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Limitations of ligand-only approaches for predicting the reactivity of covalent inhibitors

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Abstract: Covalent inhibition has undergone a resurgence and is an important modern-day drug design and chemical biology approach. To avoid off-target interactions, and to fine tune reactivity, the ability to accurately predict reactivity is vitally important for the design and development of safer and more effective covalent drugs. Several ligand-only metrics have been proposed that promise quick and simple ways of determining covalent reactivity. In particular, we examine proton affinity and reaction energies calculated with the density functional B3LYP-D3/6-311+G**//B3LYP-D3/6-31G* method to assess the reactivity of a series of α,β-unsaturated carbonyl compounds that form covalent adducts with cysteine. We demonstrate that, whilst these metrics correlate well with experiment for a diverse range of covalent fragments, these approaches fail for predicting the reactivity of drug-like compounds. We conclude that ligand-only metrics such as proton affinity and reaction energies do not capture determinants of reactivity in situ and fail to account for important factors such as conformation, solvation and intramolecular interactions.

Predicting covalent reactivity is an important goal in targeted covalent inhibitor (TCI) design. In contrast to conventional reversible inhibitors, TCIs form a covalent attachment with their target, resulting in several advantageous properties. These include increased potency, selectivity, and residence time all leading to distinct pharmacodynamic properties. The half-life of the covalent complex (which may be long, compared with the clearance of the free TCI and synthesis rate of the target protein) can offer pharmacokinetic advantages. Efficacy can be extended beyond what would be expected due to the half-life of the free drug alone in plasma. There are now several recently marketed covalent kinase inhibitors, for example (Figure 1). However, toxicity concerns are an important consideration when pursuing a covalent mechanism of action in drug design, owing to the intrinsic reactivity of TCIs. In addition, clearance and cross-reactivity/selectivity can be problematic. A typical approach is the addition of a reactive covalent warhead onto a potent reversible inhibitor of the target, in the expectation that selectivity will be improved by the optimized fit of the reversible molecule for the particular protein target. Covalent warheads can target various amino acids including cysteine, serine, tyrosine and lysine. Amongst these, acrylamides are often used to target cysteine residues; acrylamide reactivity can be modulated via the amine and β-carbon substituents of the motif.

As the field of TCIs has blossomed over recent years, some model/simplified experimental and computational approaches have been proposed to assist in predicting optimal reactivity. Experimental approaches have been used to estimate the general reactivity of covalent fragments with sulfur-containing species representing cysteine, typically glutathione (GSH). This is often reported as GSH $t_{1/2}$, the half-life of the reaction of a covalent compound with glutathione. The GSH trapping assessment is readily accessible within drug discovery due to its legacy as a model for reactive toxicity, particularly of metabolites. Therefore, such data is relatively easy to generate compared to protein binding reactivity or kinetics and serves as a convenient metric for comparing warhead reactivity with cysteine in TCI design.

![Figure 1. Recently approved acrylamide containing covalent kinase inhibitors and alternative covalent warheads. Ibrutinib targets BTK and is used as a treatment for multiple B cell cancers. Afatinib and osimertinib both target EGFR and are treatments for non-small cell lung carcinoma.](image-url)

