prime obtained by this analysis is larger than the observed MUE reflecting the non-Gaussian, fat-tailed error 230 distributions of Prime results. 231

²³² Is the impact of point mutations on drug binding equally well-predicted for the six TKIs?

The impact of point mutations on drug binding are not equally well predicted for the six TKIs. Figure 4 233 expands the results in *Figure 3* on a TKI-by-TKI basis to dissect the particular mutations in the presence of 234 a specific TKI. Prime and FEP+ correctly predicted that most mutations in this dataset (N=26) do not raise 235 resistance to axitinib, though FEP+ predicted 4 false positives compared with 3 false positives by Prime. The 236 MUE and RMSE of FEP+ was excellent for this inhibitor, $0.70_{0.50}^{0.93}$ kcal/mol and $0.91_{0.64}^{1.14}$ kcal/mol respectively. 237 While the classification results for bosutinib (N=21) were equally well predicted by Prime as by FEP+, FEP+ was 238 still able to achieve superior, but not highly significant, predictive performance for the quantitative metrics 239 MUE and RMSE, which were $0.96^{1.42}_{0.55}$ kcal/mol and $1.41^{1.97}_{0.77}$ kcal/mol respectively (FEP+) and $1.13^{1.83}_{0.60}$ kcal/mol and 240 $1.80_{0.92}^{2.62}$ kcal/mol respectively (Prime). For dasatinib, FEP+ achieved an MUE and RMSE of $0.76_{0.49}^{1.13}$ kcal/mol and 241 1.07^{1.57}_{0.59} kcal/mol respectively whereas the results were, as expected, less quantitatively predictive for Prime 242 (N=20). The results for imatinib were similar to those of dasatinib above, where the MUE and RMSE for FEP+ 243 were $0.82_{0.53}^{1.15}$ kcal/mol and $1.09_{0.69}^{1.43}$ kcal/mol respectively (N=20). Nilotinib, a derivative of imatinib, led to nearly 244 identical quantitative performance results for FEP+ with an MUE and RMSE of $0.82^{1.12}_{0.57}$ kcal/mol and $1.06^{1.39}_{0.69}$ 245 kcal/mol respectively (N=21). Similar to axitinib, ponatinib presented an interesting case because there were 246 no mutations in this dataset that raised resistance to it. Despite the wide dynamic range in the computed 247 values of $\Delta\Delta G$ for other inhibitors, FEP+ correctly predicted a very narrow range of $\Delta\Delta Gs$ for this drug. This 248 is reflected in the MUE and RMSE of 0.87^{1.16}_{0.62} kcal/mol and 1.09^{1.46}_{0.70} kcal/mol respectively, which are in-line with 249 the MUEs and RMSEs for the other TKIs. 250

²⁵¹ Understanding the origin of mispredictions

Resistance mutations that are mispredicted as susceptible (false negatives) are particularly critical because 252 they might mislead the clinician or drug designer into believing the inhibitor will remain effective against 253 the target. Which resistance mutations did FEP+ mispredict as susceptible? Nine mutations were classified 254 by FEP+ to be susceptible when experimentally measured ΔplC_{so} data indicate the mutations should have 255 increased resistance according to our 10-fold affinity cutoff for resistance. Notably, the 95% confidence 256 intervals for five of these mutations spanned the 1.36 kcal/mol threshold, indicating these misclassifications 257 are not statistical significant when the experimental error and statistical uncertainty in FEP+ are accounted for: 258 bosutinib/L248R ($\Delta\Delta G_{FEP+}$ =1.32^{1.94}_{0.70} kcal/mol), imatinib/E255K ($\Delta\Delta G_{FEP+}$ =0.43^{3.05}_{-2.19} kcal/mol), imatinib/Y253F 259 $(\Delta\Delta G_{FEP+}=0.95^{1.64}_{0.26}$ kcal/mol), and nilotinib/Y253F ($\Delta\Delta G_{FEP+}=0.89^{1.69}_{0.09}$ kcal/mol). The bosutinib/V299L mutation 260 was also not significant because the experimental $\Delta\Delta G$, $1.70_{1.08}^{2.33}$ kcal/mol, included the 1.36 kcal/mol cutoff; the value of $\Delta\Delta G$ predicted by FEP+ for this mutation was $0.91_{0.79}^{1.02}$ kcal/mol, the upper bound of the predicted 261 262 value was within 0.06 kcal/mol of the lower bound of the experimental value. 263

Four mutations, however, were misclassified to a degree that is statistically significant given their 95% con-

fidence intervals: dasatinib/T315A. bosutinib/T315I. imatinib/E255V. and nilotinib/E255V. For dasatinib/T315A. 265 although the T315A mutations for bosutinib, imatinib, nilotinib, and ponatinib were correctly classified as 266 susceptible, the predicted free energy changes for these four TKIs were consistently much more negative 267 than the corresponding experimental measurements, just as for dasatinib/T315A, indicating there might be a 268 generic driving force contributing to the errors in T315A mutations for these five TKIs. Abl is known to be able 269 to adopt many different conformations (including DFG-in and DFG-out), and it is very likely that the T315A 270 mutation will induce conformational changes in the app protein [50], which was not adequately sampled in 271 the relatively short simulations, leading to the errors for T315A mutations for these TKIs. By comparison, 272 the T315I mutations for axitinib, bosutinib, imatinib, nilotinib, and ponatinib were all accurately predicted 273 with the exception of bosutinib/T315I being the only misprediction, suggesting an issue specific to bosutinib. 274 The complex electrostatic interactions between the 2.4-dichloro-5-methoxyphenyl ring in bosutinib and the 275 adjacent positively charged amine of the catalytic Lys271 may not be accurately captured by the fixed-charge 276 OPLS3 force field, leading to the misprediction for bosutinib/T315I mutation. 277 Insufficient sampling might also belie the imatinib/E255V and nilotinib/E255V mispredictions because 278 they reside in the highly flexible P-loop. Since E255V was a charge change mutation, we utilized a workflow 279 that included a transmutable explicit ion (see Methods). The distribution of these ions in the simulation box 280

around the solute might not have converged to their equilibrium state on the relatively short timescale of
 our simulations (5 ns), and the insufficient sampling of ion distributions coupled with P-loop motions might
 lead to misprediction of these two mutations.

²⁸⁴ How accurately can the impact of mutations be predicted for docked TKIs?

To assess the potential for utilizing physics-based approaches in the absence of a high-resolution experimen-285 tal structure, we generated models of Abl bound to two TKIs—erlotinib and gefinitib—for which co-crystal 286 structures with wild-type kinase are not currently available. In *Figure 5*, we show the Abl:erlotinib and 287 Abl:gefitinib complexes that were generated using a docking approach (Glide-SP, see Methods). These two 288 structures were aligned against the co-crystal structures of EGFR:erlotinib and EGFR:gefinitib to highlight the 289 structural similarities between the binding pockets of Abl and EGFR and the TKI binding mode in Abl versus 290 EGFR. As an additional test of the sensitivity of FEP+ to system preparation, a second set of Abl:erlotinib and 291 Abl:gefitinib complexes was generated in which crystallographic water coordinates were transferred to the 292 docked inhibitor structures (see Methods). 293 Alchemical free-energy simulations were performed on 13 mutations between the two complexes; 7 294

²⁹⁴ mutations for erlotinib and 6 mutations were performed on 15 mutations between the two complexes, 7 ²⁹⁵ mutations for erlotinib and 6 mutations for gefitinib. The quantitative accuracy of FEP+ in predicting the ²⁹⁶ value of $\Delta\Delta G$ was excellent—MUE and RMSE of $0.58^{0.86}_{0.35}$ kcal/mol and $0.80^{1.09}_{0.44}$ kcal/mol respectively if crystal ²⁹⁷ waters are omitted, and $0.50^{0.78}_{0.26}$ kcal/mol and $0.69^{0.97}_{0.35}$ kcal/mol if crystal waters were restored after docking. ²⁹⁸ Encouragingly, these results indicate that our initial models of Abl bound to erlotinib and gefitinib were ²⁹⁹ reliable because the accuracy and dependability of our FEP+ calculations were not sensitive to crystallographic ³⁰⁰ waters. Our secondary concern was the accuracy with which the approach classified mutations as resistant ³⁰¹ or susceptible.

While the results presented in (*Figure 5*) indicate that FEP+ is capable of achieving good quantitative 302 accuracy when a co-crystal structure is unavailable, it is important to understand why a mutation was 303 predicted to be susceptible but was determined experimentally to be resistant. F317I was the one mutation 304 that increased resistance to erlotinib (or gefitinib) because it destabilized binding by more than 1.36 kcal/mol-305 1.35^{1.67}_{1.03} kcal/mol (gefitinib) and 1.58^{1.90}_{1.26} kcal/mol (erlotinib), but the magnitude of the experimental uncertainty 306 means we are unable to confidently discern whether this mutation induces more than 10-fold resistance 307 to either TKI. Therefore, the one misclassification by FEP+ in *Figure 5* is not statistically significant and the 308 classification metrics presented there underestimate the nominal performance of this alchemical free-energy 309 method. 310

311 Discussion

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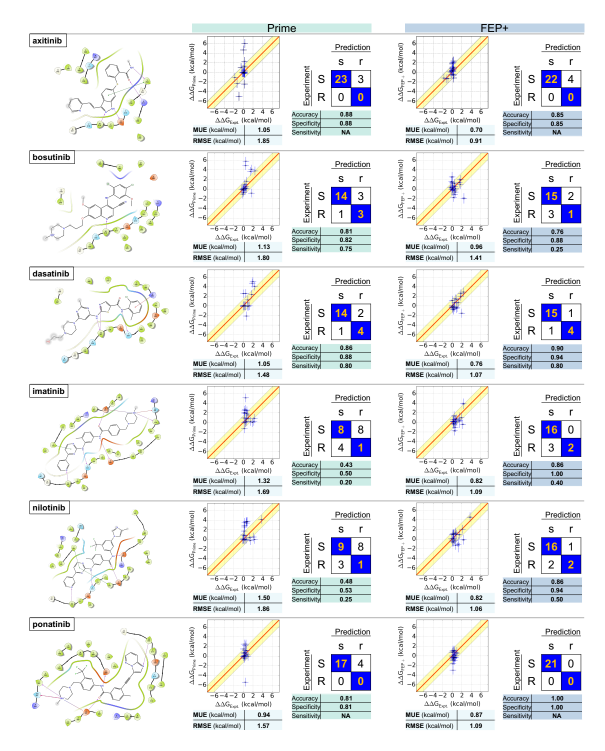


Figure 4. Physical modeling accuracy in computing the impact of clinical Abl mutations on selective inhibitor binding. Ligand interaction diagrams for six selective FDA-approved tyrosine kinase inhibitors (TKIs) for which co-crystal structures with Abl were available (left). Comparisons for clinically-observed mutations are shown for FEP+ (right) and Prime (left). For each ligand, computed vs. experimental binding free energies ($\Delta\Delta G$) are plotted with MUE and RMSE (units of kcal/mol) depicted below. Truth tables are shown to the right. Rows denote *true* susceptible (S, $\Delta\Delta G \le 1.36$ kcal/mol) or resistant (R, $\Delta\Delta G > 1.36$ kcal/mol) experimental classes using a 1.36 kcal/mol. Correct predictions populate diagonal elements (orange text), incorrect predictions populate off-diagonals. Accuracy, specificity, and sensitivity for two-class classification are shown below the truth table. Elliptical point sizes and error bars in the scatter plots depict estimated uncertainty/variability and error respectively ($\pm\sigma$) of FEP+ values (vertical size) and experimental values (horizontal size). Note: The sensitivity for axitinib and ponatinib is NA, because there is no resistant mutation for these two drugs.

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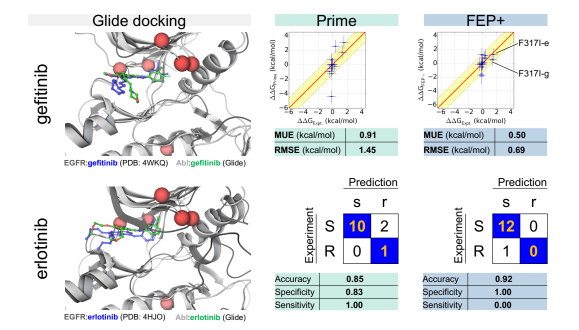


Figure 5. Predicting resistance mutations using FEP+ for inhibitors for which co-crystal structures with wild-type kinase are not available. The docked pose of Abl:erlotinib is superimposed on the co-crystal structure of EGFR:erlotinib; erlotinib docked to Abl (light gray) is depicted in green and erlotinib bound to EGFR (dark gray) is depicted in blue. The docked pose of Abl:gefitinib is superimposed on the co-crystal structure of EGFR:gefitinib; gefitinib docked to Abl (light gray) is depicted in green and erlotinib bound to EGFR (dark gray) is depicted in blue. The docked pose of Abl:gefitinib is superimposed on the co-crystal structure of EGFR:gefitinib; gefitinib docked to Abl (light gray) is depicted in green and gefitinib bound to EGFR (dark gray) is depicted in blue. The locations of clinical mutants for each inhibitor are highlighted (red spheres). The overall RMSEs and MUEs for Prime (center) and FEP+ (right) and two-class accuracies are also shown in the figure. Computed free energy changes due to the F317I mutation for erlotinib (-e) and gefitinib (-g) are highlighted in the scatter plot. FEP+ results are based on the docked models prepared with crystal waters added back while the Prime (an implicit solvent model) results are based on models without crystallographic water.

³¹² Physics-based modeling can reliably predict when a mutation elicits resistance to therapy

³¹³ The results presented in this work are summarized in *Table 2*. The performance metrics summarized in

Table 2 indicates that the set of 131 mutations for the six TKIs in which co-crystal structures were available is
 on par with the complete set (144 mutations), which included results based on AbI:TKI complexes generated
 from docking models. The performance results for the 13 mutations for the two TKIs (erlotinib and gefitinib)
 in which co-crystal structures were unavailable exhibited good quantitative accuracy (MUE and RMSE) and

³¹⁸ good classification power.

³¹⁹ Overall (N=144), the MM-GBSA approach Prime classified mutations with good accuracy $(0.73^{0.80}_{0.66})$ and ³²⁰ specificity $(0.76^{0.84}_{0.69})$ while the alchemical approach FEP+ was a significant improvement in classification ³²¹ accuracy $(0.88^{0.93}_{0.82})$ and specificity $(0.94^{0.98}_{0.89})$. The quantitative accuracy with which Prime was able to predict the ³²² experimentally measured change in AbI:TKI binding (N=142) characterized by RMSE and MUE was $1.70^{1.98}_{1.40}$ ³²³ kcal/mol and $1.14^{1.35}_{0.93}$ kcal/mol respectively. In stark contrast, the quantitative accuracy of FEP+ was statistically ³²⁴ superior to Prime with an RMSE and an MUE of $1.07^{1.26}_{0.89}$ kcal/mol and $0.79^{0.92}_{0.67}$ kcal/mol respectively.

From the perspective of a clinician, classification rate would be an important metric to measure the 325 predictive power of technologies such as Prime and FEP+. To test the hypothesis that reducing the large 326 spread in Prime predictions could improve its classification rate, we scaled the computed relative free 32 energies (by 1/2, 1/3, and by 0.23, which was the optimal factor that gives lowest RMSE) and recalculated 328 the classification metrics (Table 58). As expected, the MUE and RMSE were improved but the specificity of 329 Prime was drastically diminished; as MUE and RMSE improved, it became increasingly unable to identify 330 resistance mutations. Scaling FEP+ eliminated its sensitivity and a naïve model (where all free energies were 331 set to 0.00 kcal/mol) had zero sensitivity. Lastly, we constructed a consensus model in which free energies 332 were a weighted average of scaled Prime and FEP+. However, this model also had no sensitivity. It appears 333 difficult to improve upon the predictive power of FEP+ by statistical operations. 334

To address the impact of picking a cutoff to classify predicted free energies as resistant or sensitizing, we 335 computed ROC curves for the various predicted datasets: Prime (scaled and non-scaled), FEP+ (scaled and 336 non-scaled), naïve model, and consensus model (constructed from scaled Prime and scaled FEP+, see above). 337 ROC curves are independent of a linear transformation on the predicted dataset. Therefore, ROC curves 338 and ROC-AUCs for scaled and non-scaled Prime were identical, as well as scaled and non-scaled FEP+. ROC 339 curves for these six sets of predictions are presented in Supplementary Figure S3. ROC-AUC for FEP+ was 340 $0.75_{0.61}^{0.90}$ (n=144); ROC-AUC for Prime was $0.66_{0.52}^{0.81}$ (n=144); ROC-AUCs for the naïve model and consensus model 341 were $0.50^{0.50}_{0.50}$ (n=144) and $0.78^{0.90}_{0.67}$ (n=144) respectively. These results show that Prime apparently has poor 342 discriminatory power (ROC-AUC in [0.6,0.7]) while FEP+ apparently has fair discriminatory power (ROC-AUC 343 in [0.7,0.8]). 344

³⁴⁵ Hierarchical Bayesian model estimates global performance (N=144)

A hierarchical Bayesian approach was developed to estimate the intrinsic accuracy of the models when the noise in the experimental and predicted values of $\Delta\Delta G$ was accounted for. Utilizing this approach, the MUE and RMSE for Prime was found to be $1.39_{1.23}^{1.58}$ kcal/mol and $1.75_{1.55}^{1.98}$ kcal/mol (N=142) respectively. The accuracy, specificity, and sensitivity of Prime was found using this method to be $0.74_{0.71}^{0.76}$, $0.75_{0.73}^{0.77}$, and $0.59_{0.40}^{0.78}$ (N=144) respectively. The MUE and RMSE of FEP+ was found to be $0.76_{0.66}^{0.87}$ kcal/mol and $0.95_{0.82}^{1.09}$ kcal/mol (N=142) respectively, which is significantly better than Prime. Likewise, a clearer picture of the true classification accuracy, specificity, and sensitivity of FEP+ was found— $0.90_{0.86}^{0.90}$, $0.92_{0.95}^{0.95}$, and $0.68_{1.00}^{1.00}$ respectively.

Examining the physical and chemical features of outliers

³⁵⁴ Current alchemical approaches neglect effects that will continue to improve accuracy

³⁵⁵ The high accuracy of FEP+ is very encouraging, and the accuracy can be further improved with more accurate

³⁵⁶ modeling of a number of physical chemical effects not currently considered by the method. While highly

optimized, the fixed-charged OPLS3 [25] force field can be further improved by explicit consideration of

polarizability effects [51], as hinted by some small-scale benchmarks [52]. These features could be especially

³⁵⁹ important for bosutinib, whose 2,4-dichloro-5-methoxyphenyl ring is adjacent to the positively charged amine

³⁶⁰ of the catalytic Lys271. Many simulation programs also utilize a long-range isotropic analytical dispersion

Dataset	Method	N _{quant}	MUE	RMSE	N _{class}	Accuracy	Specificity	Sensitivity
			(kcal/mol)	(kcal/mol)				
all	FEP+	142	$0.79_{0.67}^{0.92}$	$1.07^{1.26}_{0.89}$	144	$0.88_{0.82}^{0.93}$	$0.94_{0.89}^{0.98}$	$0.47^{0.69}_{0.25}$
all	Prime	142	$1.14_{0.93}^{1.35}$	$1.70^{1.98}_{1.40}$	144	$0.73_{0.66}^{0.80}$	$0.76_{0.69}^{0.84}$	$0.53_{0.30}^{0.76}$
xtals	FEP+	129	$0.82_{0.69}^{0.95}$	$1.11_{0.91}^{1.30}$	131	$0.87_{0.81}^{0.92}$	$0.93_{0.88}^{0.97}$	$0.50_{0.29}^{0.74}$
xtals	Prime	129	$1.16_{0.96}^{1.37}$	$1.72^{2.00}_{1.41}$	131	$0.72_{0.64}^{0.79}$	$0.75_{0.67}^{0.83}$	$0.50_{0.25}^{0.73}$
axitinib	FEP+	26	$0.70_{0.50}^{0.93}$	$0.91^{1.14}_{0.64}$	26	$0.85_{0.69}^{0.96}$	$0.85_{0.69}^{0.96}$	NA
axitinib	Prime	26	$1.05^{1.71}_{0.53}$	$1.85^{2.61}_{0.96}$	26	$0.88^{1.00}_{0.73}$	$0.88^{1.00}_{0.73}$	NA
bosutinib	FEP+	21	$0.96^{1.42}_{0.55}$	$1.41^{1.97}_{0.77}$	21	$0.76^{0.95}_{0.57}$	$0.88^{1.00}_{0.71}$	$0.25^{1.00}_{0.00}$
bosutinib	Prime	21	$1.13^{1.83}_{0.60}$	$1.80^{2.62}_{0.92}$	21	$0.81_{0.62}^{0.95}$	$0.82^{1.00}_{0.62}$	$0.75^{1.00}_{0.00}$
dasatinib	FEP+	20	$0.76^{1.13}_{0.49}$	$1.07^{1.57}_{0.59}$	21	$0.90^{1.00}_{0.76}$	$0.94^{1.00}_{0.79}$	$0.80^{1.00}_{0.33}$
dasatinib	Prime	20	$1.05^{1.54}_{0.61}$	$1.48^{1.92}_{0.95}$	21	$0.86^{1.00}_{0.71}$	$0.88^{1.00}_{0.69}$	$0.80^{1.00}_{0.33}$
imatinib	FEP+	20	$0.82^{1.15}_{0.53}$	$1.09^{1.43}_{0.69}$	21	$0.86^{1.00}_{0.71}$	$1.00^{1.00}_{1.00}$	$0.40_{0.00}^{0.83}$
imatinib	Prime	20	$1.32_{0.91}^{1.81}$	$1.69^{2.26}_{1.15}$	21	$0.43_{0.24}^{0.67}$	$0.50_{0.25}^{0.75}$	$0.20_{0.00}^{0.67}$
nilotinib	FEP+	21	$0.82^{1.12}_{0.57}$	$1.06_{0.69}^{1.39}$	21	$0.86^{1.00}_{0.67}$	$0.94_{0.80}^{1.00}$	$0.50^{1.00}_{0.00}$
nilotinib	Prime	21	$1.50^{1.97}_{1.06}$	$1.86_{1.43}^{2.25}$	21	$0.48_{0.24}^{0.67}$	$0.53_{0.29}^{0.75}$	$0.25^{1.00}_{0.00}$
ponatinib	FEP+	21	$0.87^{1.16}_{0.62}$	$1.09^{1.46}_{0.70}$	21	$1.00^{1.00}_{1.00}$	$1.00^{1.00}_{1.00}$	NA
ponatinib	Prime	21	$0.94_{0.50}^{1.54}$	$1.57^{2.44}_{0.69}$	21	$0.81_{0.62}^{0.95}$	$0.81_{0.62}^{0.95}$	NA
Glide	FEP+	13	$0.50_{0.26}^{0.78}$	$0.69_{0.35}^{0.97}$	13	$0.92^{1.00}_{0.77}$	$1.00^{1.00}_{1.00}$	$0.00^{0.00}_{0.00}$
Glide	Prime	13	0.91 ^{1.56} _{0.39}	$1.45_{0.54}^{2.22}$	13	$0.85^{1.00}_{0.62}$	$0.83^{1.00}_{0.58}$	$1.00^{1.00}_{0.00}$

Table 2. Summary of FEP+ and Prime statistics in predicting mutational resistance or sensitivity to FDA-approved	
TKIs.	