With regard to computational chemistry methods, modeling the reaction in the protein-ligand complex with QM/MM methods has shown success but remains complex and relatively time-consuming for drug discovery. Instead, one approach has
been to reduce the covalent reaction between acrylamides and their protein target to a simple ligand-only reactivity metric. Ligand-only methods have shown some success in predicting reactivity trends for small covalent fragments.\textsuperscript{16-19} Examples of these methods include using calculated proton affinity (PA), reaction energies ($\Delta E_{\text{react}}$), kinetic barrier heights ($\Delta G^2$), electrophilicity index and NMR chemical shifts to predict covalent reactivity. Recent work from Houk \textit{et al.} elaborated the full reaction mechanism for a small set of $\alpha,\beta$-unsaturated carbonyl compounds illustrating that both kinetics and thermodynamics are important determinants of covalent reactivity of Michael acceptors with methyl thiolate.\textsuperscript{19} Complementary to this, Lonsdale \textit{et al} used density functional theory (DFT) to calculate reaction energies of a diverse set of covalent fragments. Whilst they set out calculating some QM transition state (TS) barriers, most of their work focused on calculating the energy difference between reactants and adduct equivalent to species 1 and 5 ($\Delta E_{\text{react}}$ in Figure 2). They found these to correlate well with GSH reactivity ($R^2=0.69$).\textsuperscript{17} Furthermore, Flanagan \textit{et al} found that calculated reaction barrier heights ($\Delta G^2$) correlate strongly with GSH reactivity ($R^2=0.92$).\textsuperscript{20} $\Delta G^2$ is the energy difference between species 1 and the transition state in Figure 2, and reflects the overall reactivity rate (GSH $t_{1/2}$) of the reaction. PA provides an estimation of the free energy difference between the covalent-thiol adduct and its corresponding conjugate base (4 and 5 respectively, $\Delta G_{\text{PA}}$ in Figure 2). Krishnan \textit{et al} found that PA values correlate well with $\beta$-elimination rates of covalent fragments from sulfur containing species, which vary based on the acidity of the $\alpha$-carbon adjacent to the carbonyl group of the acrylamide.\textsuperscript{16} More recently, the electrophilicity index, derived from QM calculations has been shown to correlate well with experimental data. However, the authors note the difficulty in reactivity prediction using this method for larger compounds and non-terminal acrylamides.\textsuperscript{21}

Proton affinities and reaction energies ($\Delta E_{\text{react}}$) are particularly attractive reactivity metrics, as their computational prediction is relatively straightforward from quantum mechanical (QM) calculations.\textsuperscript{22} PA and $\Delta E_{\text{react}}$ are reactivity metrics that provide insight into the thermodynamic stability of the reaction. Although the TS is not explicitly used, PA and $\Delta E_{\text{react}}$ calculations give a good approximation to overall experimental reactivity as the enolate intermediate 4 that results from thio-Michael addition between cysteine and an acrylamide (Figure 2) is close in energy and geometry to the TS.\textsuperscript{17} When considering an appropriate computational protocol to predict reactivity for drug molecules, the ability to consistently and reliably generate results for various input molecules is crucial because such an approach needs to be applicable to libraries of hundreds of analogues. The automated QM calculation of transition states presents a challenge because of the difficulty to reach geometrical convergence for saddle point structures whereas calculating PA and $\Delta E_{\text{react}}$ is more straightforward. However, new tools for TS searches are emerging and have been applied to thio-Michael reactivity, although the authors note the difficulty due to the typically flat nature of the potential energy surface (PES) around the TS.\textsuperscript{23} In addition, it is difficult to accurately represent solvation changes with continuum solvation models, particularly for reactions involving charged intermediates, although good results can be achieved with explicit treatment of solvent.\textsuperscript{24}

Our aim here is to investigate if two common ligand-only reactivity metrics, PA and $\Delta E_{\text{react}}$, can accurately and reliably be used to predict biologically relevant covalent reactivity. We use a large number of compounds ranging from small covalent fragments, to large drug-like molecules to demonstrate how the simple ligand-only reactivity trends based on PA and $\Delta E_{\text{react}}$ compare favorably with GSH reactivity for similar small fragments but break down for larger drug like molecules.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{A schematic view of thiol addition to a generic $\alpha,\beta$-unsaturated carbonyl compound. The energy profile highlights the chemical species commonly used in ligand-only reactivity metrics: reaction energy ($\Delta E_{\text{react}}$), reaction barrier height ($\Delta G^2$) and proton affinity ($\Delta G_{\text{PA}}$)}
\end{figure}