N_{quant}: Number of mutations for which quantitative metrics were evaluated; N_{class}: Number mutations for which classification metrics were evaluated; All: All mutations; xtals: All mutations for which co-crystal structures were available; Glide: erlotinib and gefitinib

Accuracy, specificity, and sensitivity were computed to assess two-class prediction performance:

resistant ($\Delta\Delta G > 1.36$ kcal/mol) or susceptible ($\Delta\Delta G \le 1.36$ kcal/mol).

95% CIs (sub-/superscripts) were estimated from 1000 bootstrap replicates. Note: The sensitivity for axitinib and ponatinib is NA, because there is no resistant mutation for these two drugs.

correction intended to correct for the truncation of dispersion interactions at finite cutoff, which can induce 361 an error in protein-ligand binding free energies that depends on the number of ligand heavy atoms being 362 modified [53]; recently, efficient Lennard-Jones PME methods [54, 55] and perturbation schemes [53] have 363 been developed that can eliminate the errors associated with this truncation. While the currently employed 364 methodology for alchemical transformations involving a change in system charge (see Methods) reduces 365 artifacts that depend on the simulation box size and periodic boundary conditions, the explicit ions that were 366 included in these simulations may not have sufficiently converged to their equilibrium distributions in these 367 relatively short simulations. Kinases and their inhibitors are known to possess multiple titratable sites with 368 either intrinsic or effective pK_as near physiological pH, while the simulations here treat protonation states 369 and proton tautomers fixed throughout the bound and unbound states; the accuracy of the model can be 370 further improved with the protonation states or tautomers shift upon binding or mutation considered [56, 57]. 371 Similarly, some systems display significant salt concentration dependence [58], while the simulations for 372 some systems reported here did not rigorously mimic all aspects of the experimental conditions of the cell 373 viability assays. 374

$_{375}$ Experimentally observed IC₅₀ changes can be caused by other physical mechanisms

³⁷⁶ While we have shown that predicting the direct impact of mutations on the binding affinity of ATP-competitive

³⁷⁷ tyrosine kinase inhibitors for a single kinase conformation has useful predictive capacity, many additional

- physical effects that can contribute to cell viability are not currently captured by examining only the predicted
- ³⁷⁹ change in inhibitor binding affinity. For example, kinase missense mutations can also shift the populations

of kinase conformations (which may affect ATP and inhibitor affinities differentially), modulate ATP affinity,

- modulate affinity for protein substrate, or modulate the ability of the kinase to be regulated or bounded
- ³⁸² by scaffolding proteins. These physical mechanisms might affect the IC₅₀s of cell viability assays but not
- necessarily the binding affinity of the inhibitors. While many of these effects are in principle tractable by
- ³⁸⁴ physical modeling in general (and alchemical free energy methods in particular), it is valuable to examine our
- ³⁸⁵ mispredictions and outliers to identify whether any of these cases is likely to induce resistance (as observed
- $_{386}$ by ΔpIC_{50} shifts) by one of these alternative mechanisms.
- ³⁸⁷ Other physical mechanisms of resistance are likely similarly computable.

A simple threshold of 10-fold TKI affinity change is a crude metric for classifying resistance or susceptibility 388 due to the myriad biological factors that contribute to the efficacy of a drug in a person. Except for affecting 389 the binding affinity of inhibitors, missense mutations can also cause drug resistance through other physical 390 mechanisms including induction of splice variants or alleviation of feedback. While the current study only 391 focused on the effect of mutation on drug binding affinity, resistance from these other physical mechanisms 392 could be similarly computed using physical modeling. For example, some mutations are known to activate 393 the kinase by increasing affinity to ATP, which could be computed using the same thermodynamic cycle 394 utilized here for inhibitors. 395

Conclusion

Revolutionary changes in computing power—especially the arrival of inexpensive graphics processors 397 (GPUs)—and software automation have enabled alchemical free-energy calculations to impact drug discovery 398 and life sciences projects in previously unforeseen ways. In this communication, we tested the hypothesis 399 that FEP+, a fully-automated relative-alchemical free-energy workflow, had reached the point where it can 400 accurately and reliably predict how clinically-observed mutations in Abl kinase alter the binding affinity of 401 eight FDA-approved TKIs. To establish the potential predictive impact of current-generation alchemical free 402 energy calculations—which incorporate entropic and enthalpic effects and the discrete nature of aqueous 403 solvation—compared to a simpler physics-based approach that also uses modern forcefields but scores a 404 single minimized conformation, we employed a second physics-based approach (Prime). This simpler physics-405 based model, which uses an implicit model of solvation to score the energetic changes in interaction energy 406 that arise from the mutation, was able to capture a useful amount of information to achieve substantial 407 predictiveness with an MUE of $1.14_{0.93}^{1.35}$ kcal/mol (N=142), RMSE of $1.70_{1.40}^{1.98}$ kcal/mol respectively (N=142), and 408 classification accuracy of 0.73^{0.80}_{0.66} (N=144). Surpassing these good results, we went on to demonstrate that 409 FEP+ is able to achieve superior predictive performance— MUE of 0.79^{0.92}_{0.67} kcal/mol (N=142), RMSE of 1.07^{1.26}_{0.89} 410 kcal/mol (N=142), and classification accuracy of $0.88_{0.82}^{0.93}$ (N=144). While future enhancements to the workflows 411 for Prime and FEP+ to account for additional physical and chemical effects are likely to improve predictive 412 performance further, the present results are of sufficient quality and achievable on a sufficiently rapid 413 timescale (with turnaround times ~6 hours/calculation) to impact research projects in drug discovery and the 414 life sciences. With exponential improvements in computing power, we anticipate the domains of applicability 415 for alchemical free-energy methods such as FEP+ will take on increasingly integrated roles to impact projects. 416 This work illustrates how the domain of applicability for alchemical free-energy methods is much larger 417 than previously appreciated, and might further be found to include new areas as research progresses: 418 aiding clinical decision-making in the selection of first- or second-line therapeutics guided by knowledge 419 of likely subclonal resistance; identifying other selective kinase inhibitors (or combination therapies) to 420 which the mutant kinase is susceptible; supporting the selection of candidate molecules to advance to 421 clinical trials based on anticipated activity against likely mutations; facilitating the enrollments of patients in 422 mechanism-based basket trials; and generally augmenting the armamentarium of precision oncology. 423

424 Methods

425 System preparation

All system preparation utilized the Maestro Suite (Schrödinger) version 2016-4. Comparative modeling to add

427 missing residues using a homologous template made use of the Splicer tool, while missing loops modeled

vithout a template used Prime. All tools employed default settings unless otherwise noted. The Abl wild-type

sequence used in building all Abl kinase domain models utilized the ABL1_HUMAN Isoform IA (P00519-1)

⁴³⁰ UniProt gene sequence spanning S229–K512. Models were prepared in non-phosphorylated form. We used ⁴³¹ a residue indexing convention that places the Thr gatekeeper residue at position 315 to match common

usage; an alternate indexing convention utilized in experimental X-ray structures for Abl:imatinib (PDB: 10PI)

433 [59] and Abl:dasatinib (PDB: 4XEY) [60] was adjusted to match our convention.

Complexes with co-crystal structures. Chain B of the experimental structure of Abl:axitinib (PDB: 434 4WA9) [44] was used, and four missing residues at the N- and C-termini were added using homology 435 modeling with PDB 3IK3 [61] as the template following alignment of the respective termini of the kinase 436 domain. Chain B was selected because chain A was missing an additional 3 and 4 residues at the N- and 437 C-termini, respectively, in addition to 3- and 20-residue loops, both of which were resolved in chain B. All 438 missing side chains were added with Prime. The co-crystal structure of Abl:bosutinib (PDB: 3UE4) [62] was 439 missing 4 and 10 N- and C-terminal residues respectively in chain A that were built using homology modeling 440 with 3IK3 as the template. All loops were resolved in chain A (chain B was missing two residues in the P-loop 441 O252 and Y253). All missing side chains were added with Prime. The co-crystal structure of Abl:dasatinib 442 (PDB: 4XEY) [60] was missing 2 and 9 N- and C-terminal residues, respectively, that were built via homology 443 modeling using 3IK3 as the template. A 3 residue loop was absent in chain B but present in chain A: chain 444 A was chosen. The co-crystal structure of Abl:imatinib (PDB: 10Pl) [59] had no missing loops. Chain B was 445 used because chain A was missing two C-terminal residues that were resolved in chain B. A serine was 446 present at position 336 (index 355 in the PDB file) and was mutated to asparagine using Prime to match 447 the human wild-type reference sequence (P00519-1). The co-crystal structure of Abl:nilotinib (PDB: 3CS9) 448 [63] contained four chains in the asymmetric unit all of which were missing at least one loop. Chain A was 449 selected because its one missing loop involved the fewest number of residues of the four chains: chain A 450 was missing 4 and 12 N- and C-terminal residues, respectively, that were built using homology modeling 451 with 3IK3 as the template. A 4-residue loop was missing in chain A (chain B and C were missing two loops. 452 chain D was missing a five residue loop) that was built using Prime. The co-crystal structure of Abl:ponatinib 453 (PDB: 30X7) [64] contained only one chain in the asymmetric unit. It had two missing loops, one 4 residues 454 (built using Prime) and one 12 residues (built using homology modeling with 30Y3 [64] as the template). 455 Serine was present at position 336 and was mutated to Asn using Prime to match the human wild-type 456 reference sequence (P00519-1). Once the residue composition of the six Abl:TKI complexes were normalized 457 to have the same sequence, the models were prepared using Protein Preparation Wizard. Bond orders 458 were assigned using the Chemical Components Dictionary and hydrogen atoms were added. Missing side 459 chain atoms were built using Prime. Termini were capped with N-acetyl (N-terminus) and N-methyl amide 460 (C-terminus). If present, crystallographic water molecules were retained. Residue protonation states (e.g. 461 Asp381 and Asp421) were determined using PROPKA [65] with a pH range of 5.0–9.0. Ligand protonation 462 state was assigned using PROPKA with pH equal to the experimental assay. Hydrogen bonds were assigned 463 by sampling the orientation of crystallographic water, Asn and Gln flips, and His protonation state. The 464 positions of hydrogen atoms were minimized while constraining heavy atoms coordinates. Finally, restrained 465 minimization of all atoms was performed in which a harmonic positional restraint (25.0 kcal/mol/Å²) was 466 applied only to heavy atoms. *Table S9* summarizes the composition of the final models used for FEP. 467

Complexes without co-crystal structures. Co-crystal structures of Abl bound to erlotinib or gefitinib 468 were not publicly available. To generate models of these complexes, Glide-SP [66] was utilized to dock 469 these two compounds into an Abl receptor structure. Co-crystal structures of these two compounds bound 470 to EGFR were publicly available and this information was used to obtain initial ligand geometries and to 471 establish a reference binding mode against which our docking results could be structurally scored. The Abl 472 receptor structure bound to bosutinib was used for docking because its structure was structurally similar to 473 that of EGFR in the erlotinib- (PDB: 4HIO) [67] and gefitinib-bound (PDB: 4WKO) [68] co-crystal structures. 474 Abl was prepared for docking by using the Protein Preparation Wizard (PPW) with default parameters. 475 Crystallographic waters were removed but their coordinates retained for a subsequent step in which they 476

were optionally reintroduced. Erlotinib and gefitinib protonation states at $pH7.0\pm2.0$ were determined using 477 Epik [69]. Docking was performed using the Glide-SP workflow. The receptor grid was centered on bosutinib. 478 The backbone NH of Met318 was chosen to participate in a hydrogen bonding constraint with any hydrogen 479 bond donor on the ligand. The hydroxyl of T315 was allowed to rotate in an otherwise rigid receptor. Ligand 480 docking was performed with enhanced sampling; otherwise default settings were used. Epik state penalties 481 were included in the scoring. The 16 highest ranked (Glide-SP score) poses were retained for subsequent 482 scoring. To determine the docked pose that would be subsequently used for free energy calculations, the 483 ligand heavy-atom RMSD between the 16 poses and the EGFR co-crystal structures (PDB IDs 4HIO and 4WKO) 484 was determined. The pose in which erlotinib or gefitinib most structurally resembled the EGFR co-crystal 485 structure (lowest heavy-atom RMSD) was chosen as the pose for subsequent FEP+. Two sets of complex 486 structures were subjected to free energy calculations to determine the effect of crystal waters: In the first 487 set, without crystallographic waters, the complexes were prepared using Protein Prep Wizard as above. In 488 the second set, the crystallographic waters removed prior to docking were added back, and waters in the 489 binding pocket that clashed with the ligand were removed. 490

⁴⁹¹ Force field parameter assignment

The OPLS3 forcefield [25] version that shipped with Schrödinger Suite release 2016-4 was used to parameterize the protein and ligand. Torsion parameter coverage was checked for all ligand fragments using Force Field Builder. The two ligands that contained a fragment with a torsion parameter not covered by OPLS3 were axitinib and bosutinib; Force Field Builder was used to obtain these parameters. SPC parameters [70] were used for water. For mutations that change the net change of the system, counterions were included to neutralize the system with additional Na+ and Cl- ions added to achieve 0.15 M excess to mimic the solution conditions of the experimental assay.

⁴⁹⁹ Prime (MM-GBSA)

Prime was used to predict the geometry of mutant side chains and to calculate relative changes in free energy using MM-GBSA single-point estimates [39]. VSGB [71] was used as the implicit solvent model to calculate the solvation free energies for the four states (complex/wild-type, complex/mutant, apo protein/wild-type, and apo protein/mutant) and $\Delta\Delta G$ calculated using the thermodynamic cycle depicted in *Figure 1*b. Unlike FEP (see below), which simulates the horizontal legs of the thermodynamic cycle, MM-GBSA models the vertical legs by computing the interaction energy between the ligand and protein in both wild-type and mutant states, subtracting these to obtain the $\Delta\Delta G$ of mutation on the binding free energy.

⁵⁰⁷ Alchemical free energy perturbation calculations using FEP+

Alchemical free energy calculations were performed using the FEP+ tool in the Schrödinger Suite version 508 2016-4, which offers a fully automated workflow requiring only an input structure (wild-type complex) and 509 specification of the desired mutation. The default protocol was used throughout: It assigns protein and 510 ligand force field parameters (as above), generates a dual-topology [72] alchemical system for transforming 511 wild-type into mutant protein (whose initial structure is modeled using Prime), generates the solvent-leg 512 endpoints (wild-type and mutant apo protein), and constructs intermediate windows spanning wild-type 513 and mutant states. Simulations of the apo protein were setup by removing the ligand from the prepared 514 complex (see System Preparation) followed by an identical simulation protocol as that used for the complex. 515 Charge-conserving mutations utilized 12 λ windows (24 systems) while charge-changing mutations utilized 24 516 λ windows (48 systems). Each system was solvated in an orthogonal box of explicit solvent (SPC water [70]) 517 with box size determined to ensure that solute atoms were no less than 5 Å (complex leg) or 10 Å (solvent leg) 518 from an edge of the box. For mutations that change the net charge of the system, counterions were included 519 to neutralize the charge of the system, and additional Na+ and Cl- ions added to achieve 0.15 M excess 520 NaCl to mimic the solution conditions of the experimental assay. The artifact in electrostatic interactions for 521 charge change perturbations due to periodic boundary conditions in MD simulations are corrected based on 522 the method proposed by Rocklin et al. [73]. 523

524 System equilibration was automated. It followed the default 5-stage Desmond protocol: (i) 100 ps with 525 1 fs time steps of Brownian dynamics with positional restraints of solute heavy atoms to their initial geometry

using a restraint force constant of 50 kcal/mol/Å²; this Brownian dynamics integrator corresponds to a 526 Langevin integrator in the limit when $\tau \rightarrow 0$, modified to stabilize equilibration of starting configurations 527 with high potential energies; particle and piston velocities were clipped so that particle displacements were 528 limited to 0.1 Å, in any direction, (ii) 12 ps MD simulations with 1 fs time step using Langevin thermostat at 529 10 K with constant volume, using the same restraints; (iii) 12 ps MD simulations with 1 fs time step using 530 Langevin thermostat and barostat [74] at 10 K and constant pressure of 1 atmosphere, using the same 531 restraints; (iv) 12 ps MD simulations with 1 fs time step using Langevin thermostat and barostat at 300 K 532 and constant pressure of 1 atmosphere, using the same restraints; (v) a final unrestrained equilibration 533 MD simulation of 240 ps with 2 fs time step using Langevin thermostat and barostat at 300 K and constant 534 pressure of 1 atmosphere. Electrostatic interactions were computed with particle-mesh Ewald (PME) [75] 535 and a 9 Å cutoff distance was used for van de Waals interactions. The production MD simulation was 536 performed in the NPT ensemble using the MTK method [76] with integration time steps of 4 fs. 4 fs. and 8 fs 537 respectively for the bonded, near, and far interactions following the RESPA method [77] through hydrogen 538 mass repartitioning [78]. Production FEP+ calculations utilized Hamiltonian replica exchange with solute 539 tempering (REST) [79], with automated definition of the REST region. Dynamics were performed with 540 constant pressure of 1 atmosphere and constant temperature of 300 K for 5 ns in which exchanges between 541 windows was attempted every 1.2 ps. 542

Because cycle closure could not be used to reduce statistical errors via path redundancy [79], we instead performed mutational free energy calculations in triplicate by initializing dynamics with different random seeds. The relative free energies for each mutation in each independent run were calculated using BAR [80, 81] The reported $\Delta\Delta G$ was computed as the mean of the computed $\Delta\Delta G$ from three independent simulations. Triplicate simulations were performed in parallel using four NIVIDA Pascal Architecture GPUs per alchemical free-energy simulation (12 GPUs in total), requiring ~6 hours in total to compute $\Delta\Delta G$.

⁵⁴⁹ Obtaining $\Delta\Delta G$ from ΔpIC_{50} benchmark set data

⁵⁵⁰ Reference relative free energies were obtained from three publicly available sources of ΔplC_{50} data (*Table 1*).

551 Under the assumption of Michaelis-Menten binding kinetics (pseudo first-order, but relative free energies are

⁵⁵² likely consistent), the inhibitor is competitive with ATP (*Equation 1*). This assumption has been successfully

 $_{\rm 553}$ $\,$ used to estimate relative free energies [37, 82–84] using the relationship between IC $_{\rm 50}$ and competitive

554 inhibitor affinity K_i ,

$$IC_{50} = \frac{K_i}{1 + \frac{[S_0]}{K_{12}}}.$$
(1)

If the Michaelis constant for ATP (K_M) is much larger than the initial ATP concentration S_0 , the relation in **Equation 1** will tend towards the equality $IC_{50} = K_i$. The relative change in binding free energy of AbI:TKI binding due to protein mutation is simply,

$$\Delta \Delta G = -RT \ln \frac{\mathrm{IC}_{50,WT}}{\mathrm{IC}_{50,mut}}$$
⁽²⁾

where $IC_{50,WT}$ is the IC_{50} value for the TKI binding to the wild-type protein and $IC_{50,mut}$ is the IC_{50} value for the mutant protein. *R* is the ideal gas constant and *T* is taken to be room temperature (300 K). As alluded to above, relating ΔpIC_{50} s to $\Delta \Delta Gs$ assumes that the Michaelis constant for ATP is much larger than the initial concentration of ATP, and that the experimentally observed ΔpIC_{50} change is solely from

changes in kinase:TKI binding affinity. In practice, not all of these assumptions may hold. For example, the experimentally observed ΔpIC_{50} might depend on the metabolism of drugs, and for drugs with different mechanisms of action than directly binding to the kinase binding pocket (e.g., binding to the transition structures of kinases, target gene amplification, up-/down-regulation of positive-/negative-feedback effectors, diminished synergism of pro-apoptotic machinery, decoupling of the target from cell survival circuits) [85, 86], their inhibition ability might not correlate well with binding affinity. However, the comparison between

 $_{568}$ ΔplC_{50} and ΔK_p is presented in *Figure 2*d, and this comparison indicates the assumptions we used to relate

 $_{569}$ ΔpIC_{50} to $\Delta \Delta G$ are reasonable for the dataset we studied.

570 Assessing prediction performance

- 571 Quantitative accuracy metrics
- 572 Mean unsigned error (MUE) was calculated by taking the average absolute difference between predicted and
- 573 experimental estimates of ΔΔG. Root-mean square error (RMSE) was calculated by taking the square root
- of the average squared difference between predicted and experimental estimates of $\Delta\Delta G$. MUE depends
- ⁵⁷⁵ linearly on errors such that large and small errors contribute equally to the average value, while RMSE
- 576 depends quadratically on errors, magnifying their effect on the average value.