\textbf{Methods}

A previous study used QM calculations to calculate proton affinities with the B3LYP functional and the 6-311+G(d) basis set.\textsuperscript{16} Here we use a modified protocol, appropriate for proton affinity calculations of larger compounds. We performed gas phase geometry optimizations at the B3LYP-D3/6-31G(d) level of theory, followed by single point energies in solution using B3LYP-D3/6-311+G(d,p) level of theory. This has been shown to be sufficient for the calculation of proton affinity values for simple organic molecules of this type.\textsuperscript{16,25,26} The inclusion of dispersion corrections to B3LYP has been recommended in QM calculations,\textsuperscript{27} particularly when modelling cysteine reactivity.\textsuperscript{28} Although B3LYP has known limitations in modelling reactivity,\textsuperscript{29-32} it provides an acceptable balance of accuracy and speed in this context for predicting relative reactivities. We have also tested the M06-2X functional, but found it made no appreciable difference to the results (Supporting Information, Section 4). We included solvation by using the Poisson-Boltzmann Finite (PBF) element implicit water solvation model using a probe radius of 1.4 Å.\textsuperscript{33,34} which has been shown to reproduce experimental solvation energies for anionic species with good accuracy.\textsuperscript{35} All QM calculations were performed with Jaguar v8.5-13.\textsuperscript{36}
For larger, drug-like molecules, conformational sampling becomes important. An assessment of the variation of PA with respect to the conformation of each molecule was performed. Six representative molecules were chosen. A conformational search was performed on each using the Macromodel tool in Maestro v2018.3. The mixed torsional/low-mode sampling approach was used with default settings, using water as a solvent. An energy window of 5 kcal/mol was applied to discard higher energy conformations. This search resulted in over 2100 conformations for the 6 molecules combined. Two distinct conformations per molecule were extracted: the lowest energy conformation; and the next lowest energy with a root mean square deviation (RMSD) > 2.0 Å with respect to the first. This RMSD cutoff is typically used to distinguish structurally different conformations of drug-like molecules. These conformations were submitted to QM minimization at the B3LYP-D3/6-311+G(d,p) level of theory again using the PBF solvation model. All outputs were submitted to frequency calculations to confirm stationary points as minima, and to obtain zero-point energy for thermal and entropic corrections. These two minima were then used as inputs for the PA calculations as described above. PAs were calculated according to Figure 2. This involved calculating the difference in energy of the two states, 4 and 5 to obtain $\Delta G_{PA}$. These $\Delta G_{PA}$ values were then normalized relative to the most acidic compound within each data set, providing what we refer to as $\Delta G_{PA}^\ast$ (kcal mol$^{-1}$). Reaction energy ($\Delta E_{\text{react}}$) values were calculated by taking the difference in energy between states 1 and 5. Since we already had optimized structures for state 5, state 1 was generated simply by breaking the S-C bond to minimize towards the local reactant energy minimum. Geometry optimization was performed at the B3LYP-D3/6-31G(d) level in the gas phase, followed by single point energy calculations in aqueous solution at the B3LYP-D3/6-31G(d,p) level of theory.

![Graph](image-url)

**Figure 3.** Plot of GSH reactivity vs relative proton affinity ($\Delta G_{PA}$) for data set 1, $R^2=0.79$ turquoise squares, data set 2, $R^2=0.90$ magenta circles, data set 3, $R^2=0.75$ blue triangles. Representative structures from each data set are shown. There is a positive correlation observed between $\Delta G_{PA}$ and log $k_{GSH}$ for each data set. More acidic compounds (represented by a more positive $\Delta G_{PA}$ value) have faster experimental reactivity rates.

### Results

A literature search was performed to identify data sets that contained small molecule compounds bearing acrylamide functionality and associated GSH reactivity data. The fragments identified had a molecular weight range of 133 to 271 g mol$^{-1}$, and between 1 and 6 rotatable bonds. Using the method outlined above, we calculated proton affinity values for three data sets containing a total of 37 unique compounds. For each data set, a plot of the relative proton affinity ($\Delta G_{PA}$) against the GSH reactivity rate ($\log k$) reveals a strong correlation: data set 1: $R^2=0.79$; data set 2: $R^2=0.90$; and data set 3: $R^2=0.75$ (Figure 3). A more negative $\Delta G_{PA}$ value corresponds to a more acidic compound and as expected these acidic compounds also have a faster GSH reactivity rate.

To test our approach for larger, more conformationally complex compounds, we performed a literature search for data sets that contained `drug-like` compounds, bearing structural similarity to already approved covalent drugs on the market that also had published GSH reactivity data. The compounds had higher MW and increased flexibility, MW in the range 352 to 659 g mol$^{-1}$, and number of rotatable bonds from 6 to 13. We retrieved 3 data sets that contained 6, 11, and 9 compounds respectively. The first data set was based on a combination of anilinoquinoxilines and anilinoquinazolines EGFR inhibitors, the second on a series of bis-anilino-pyrimidine EGFR inhibitors, and the third were based on the chemical scaffold of pyrazolopyrimidine BTK inhibitors containing an inverted cyanoacrylamide warhead.