577 Truth tables

- Two-class truth tables were constructed to characterize the ability of Prime and FEP+ to correctly classify mutations as susceptible ($\Delta\Delta G \le 1.36$ kcal/mol) or resistant ($\Delta\Delta G > 1.36$ kcal/mol), where the 1.36 kcal/mol threshold represents a 10-fold change in affinity. Accuracy was calculated as the fraction of all predictions that were correctly classified as sensitizing, neutral, or resistant. Sensitivity and specificity were calculated using a binary classification of resistant ($\Delta\Delta G > 1.36$ kcal/mol) or susceptible ($\Delta\Delta G \le 1.36$ kcal/mol). Specificity was calculated as the fraction of correctly predicted non-resistant mutations out of all truly susceptible mutations **S**. Sensitivity was calculated as the fraction of correctly predicted resistant mutations out of all
- $_{ss}$ truly resistant mutations, **R**. The number of susceptible mutations was 113 for axitinib, bosutinib, dasatinib,
- imatinib. nilotinib and ponatinib. and 12 for erlotinib and gefitinib; the number of resistant mutations **R** was
- ⁵⁸⁷ 18 for axitinib, bosutinib, dasatinib, imatinib, nilotinib, and ponatinib, and 1 for erlotinib and gefitinib.

588 Consensus model

- ⁵⁸⁹ First, Prime and FEP+ (n=142) were scaled by minimizing their RMSE to experiment by optimizing slope using
- ⁵⁹⁰ linear regression. The resulting (minimum) RMSE was used in a subsequent step to combine the scaled FEP+
- and scaled Prime free energies with inverse-variance weighted averaging.

592 ROC

⁵⁹³ A ROC curve was generated by computing the true positive rate (sensitivity) and the true negative rate ⁵⁹⁴ (specificity) when the classification cutoff differentiating resistant from sensitizing mutations is changed for ⁵⁹⁵ (only) the predicted values of $\Delta\Delta G$. Cutoffs were chosen by taking the minimum and maximum value of $\Delta\Delta G$ ⁵⁹⁶ for a data set (Prime or FEP+), and iteratively computing specificity and sensitivity in steps of 0.001 kcal/mol, ⁵⁹⁷ which by this definition will be in the range [0,1]. Experimental positives and negatives were classified with ⁵⁹⁸ the 1.36 kcal/mol cutoff. ROC-AUC was computed using the trapezoidal rule.

⁵⁹⁹ Estimating uncertainties of physical-modeling results

⁶⁰⁰ 95% symmetric confidence intervals (Cl, 95%) for all performance metrics were calculated using bootstrap by

resampling all datasets with replacement, with 1000 resampling events. Confidence intervals were estimated

for all performance metrics and reported as $x_{x_{low}}^{x_{high}}$ where x is the mean statistic calculated from the complete

⁶⁰³ dataset (e.g. RMSE), and x_{low} and x_{high} are the values of the statistic at the 2.5th and 97.5th percentiles of the

value-sorted list of the bootstrap samples. Uncertainty for $\Delta\Delta$ Gs was computed by the standard deviation

⁶⁰⁵ between three independent runs (using different random seeds to set initial velocities), where the 95% CI

was $[\Delta\Delta G - 1.96 \times \sigma_{FEP+}, \Delta\Delta G + 1.96 \times \sigma_{FEP+}]$ kcal/mol. 1 σ used in plots for FEP+ and experiment; 0 σ for Prime.

⁶⁰⁷ Bayesian hierarchical model to estimate intrinsic error

⁶⁰⁸ We used Bayesian inference to estimate the true underlying prediction error of Prime and FEP+ by making

⁶⁰⁹ use of known properties of the experimental variability (characterized in *Figure 2*) and statistical uncertainty

estimates generated by our calculations under weak assumptions about the character of the error.

⁶¹¹ We presume the true free energy differences of mutation *i*, $\Delta\Delta G_i^{\text{true}}$, comes from a normal background ⁶¹² distribution of unknown mean and variance,

$$\Delta \Delta G_i^{\text{true}} \sim \mathcal{N}(\mu_{\text{mut}}, \sigma_{\text{mut}}^2) \ i = 1, \dots, M$$
(3)

 $_{613}$ where there are M mutations in our dataset. We assign weak priors to the mean and variance

$$\mu_{\rm mut} \sim U(-6,+6) \tag{4}$$

$$\sigma_{\rm mut} \propto 1$$
 (5)

614 where we limit $\sigma > 0$.

⁶¹⁵ We presume the true computational predictions (absent statistical error) differ from the (unknown) ⁶¹⁶ true free energy difference of mutation $\Delta\Delta G_i^{true}$ by normally-distributed errors with zero bias but standard ⁶¹⁷ deviation equal to the RMSE for either Prime or FEP+, the quantity we are focused on estimating:

$$\Delta \Delta G_{i,\text{Prime}}^{\text{true}} \sim \mathcal{N}(\Delta \Delta G_i^{\text{true}}, \text{RMSE}_{\text{Prime}}^2)$$
(6)

$$\Delta \Delta G_{i,\text{FEP}+}^{\text{true}} \sim \mathcal{N}(\Delta \Delta G_{i}^{\text{true}}, \text{RMSE}_{\text{FEP}+}^{2})$$
(7)

In the case of Prime, since the computation is deterministic, we actually calculate $\Delta\Delta G_{Prime}^{true}$ for each mutant. For FEP+, however, the computed free energy changes are corrupted by statistical error, which we also presume to be normally distributed with standard deviation $\sigma_{calculate}$

$$\Delta \Delta G_{i,\text{FEP+}} \sim \mathcal{N}(\Delta \Delta G_{i,\text{FEP+}}, \sigma_{i,\text{FEP+}}^2)$$
(8)

where $\Delta\Delta G_{i,\text{FEP+}}$ is the free energy computed for mutant *i* by FEP+, and $\sigma_{i,\text{FEP+}}$ is the corresponding statistical error estimate.

The experimental data we observe is also corrupted by error, which we presume to be normally distributed with standard deviation σ_{exp} :

$$\Delta \Delta G_{i,\exp} \sim \mathcal{N}(\Delta \Delta G_i, \sigma_{\exp}^2)$$
(9)

⁶²⁵ Here, we used an estimate of K_d - and IC₅₀-derived $\Delta\Delta G$ variation derived from the empirical RMSE of 0.81 ⁶²⁶ kcal/mol, where we took $\sigma_{exp} \approx 0.81/\sqrt{2} = 0.57$ kcal/mol to ensure the difference between two random ⁶²⁷ measurements of the same mutant would have an empirical RMSE of 0.81 kcal/mol.

Under the assumption that the true $\Delta\Delta G$ is normally distributed and the calculated value differs from

the true value via a normal error model, it can easily be shown that the MUE is related to the RMSE via

MUE =
$$\int dx_{\text{true}} p(x_{\text{true}}) \int dx_{\text{calc}} p(x_{\text{calc}} | x_{\text{true}}) | x_{\text{calc}} - x_{\text{true}} |$$
(10)

$$= \int dx_{\rm true} \frac{1}{\sqrt{2\pi\sigma_{\rm true}^2}} e^{-\frac{(x_{\rm true}-\mu_{\rm true})^2}{2\sigma_{\rm true}^2}} \int dx_{\rm calc} \frac{1}{\sqrt{2\pi\sigma_{\rm calc}^2}} e^{-\frac{(x_{\rm calc}-\mu_{\rm true})^2}{2\sigma_{\rm calc}^2}} |x_{\rm calc} - x_{\rm true}|$$
(11)

$$= \sqrt{\frac{2}{\pi}} \text{RMSE}$$
(12)

The model was implemented using PyMC3 [87], observable quantities were set to their computed or experimental values, and 5000 samples drawn from the posterior (after discarding an initial 500 samples to burn-in) using the default NUTS sampler. Expectations and posterior predictive intervals were computed from the marginal distributions obtained from the resulting traces.

634 Data availability

Compiled experimental datasets, input files for Prime and FEP+ and computational results can be found at
 the following URL: https://goo.gl/6cC8Bu

637 Code availability

⁶³⁸ Scripts used for statistics analysis (including the Bayesian inference model) can be found at the following

639 URL: https://goo.gl/6cC8Bu

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- 647 **Disclosures**
- JDC is a member of the Scientific Advisory Board for Schrödinger Inc.
- **Author Contributions**
- ⁶⁵⁰ KH, JDC, CN, RA, and LW designed the research; KH, SA, TS, and LW identified experimental datasets; KH and
- LW performed the simulations; KH, CN, SKA, SR, TS, RA, JDC, and LW analyzed the data; KH, JDC, SKA, and LW
- ⁶⁵² wrote the paper.
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⁸⁹⁶ Supplementary Information

897 TITLE

- ⁸⁹⁸ Predicting resistance of clinical Abl mutations to targeted kinase inhibitors using alchemical free-energy
- 899 calculations

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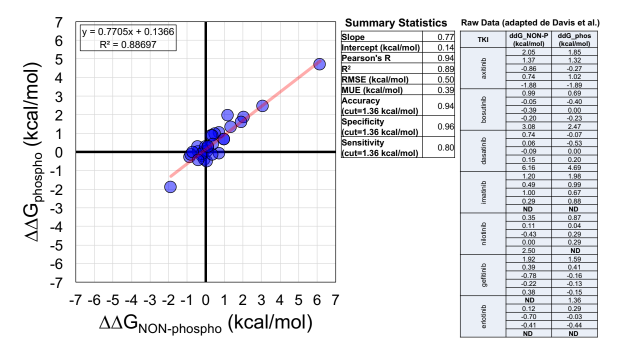


Figure S1. Comparison of 31 mutations for which phosphorylated and non-phosphorylated ΔK_d s were available. Scatter plot compares $\Delta\Delta$ Gs (derived from the ΔK_d s) and contains the best-fit line with slope 0.77 and intercept 0.14. Summary statistics for this comparison are also shown. The raw $\Delta\Delta$ Gs used for this comparison were adapted from [46]; kino-bead data for ponatinib was not available.

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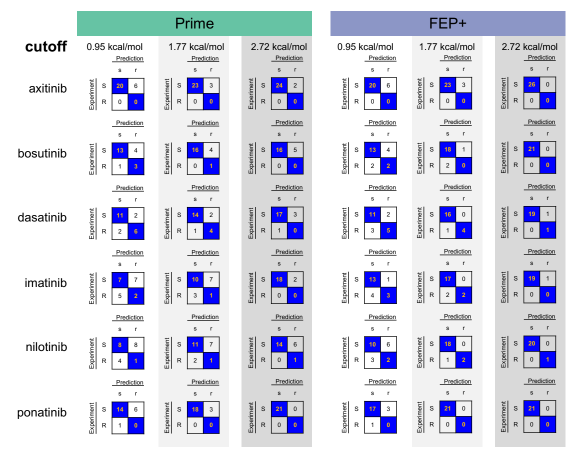


Figure S2. TKI-by-TKI truth tables with increasingly large classification cutoffs. Truth tables for the six TKIs (axitinib, bosutinib, dasatinib, imatinib, nilotinib, and ponatinib) using Prime (left, green) and FEP+ (right, blue) with classification cutoff values defining whether mutations are susceptible (S, experiment; s, prediction) or resistant (R, experiment; r, prediction). A mutation is susceptible if $\Delta\Delta G \leq$ cutoff or resistant if $\Delta\Delta G >$ cutoff.

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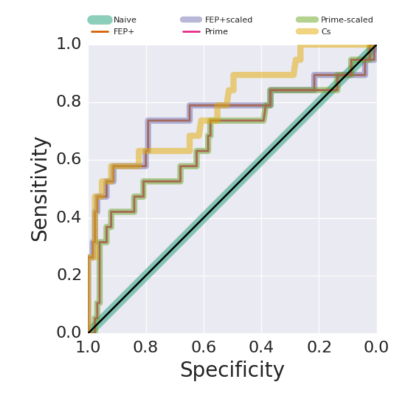


Figure S3. ROC curves for non-scaled and scaled FEP+, **non-scaled and scaled Prime**, **a consensus model and a naïve model**. ROC-AUC for scaled and non-scaled FEP+ was $0.75_{0.61}^{0.90}$ (n=144); ROC-AUC for scaled and non-scaled Prime was $0.66_{0.52}^{0.81}$ (n=144); ROC-AUCs for the naïve model and consensus model were $0.50_{0.50}^{0.50}$ (n=144) and $0.78_{0.67}^{0.00}$ (n=144) respectively. Optimal scaling factors (a=0.34 for FEP+; a=0.23 for Prime) obtained using linear regression (m=142) were applied to the full dataset (n=144), which was used in this ROC analysis. ROC-AUC interpretations: [0.50,0.60], failure; [0.60,0.70], poor; [0.70,0.80], fair; [0.80,0.90], good; [0.90,1.00], excellent.

Mutation	$\Delta\Delta G$	$\Delta\Delta G$	dasatinib ΔΔG	$\Delta\Delta G$	$\Delta\Delta G$	ponatinib ∆∆G (kcal/mol)	ΔΔG	$\Delta\Delta G$	Clinical-Observation
M244V	-0.11	0.43	0.00	0.21	-0.13	0.00	nd	nd	A
L248R	0.31	1.50	0.65	2.33	2.15	0.58	nd	nd	B
L248V	0.32	0.56	0.55	0.64	0.33	0.17	nd	nd	A,C
G250E	0.27	0.11	0.41	1.01	0.60	0.30	nd	nd	A,C,D
Q252H	0.20	nd	nd	nd	nd	nd	-0.44	-0.13	A
Y253F	0.26	-0.34	0.24	1.90	1.48	0.30	-0.17	0.00	С
Y253H	0.03	nd	nd	nd	nd	nd	nd	nd	A,C,D
E255K	0.26	0.56	0.90	1.50	1.27	0.41	-0.11	-0.11	A,C,D
E255V	0.30	0.66	1.02	2.22	2.36	1.00	nd	nd	A,C
D276G	0.18	nd	nd	nd	nd	nd	nd	nd	C
E279K	-0.03	nd	nd	nd	nd	nd	nd	nd	С
E292L	0.03	nd	nd	nd	nd	nd	nd	nd	E
V299L	-0.88	1.70	1.24	0.23	0.28	0.17	nd	nd	С
T315A	-0.45	0.32	2.02	0.51	0.72	0.17	nd	nd	С
T315I	-1.27	2.45	5.08	2.32	3.75	0.41	nd	-0.15	C,D
T315V	-1.73	nd	nd	nd	nd	nd	nd	nd	В
F317C	nd	0.50	1.86	0.28	0.04	0.00	nd	nd	A ^g
F317I	nd	0.71	1.79	0.17	0.30	0.51	1.35	1.58	С
F317L	0.23	0.09	0.96	0.72	0.20	0.17	0.29	0.40	C,D
F317R	0.27	nd	nd	nd	nd	nd	nd	nd	В
F317V	0.28	1.72	2.36	0.97	0.33	0.72	nd	nd	С
M343T	0.21	nd	nd	nd	nd	nd	nd	nd	F^h
M351T	-0.24	0.19	0.00	0.42	0.00	0.17	0.05	-0.08	A,C,D
E355A	nd	0.02	0.24	0.47	0.11	0.51	nd	nd	C
F359C	nd	-0.01	0.00	0.77	0.68	0.41	nd	nd	С
F359I	0.10	0.04	0.24	0.28	0.86	0.77	nd	nd	A
F359V	0.07	-0.11	0.00	0.32	0.60	0.17	nd	nd	A,C
L384M	0.06	nd	nd	nd	nd	nd	nd	nd	F ⁱ
H396R	0.25	-0.10	0.00	0.40	0.25	0.17	nd	nd	A ^j
F486S	0.05	nd	nd	nd	nd	nd	nd	nd	A ^k
E459K	nd	0.35	0.41	0.66	0.55	0.30	nd	nd	C

Table S1. $\Delta\Delta G$ data derived from publicly available ΔplC_{50} measurements and sources of mutation clinical-observation

A: Gruber et al. ([88])

B: Redaelli et al. ([89])

C: Cortes et al. ([90])

D: Branford et al. ([91])

E: Press et al. ([92])

F: Shah et al. ([3])

 g : F317C observed with Δ 27-183

^{*h*}: M343T observed as compound mutation with H396R

^{*i*}: L384M observed as compound mutation with M343T

^j: H396R observed as compound mutation with F486S

^k: F486S observed as compound mutation with H396R

	Expt. IC ₅₀	Expt. ΔΔG	Prime ΔΔG	FEP+ _{Run1} ΔΔG	FEP+ _{Run1} BAR err	FEP+ _{Run2} ΔΔG	FEP+ _{Run2} BAR err	$FEP+_{Run3} \Delta\Delta G$	FEP+ _{Run3} BAR err	$\Delta\Delta G_{Av}$	SE
	50			(kcal/mol)						110	
wild-type	823										
M244V	690	-0.11	-0.10	-0.40	0.41	-0.35	0.41	-0.43	0.41	-0.39	0.02
L248R	1393	0.31	-0.06	2.13	0.43	2.42	0.45	2.46	0.43	2.34	0.10
L248V	1399	0.32	6.02	-1.32	0.41	-1.04	0.42	-1.22	0.42	-1.19	0.08
G250E	1295	0.27	0.31	-0.35	0.41	-0.71	0.41	-0.74	0.41	-0.60	0.13
Q252H	1155	0.20	-0.18	0.07	0.43	0.30	0.42	0.29	0.43	0.22	0.08
Y253F	1275	0.26	1.11	0.77	0.43	0.23	0.43	1.15	0.45	0.72	0.27
Y253H	867	0.03	4.65	1.14	0.47	0.38	0.49	-0.19	0.45	0.44	0.39
E255K	1282	0.26	0.12	1.30	0.44	0.63	0.43	1.10	0.44	1.01	0.20
E255V	1350	0.30	-0.29	0.98	0.42	1.04	0.42	1.26	0.43	1.09	0.09
D276G	1105	0.18	-0.01	0.03	0.42	0.64	0.42	0.44	0.43	0.37	0.18
E279K	778	-0.03	-0.15	0.06	0.42	-0.22	0.43	1.27	0.43	0.37	0.46
E292L	863	0.03	-0.00	0.53	0.43	0.35	0.42	0.31	0.42	0.40	0.07
V299L	188	-0.88	-5.00	-1.08	0.42	-1.39	0.42	-1.37	0.42	-1.28	0.10
T315A	389	-0.45	0.99	0.09	0.43	0.24	0.47	0.31	0.42	0.21	0.06
T315I	98	-1.27	-2.30	-1.26	0.42	-1.50	0.45	-1.39	0.43	-1.38	0.07
T315V	45	-1.73	-1.07	-1.10	0.41	-1.32	0.42	-1.15	0.48	-1.19	0.07
F317L	1220	0.23	1.29	-0.64	0.41	-0.10	0.41	-0.38	0.41	-0.37	0.16
F317R	1286	0.27	-2.46	2.64	0.46	2.27	0.51	1.38	0.47	2.10	0.37
F317V	1320	0.28	2.29	0.45	0.42	0.70	0.42	0.75	0.42	0.63	0.09
M343T	1175	0.21	-0.04	-0.26	0.54	-0.50	0.53	-0.58	0.50	-0.45	0.10
M351T	553	-0.24	-0.07	-0.25	0.41	-0.03	0.41	0.37	0.41	0.03	0.18
F359I	975	0.10	-0.04	1.89	0.41	1.60	0.42	1.78	0.41	1.76	0.08
F359V	933	0.07	-0.07	2.68	0.42	1.55	0.42	1.64	0.41	1.96	0.36
L384M	916	0.06	-0.01	-0.07	0.41	0.27	0.41	0.23	0.41	0.14	0.11
H396R	1247	0.25	-0.02	0.36	0.42	1.23	0.41	0.65	0.42	0.75	0.26
F486S	897	0.05	-0.09	0.65	0.47	1.14	0.46	0.44	0.48	0.74	0.21

Table S2. Axitinib: experimental IC $_{50}$ values and alchemical free-energy $\Delta\Delta Gs$ for each mutation.

BAR err: Bennett Acceptance Ratio error.

 $\Delta \Delta \mathbf{G}_{Av}$: Average of three independent FEP+ runs.

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Table S3. Bosutinib: experimental IC $_{50}$ values and alchemical free-energy $\Delta\Delta Gs$ for each mutation.

	_	_									
	Expt.	Expt.	Prime	FEP+ _{Run1}	FEP+ _{Run1}	FEP+ _{Run2}	FEP+ _{Run2}	FEP+ _{Run3}	FEP+ _{Run3}		
	IC ₅₀	ΔΔG	ΔΔG	ΔΔG	BAR err	ΔΔG	BAR err	ΔΔG	BAR err	$\Delta\Delta G_{Av}$	SE
	(nM)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)
wild-type	71										
M244V	147	0.43	0.02	-0.28	0.41	-0.11	0.41	-0.08	0.41	-0.16	0.06
L248R	874	1.50	3.67	1.00	0.43	1.63	0.43	1.33	0.43	1.32	0.18
L248V	182	0.56	5.77	0.37	0.41	0.72	0.42	0.38	0.42	0.49	0.12
G250E	85	0.11	-0.30	0.28	0.43	0.63	0.43	-1.07	0.43	-0.05	0.52
Y253F	40	-0.34	-0.03	0.21	0.45	0.02	0.43	0.95	0.43	0.39	0.28
E255K	181	0.56	0.49	-1.01	0.43	-1.30	0.43	-1.01	0.43	-1.11	0.10
E255V	214	0.66	0.11	-0.47	0.42	-0.51	0.43	-0.91	0.43	-0.63	0.14
V299L	1228	1.70	-0.85	0.97	0.43	0.90	0.42	0.85	0.42	0.91	0.03
T315A	122	0.32	1.00	-1.61	0.41	-1.61	0.41	-1.97	0.41	-1.73	0.12
T315I	4338	2.45	3.75	-2.32	0.43	-2.21	0.42	-1.26	0.42	-1.93	0.34
F317C	165	0.50	4.83	1.04	0.41	1.27	0.41	1.22	0.42	1.18	0.07
F317I	232	0.71	1.61	0.16	0.41	0.07	0.42	0.02	0.41	0.08	0.04
F317L	82	0.09	-0.71	0.05	0.41	0.47	0.41	0.24	0.41	0.25	0.12
F317V	1280	1.72	4.12	1.98	0.42	1.50	0.42	2.25	0.42	1.91	0.22
M351T	97	0.19	0.02	0.36	0.42	0.82	0.41	0.71	0.41	0.63	0.14
E355A	74	0.02	0.13	-0.20	0.44	0.13	0.43	0.27	0.43	0.07	0.14
F359C	70	-0.01	-0.09	3.02	0.42	2.51	0.42	1.97	0.43	2.50	0.30
F359I	76	0.04	-0.06	0.66	0.41	1.74	0.41	1.43	0.42	1.28	0.32
F359V	59	-0.11	-0.06	0.98	0.43	1.69	0.41	1.91	0.42	1.53	0.28
H396R	60	-0.10	-1.07	0.62	0.42	-0.07	0.42	-0.93	0.43	-0.13	0.45
E459K	127	0.35	0.26	-0.69	0.42	0.23	0.42	-0.54	0.42	-0.33	0.28

BAR err: Bennett Acceptance Ratio error.