For these larger molecules, conformational sampling could be important. Therefore, we performed a test for the conformational dependence of PA. We took the 6 compounds from data set 4, and identified the lowest energy conformation, and also the next lowest energy but structurally different conformation (>2 Å RMSD to the first, see methods) for each molecule. These conformations were then used to calculate the PA. The different conformations show significantly different PAs. Figure 4 shows there are large variations between the $\Delta G_{PA}$ calculated for the two alternative low energy conformations of each molecule. There are some examples such as compound 15 (Figure 4) where the PA for the two different conformations varies by as much as 9.3 kcal mol$^{-1}$, whereas others are closer. In addition, we calculated a Boltzmann weighted $\Delta G_{PA}$ value for all compounds in data set 4, averaged over 4 low energy conformers per molecule (Supporting Information, section 2). However, we find that there is no significant improvement in correlation between PA and experimental reactivity when using this approach (Figure S1, Supporting Information).
The strong conformational dependence of the PA suggested that it would not be an ideal metric for large compounds. We investigated this by performing PA calculations using the method outlined above for all the compounds in these new data sets, but only using the lowest energy conformation. This involved a total of 25 unique drug-like compounds across the three data sets. For each data set, the plot of the relative proton affinity ($\Delta G_{PA}$) against the GSH reactivity rate ($\log k_{GSH}$) revealed no correlation: data set 1: $R^2=0.12$; data set 2: $R^2=0.00$; and data set 3: $R^2=0.14$ (Figure 5).

As shown in Figure 3, the calculated PA for small fragments appears to be predictive for reactivity with GSH. Therefore, we examined it as a surrogate for predicting the reactivity of the larger molecules. We chose all the molecules from data set 4 and converted these to small fragments, so that only the core scaffold from the parent structure remained but each fragment was unique (Figure 6). These fragments are analogous to the types of compounds we studied in data sets 1-3. This approach is similar to the truncation algorithm used by Palazzesi et al\textsuperscript{21}, but we had to modify the approach slightly to avoid duplication of fragments. The calculated PAs for the lowest energy conformation of the full molecules and the fragments are shown in Table 1. The results suggest that the fragment calculations are less predictive for the reactivity of larger molecules and do not correlate with experiment. For instance, the most reactive molecule identified from experiment (F22) was predicted to be the...
least reactive based on the fragment $\Delta G_{PA}$ calculation. Thus, even for the simpler case of GSH compared with protein reactivity, additional parameters beyond the local atomic and electronic environment are important.

Table 1. Calculated $\Delta G_{PA}$ values at the B3LYP-D3/6-311+G** level for fragments and the corresponding parent compound, and experimental reactivity ranking. The predicted relative rank order from calculations is shown in parentheses.

<table>
<thead>
<tr>
<th>Fragment</th>
<th>Fragment $\Delta G_{PA}$</th>
<th>‘Parent’ $\Delta G_{PA}$</th>
<th>Experimental ranking</th>
</tr>
</thead>
<tbody>
<tr>
<td>F15</td>
<td>-11.6 (4)</td>
<td>-4.3 (3)</td>
<td>4</td>
</tr>
<tr>
<td>F18</td>
<td>-7.0 (5)</td>
<td>0.0 (6)</td>
<td>6</td>
</tr>
<tr>
<td>F19</td>
<td>-15.2 (1)</td>
<td>-5.4 (2)</td>
<td>2</td>
</tr>
<tr>
<td>F20</td>
<td>-13.2 (3)</td>
<td>-3.3 (5)</td>
<td>5</td>
</tr>
<tr>
<td>F21</td>
<td>-13.3 (2)</td>
<td>-10.0 (1)</td>
<td>3</td>
</tr>
<tr>
<td>F22</td>
<td>0.0 (6)</td>
<td>-3.3 (4)</td>
<td>1</td>
</tr>
</tbody>
</table>

*Experimental ranking is based on most reactive being assigned ranking of 1, and least reactive a ranking of 6.

Figure 6. Fragments of drug-like compounds from data set 4 used here to calculate fragment $\Delta G_{PA}$ values.