 $\Delta\Delta \mathbf{G}_{Av}$: Average of three independent FEP+ runs.

	Expt.	Expt.	Prime	FEP+ _{Run1}	FEP+ _{Run1}	FEP+ _{Run2}	FEP+ _{Run2}	FEP+ _{Run3}	FEP+ _{Run3}		
	IC_{50}	$\Delta\Delta G$	$\Delta\Delta G$	$\Delta\Delta G$	BAR err	$\Delta\Delta G$	BAR err	$\Delta\Delta G$	BAR err	$\Delta\Delta G_{Av}$	SE
	(nM)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)
wild-type	2										
M244V	2	0.00	-0.10	0.05	0.41	-0.37	0.41	-0.43	0.41	-0.25	0.15
L248R	6	0.65	-2.13	1.40	0.42	1.50	0.43	1.51	0.42	1.47	0.04
L248V	5	0.55	2.60	0.58	0.42	0.70	0.41	0.79	0.41	0.69	0.06
G250E	4	0.41	-0.00	-0.54	0.43	-0.31	0.43	0.01	0.44	-0.28	0.16
Y253F	3	0.24	0.00	-0.21	0.43	-0.24	0.43	-0.03	0.44	-0.16	0.07
E255K	9	0.90	-0.08	-0.30	0.43	-0.17	0.44	-1.05	0.43	-0.51	0.27
E255V	11	1.02	-0.08	0.06	0.42	-0.80	0.42	-0.12	0.42	-0.29	0.26
V299L	16	1.24	0.01	0.83	0.41	0.36	0.42	0.77	0.42	0.65	0.15
T315A	59	2.02	5.09	-1.74	0.41	-1.65	0.41	-1.23	0.41	-1.54	0.16
T315I	10000	5.08	-2.69	5.63	0.43	4.69	0.44	5.50	0.43	5.27	0.29
F317C	45	1.86	4.72	2.63	0.42	2.32	0.42	2.62	0.41	2.52	0.10
F317I	40	1.79	2.38	1.94	0.41	2.04	0.41	1.94	0.41	1.97	0.03
F317L	10	0.96	1.22	1.26	0.41	1.42	0.41	1.08	0.41	1.25	0.10
F317V	104	2.36	4.08	3.12	0.42	2.84	0.42	2.68	0.42	2.88	0.13
M351T	2	0.00	0.04	0.04	0.41	0.14	0.41	0.00	0.42	0.06	0.04
E355A	3	0.24	0.00	-0.24	0.43	-0.87	0.45	-1.25	0.44	-0.79	0.29
F359C	2	0.00	-0.03	1.24	0.42	0.68	0.41	1.38	0.42	1.10	0.21
F359I	3	0.24	-0.02	-0.50	0.42	-0.33	0.42	-1.14	0.42	-0.66	0.25
F359V	2	0.00	-0.03	-0.87	0.41	0.57	0.42	-0.62	0.41	-0.31	0.44
H396R	2	0.00	2.53	-0.76	0.43	-0.09	0.43	-0.06	0.43	-0.30	0.23
E459K	4	0.41	0.00	-0.68	0.42	-0.17	0.42	-0.07	0.41	-0.31	0.19

Table S4. Dasatinib: experimental IC $_{50}$ values and alchemical free-energy $\Delta\Delta Gs$ for each mutation.

T315I was beyond the concentration limit of the assay (10,000 nM).

BAR err: Bennett Acceptance Ratio error.

 $\Delta \Delta \mathbf{G}_{Av}$: Average of three independent FEP+ runs.

	Expt. IC ₅₀	Expt. ΔΔG	Prime ΔΔG	$FEP+_{Run1} \Delta \Delta G$	FEP+ _{Run1} BAR err	$FEP+_{Run2} \Delta\Delta G$	FEP+ _{Run2} BAR err	$FEP+_{Run3} \Delta\Delta G$	FEP+ _{Run3} BAR err	$\Delta\Delta G_{Av}$	SE
	(nM)					(kcal/mol)				110	
wild-type	201										
M244V	287	0.21	-0.08	0.15	0.41	0.43	0.41	0.17	0.41	0.25	0.09
L248R	10000	2.33	1.92	1.92	0.43	2.52	0.44	2.34	0.43	2.26	0.18
L248V	586	0.64	1.89	-1.04	0.41	-1.02	0.42	-1.20	0.41	-1.09	0.06
G250E	1087	1.01	0.92	0.16	0.41	0.02	0.41	0.12	0.41	0.10	0.04
Y253F	4908	1.90	-0.02	0.87	0.43	0.65	0.42	1.34	0.44	0.95	0.20
E255K	2487	1.50	0.25	-0.12	0.44	1.95	0.44	-0.55	0.44	0.43	0.77
E255V	8322	2.22	0.24	-0.72	0.42	-0.02	0.42	-0.53	0.43	-0.42	0.21
V299L	295	0.23	-1.29	0.66	0.41	0.26	0.42	-0.37	0.42	0.18	0.30
T315A	476	0.51	5.10	-1.39	0.41	-1.86	0.41	-2.09	0.44	-1.78	0.21
T315I	9773	2.32	0.88	4.23	0.43	4.23	0.42	3.14	0.44	3.87	0.36
F317C	324	0.28	2.10	0.27	0.42	-0.18	0.41	0.45	0.42	0.18	0.19
F317I	266	0.17	0.94	0.59	0.41	0.66	0.41	0.48	0.41	0.58	0.05
F317L	675	0.72	0.74	0.58	0.41	0.53	0.41	0.38	0.41	0.50	0.06
F317V	1023	0.97	1.57	0.71	0.42	0.79	0.42	0.80	0.41	0.77	0.03
M351T	404	0.42	-0.02	1.72	0.41	1.03	0.42	1.20	0.42	1.32	0.21
E355A	441	0.47	0.29	0.13	0.43	0.08	0.44	0.14	0.43	0.12	0.02
F359C	728	0.77	2.43	0.88	0.42	0.47	0.41	0.33	0.42	0.56	0.17
F359I	324	0.28	1.95	-0.13	0.41	-0.87	0.41	0.08	0.41	-0.31	0.29
F359V	346	0.32	2.53	-0.66	0.41	0.02	0.41	-0.27	0.42	-0.30	0.20
H396R	395	0.40	2.76	-0.39	0.41	-0.38	0.42	-0.39	0.42	-0.39	0.00
E459K	612	0.66	0.24	-0.09	0.43	-0.09	0.42	-0.08	0.42	-0.09	0.00

Table S5. Imatinib: experimental IC $_{50}$ values and alchemical free-energy $\Delta\Delta$ Gs for each mutation.

T315I was beyond the concentration limit of the assay (10,000 nM).

BAR err: Bennett Acceptance Ratio error.

 $\Delta \Delta \mathbf{G}_{Av}$: Average of three independent FEP+ runs.

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	E.unt	E. un f	During a								
	Expt.	Expt.	Prime	FEP+ _{Run1}	FEP+ _{Run1}	FEP+ _{Run2}	FEP+ _{Run2}	FEP+ _{Run3}	FEP+ _{Run3}	A A C	сг
	IC ₅₀	$\Delta\Delta G$	$\Delta\Delta G$	$\Delta\Delta G$	BAR err	$\Delta\Delta G$	BAR err	$\Delta\Delta G$	BAR err	$\Delta\Delta G_{Av}$	SE
	(nM)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)
wild-type	15										
M244V	12	-0.13	-0.11	0.15	0.41	-0.21	0.41	0.21	0.41	0.05	0.13
L248R	549	2.15	0.48	2.05	0.43	2.12	0.47	1.93	0.43	2.03	0.06
L248V	26	0.33	3.53	-0.50	0.42	-0.39	0.41	-0.92	0.41	-0.60	0.16
G250E	41	0.60	0.05	0.06	0.41	-0.27	0.41	-0.38	0.41	-0.20	0.13
Y253F	179	1.48	-0.27	1.09	0.43	0.42	0.42	1.16	0.42	0.89	0.24
E255K	127	1.27	0.41	-2.24	0.48	-1.52	0.46	0.33	0.46	-1.14	0.77
E255V	784	2.36	-0.03	0.31	0.42	-0.25	0.43	-0.55	0.43	-0.16	0.25
V299L	24	0.28	2.94	-0.18	0.41	0.21	0.41	0.15	0.41	0.06	0.12
T315A	50	0.72	3.38	-1.33	0.41	-1.31	0.41	-1.39	0.41	-1.34	0.02
T315I	8091	3.75	4.16	4.29	0.43	5.00	0.42	4.34	0.43	4.54	0.23
F317C	16	0.04	0.90	1.34	0.41	0.88	0.41	0.60	0.41	0.94	0.22
F317I	25	0.30	-0.18	1.24	0.41	1.17	0.41	0.82	0.41	1.08	0.13
F317L	21	0.20	1.74	1.03	0.41	1.07	0.41	1.09	0.41	1.06	0.02
F317V	26	0.33	0.77	1.16	0.41	0.68	0.42	1.07	0.42	0.97	0.15
M351T	15	0.00	0.09	-0.06	0.41	-0.09	0.42	-0.46	0.42	-0.20	0.13
E355A	18	0.11	-0.06	-0.46	0.43	-1.01	0.43	-0.32	0.43	-0.60	0.21
F359C	47	0.68	3.68	1.32	0.41	1.44	0.41	1.52	0.41	1.43	0.06
F359I	64	0.86	3.70	1.05	0.41	1.13	0.41	0.74	0.41	0.97	0.12
F359V	41	0.60	3.67	1.00	0.41	1.08	0.41	1.38	0.42	1.15	0.12
H396R	23	0.25	2.58	-0.07	0.42	0.21	0.42	0.03	0.42	0.06	0.08
E459K	38	0.55	-0.00	-0.17	0.42	-0.46	0.42	-0.10	0.42	-0.24	0.11

BAR err: Bennett Acceptance Ratio error.

 $\Delta\Delta \mathbf{G}_{Av}$: Average of three independent FEP+ runs.

		-									
	Expt.	Expt.	Prime	FEP+ _{Run1}	FEP+ _{Run1}	FEP+ _{Run2}	FEP+ _{Run2}	FEP+ _{Run3}	FEP+ _{Run3}		
	IC_{50}	ΔΔG	ΔΔG	ΔΔG	BAR err	ΔΔG	BAR err	ΔΔG	BAR err	$\Delta\Delta G_{Av}$	SE
	(nM)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)	(kcal/mol)
wild-type	3										
M244V	3	0.00	-0.13	0.07	0.41	-0.28	0.41	0.12	0.41	-0.03	0.13
L248R	8	0.58	2.48	1.40	0.43	0.96	0.43	1.10	0.44	1.15	0.13
L248V	4	0.17	2.48	-1.82	0.42	-1.23	0.42	-1.96	0.42	-1.67	0.22
G250E	0.021	0.30	0.17	-0.32	0.43	-0.25	0.43	-0.71	0.46	-0.43	0.14
Y253F	5	0.30	0.05	0.85	0.43	1.32	0.44	0.77	0.43	0.98	0.17
E255K	6	0.41	1.05	-0.27	0.48	-0.66	0.48	0.03	0.47	-0.30	0.20
E255V	16	1.00	-0.04	1.19	0.43	0.94	0.43	-0.41	0.43	0.57	0.50
V299L	4	0.17	-0.29	-0.56	0.41	-0.55	0.41	-1.42	0.41	-0.84	0.29
T315A	4	0.17	-0.51	-2.90	0.41	-3.15	0.41	-2.92	0.41	-2.99	0.08
T315I	6	0.41	-5.42	0.51	0.42	0.90	0.42	0.91	0.42	0.77	0.13
F317C	3	0.00	1.45	0.44	0.41	0.98	0.42	0.80	0.41	0.74	0.16
F317I	7	0.51	0.62	-0.76	0.41	-1.03	0.41	-1.02	0.41	-0.94	0.09
F317L	4	0.17	0.57	-1.08	0.41	-0.83	0.41	-0.85	0.41	-0.92	0.08
F317V	10	0.72	1.14	0.05	0.41	-0.21	0.41	0.24	0.42	0.03	0.13
M351T	4	0.17	-0.12	0.89	0.41	1.66	0.41	0.65	0.41	1.07	0.30
E355A	7	0.51	0.01	0.12	0.44	-0.52	0.44	-0.55	0.43	-0.32	0.22
F359C	6	0.41	2.12	0.25	0.42	-0.35	0.43	0.73	0.42	0.21	0.31
F359I	11	0.77	0.34	-0.66	0.41	-0.38	0.41	0.06	0.41	-0.33	0.21
F359V	4	0.17	0.74	0.11	0.41	-0.28	0.41	0.08	0.42	-0.03	0.13
H396R	4	0.17	-0.04	0.19	0.49	0.10	0.45	-1.41	0.48	-0.37	0.52
E459K	5	0.30	-0.00	-0.51	0.42	-0.78	0.42	-0.63	0.42	-0.64	0.08

Table S7. Ponatinib: experimental IC $_{50}$ values and alchemical free-energy $\Delta\Delta Gs$ for each mutation.

BAR err: Bennett Acceptance Ratio error.

 $\Delta \Delta \mathbf{G}_{Av}$: Average of three independent FEP+ runs.

Method	Scaling factor	MUE (kcal/mol)	RMSE (kcal/mol)	Accuracy	Specificity	Sensitivity
		[N=142]	[N=142]	[N=144]	[N=144]	[N=144]
Prime	1.00	$1.14_{0.94}^{1.35}$	$1.70^{1.97}_{1.40}$	$0.73_{0.65}^{0.80}$	$0.76_{0.68}^{0.83}$	$0.53_{0.29}^{0.78}$
Prime	0.50	$0.64_{0.53}^{0.76}$	$0.91^{1.06}_{0.77}$	$0.84^{0.90}_{0.78}$	$0.00^{0.95}$	$0.42_{0.20}^{0.65}$ 0.26 ^{0.47}
Prime	0.33	$0.53^{0.62}_{0.44}$	$0.76_{0.63}^{0.87}$	$0.87^{0.92}_{0.81}$	$0.90_{0.84}$ $0.96_{0.92}^{0.99}$	0.200.08
Prime	0.23	$0.49^{0.59}$	$0.73^{0.86}$	$0.86^{0.92}$	0.991.00	$0.00^{0.00}$
FEP+	1.00	$0.79_{0.40}^{0.40}$ $0.79_{0.67}^{0.91}$	$1.07_{0.89}^{1.27}$	$0.88_{0.81}^{0.93}$	$0.99_{0.97}^{0.97}$ $0.94_{0.89}^{0.98}$	$0.47^{0.72}_{0.22}$
FEP+	0.34	$0.55_{0.47}^{0.64}$	$0.78^{0.91}_{0.65}$	$0.88^{0.93}_{0.82}$	$1.00^{1.00}_{1.00}$	$0.11_{0.00}^{0.27}$
Naive	_	$0.57_{0.46}^{0.69}$	$0.87^{1.04}_{0.70}$	$0.87_{0.81}^{0.92}$	$1.00^{1.00}_{1.00}$	$0.00_{0.00}^{0.00}$
Consensus		$0.47_{0.39}^{0.56}$	$0.71_{0.59}^{0.84}$	$0.87_{0.81}^{0.92}$	$1.00_{1.00}^{1.00}$	$0.00_{0.00}^{0.00}$

Table S8. Summary of statistics of scaled predictions, a naïve model, and a consensus model.

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Table S9. Summary of the preparation of the 6 Abl:TKl co-crystal structure complexes.

Receptor Ligand Chains # Aminos Chains # Aminos E Rec. # Rec. # Ash # Lig # Lig <th></th> <th></th> <th></th> <th></th> <th>Experim</th> <th>Experimental structure</th> <th></th> <th></th> <th></th> <th>Prep</th> <th>ared mo</th> <th>Prepared model used for simulations</th> <th>or simul.</th> <th>ations</th> <th></th> <th></th>					Experim	Experimental structure				Prep	ared mo	Prepared model used for simulations	or simul.	ations		
Abl Axit A, B 305 2219 (B) 276 (B) B 131 4580 284 $Ash21$ 0 0 46 Abl Bosut A, B 305 2219 (B) 276 (B) B 131 4580 284 $Ash21$ 0 0 46 Abl Bosut A, B 152 2195 (A) 270 (A) A 89 4581 284 $Ash21$ 0 0 66 Abl Dasat A, B 0 2195 (A) 269 (A) A 0 4581 0 <th>PDB</th> <th>Receptor</th> <th>Ligand</th> <th>Chains</th> <th># Water^a</th> <th># Rec. atoms,</th> <th># Aminos,</th> <th>Chain</th> <th># Water</th> <th># Rec.</th> <th># Rec.</th> <th># Ash</th> <th># Glh</th> <th># Hip</th> <th># Lig.</th> <th>Het. atomd</th>	PDB	Receptor	Ligand	Chains	# Water ^a	# Rec. atoms,	# Aminos,	Chain	# Water	# Rec.	# Rec.	# Ash	# Glh	# Hip	# Lig.	Het. atom d
Abl Axit A, B 305 2219(B) 276(B) B 131 4580 284 Ash21 0 46 Abl Bosut A, B 152 2187(A) 270(A) A 89 4581 284 Ash21 0 0 66 Abl Bosut A, B 0 2195(A) 269(A) A 89 4581 284 Ash311 0 0 66 Abl Dasat A, B 0 2195(A) 269(A) A 0 4581 0 0 0 66 Abl Imat A, B 2336(B) 269(A) A 0 4581 0 0 0 59 Abl Imat A, B 2336(B) 288(B) B 104 4579 284 Ash31 ^e 0 0 6 9 Abl Nilot A, B, C, D 266(A) A 99 4579 284 0 0						(Chain) ^b	(Chain)			atoms	aminos				atoms	w/ proton
Abl Bosut A,B 152 2187 (A) 270 (A) A 89 4581 284 Ash21 0 06 66 Abl Dasat A,B 0 2195 (A) 269 (A) A 0 4581 284 Ash21* 0 0 59 Abl Dasat A,B 0 2195 (A) 269 (A) A 0 4581 284 Ash31* 0 0 59 Abl Imat A,B 231 2336 (B) 288 (B) B 104 4579 284 0 0 0 59 Abl Nilot A,B,C,D 266 2142 (A) 264 (A) A 99 4579 284 0 0 0 61 Abl Ponat A 89 2152 (A) 268 (A) A 89 4580 284 0 0 0 0 67	4wa9	Abl	Axit	А, В	305	2219 (B)	276 (B)		131	4580	284	Ash421	0	0	46	neutral
Abl Dasat A,B 0 2195 (A) 269 (A) A 0 4581 Ash381 Abl Dasat A,B 0 2195 (A) 269 (A) A 0 4581 284 Ash421 ^e 0 0 59 Abl Imat A,B 231 2336 (B) 288 (B) B 104 4579 284 0 0 69 Abl Nilot A,B,C,D 266 2142 (A) 264 (A) A 99 4579 284 0 0 61 Abl Ponat A 89 4580 284 0 0 0 61	3ue4	Abl	Bosut		152	2187 (A)	270 (A)	A	89	4581	284		0	0	99	NBI,4401
Abl Dasat A, B 0 2195 (A) 269 (A) A 0 4581 284 A5H21 ^s 0 0 59 Abl Imat A, B 231 2336 (B) 288 (B) B 104 4579 284 0 69 69 Abl Nilot A, B, C, D 266 2142 (A) 264 (A) A 99 4579 284 0 0 61 Abl Ponat A 89 4580 284 0 0 0 61												Ash381				
Abl Imat A,B 231 2336 (B) 288 (B) B 104 4579 284 0 0 69 Abl Nilot A,B,C,D 266 2142 (A) 264 (A) A 99 4579 284 0 0 69 61 Abl Ponat A 89 4580 284 0 0 61	4xey	Abl	Dasat	A, B	0	2195 (A)	269 (A)	A	0	4581	284	Ash421 ^c	0	0	59	neutral
Imat A, B 231 2336 (B) 288 (B) B 104 4579 284 0 0 69 69 69 61 <td></td> <td>Ash381^c</td> <td></td> <td></td> <td></td> <td></td>												Ash381 ^c				
Abl Nilot A, B, C, D 266 2142 (A) 264 (A) A 99 4579 284 0 0 6 61 Abl Ponat A 89 4579 284 0 0 6 61	1opj		lmat		231	2336 (B)	288 (B)	В	104	4579	284	0	0	0	69	N51,4767
Abl Ponat A 89 2152(A) 268(A) A 89 4580 284 0 0 67	3cs9		Nilot			2142 (A)	264 (A)	A	66	4579	284	0	0	0	61	neutral
	30XZ	Abl	Ponat	A	89	2152 (A)	268 (A)	∢	89	4580	284	0	0	0	67	N3,2155

 b Count includes N-Acety//N-terminal (6 atoms) and N-methylamide/C-terminal (6 atoms) capping groups.

^c Original index in experimental structure was Ash440, Ash400. ^d (PDB atom name), (PDB serial).

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