We also investigated how calculated $\Delta E_{\text{react}}$ values, a commonly used alternative computational reactivity metric, correlate with the experimental reactivity. $\Delta E_{\text{react}}$ values were calculated for all 68 compounds from data sets 1-6 (see methods). Generally, a poor correlation between $\Delta E_{\text{react}}$ and experimental GSH reactivity was observed (Figure 7). Across all the data sets, $\Delta G_{PA}$ generally performed better as a reactivity metric than calculated $\Delta E_{\text{react}}$ values, although neither did as well as would be needed for a drug discovery lead optimization setting. In short, four of the six data sets of close analogues showed no appreciable correlation ($R^2 < 0.5$).

![Graph showing GSH reactivity vs reaction energy](image)

Figure 7. Plot of GSH reactivity vs reaction energy, $\Delta E_{\text{react}}$ for data set 1, $R^2=0.05$ turquoise squares; data set 2, $R^2=0.68$ magenta circles; data set 3, $R^2=0.33$ blue triangles; data set 4, $R^2=0.09$ green triangles; data set 5, $R^2=0.50$ orange diamonds; data set 6, $R^2=0.00$ black pentagons.

**Discussion**

The calculation of PA values for small covalent fragments bearing $\alpha,\beta$-unsaturated functionality initially appears to be a promising covalent reactivity metric. For compounds of this type, we observe that increased acidity at the $\alpha$-carbon of the acrylamide group correlates with faster GSH reactivity rates, as expected. This reflects the stability of the enolate intermediate (compound 4 in Figure 2) in the covalent thio-Michael reaction mechanism, which leads to faster reaction. The resulting plots show that $\Delta G_{PA}$ values correlate strongly with experimental GSH reactivity for small fragment compounds. The results are similar to previously published work\textsuperscript{16} that found proton affinity to correlate with $\beta$-elimination rates with $R^2=0.96$.

For the larger and more complex compounds, no correlation is observed between $\Delta G_{PA}$ values and GSH reactivity. To understand this result, we investigated the effect of conformational differences in our protocol by examining in detail the 6 compounds in data set 4. Two sources of potential error were identified, the first being that different input conformations significantly impact the calculated PA for each compound. The difference in PA between conformer pairs obtained from the conformational search (see Methods) can be as large as 9.3 kcal mol$^{-1}$ (compound 15, data set 4, Figure 4A and 5). In this particular case, there was a rotation of the substituted acrylamide-thiol adduct leading to an additional NH-N interaction in one of the conformers that was not present in the other (Figure 4B). The second source of possible error resulted from the difference in conformation between the neutral and ionized species. After a geometry optimization, conformational changes in the ionized species were often observed, arising due to the change in hybridization ($sp^3$ to $sp^2$) of the alpha-carbon upon deprotonation causing a change in intramolecular interactions. Rotations of flexible groups attached to aromatic rings and movements of
the acrylamide-thiol adduct were also observed, suggesting the ionized species were optimized to alternate local minima from the neutral species. We used more stringent convergence criteria during our optimizations in order to address this, however, it did not significantly change the results (Figure S2, Supporting Information). These sources of error result in an overestimation of the calculated PA, suggesting an inaccurate comparison of the relative change in energy associated with proton affinity with the inclusion of conflicting conformational effects, and therefore explain the lack of correlation between PA and experimental reactivity.

An additional factor that could lead to errors in the calculation of ΔGPA values is the contribution to the free energy from low energy vibrational modes. These low energy modes are particularly important for the large, flexible compounds in data set 4, 5 and 6, and are difficult to calculate accurately and can therefore lead to errors in the free energy calculation. To investigate the affects that these modes have on our free energy values, we used the Quasi Harmonic Oscillator (QHO) approximation to scale all frequencies under 100 cm⁻¹. We found it useful in the context of PA. 

Although previous studies have reported success in using a fragment based approach to improve correlation between ligand-only reactivity metrics and experimental reactivity, we do not find it useful in the context of PA. Inspection of the experimental data and molecular structures casts further doubt on using a fragment approach to predict the reactivity of larger molecules. For instance, compounds 15 (data set 4), 23, 24 and 25 from data set 5 (Figure 8) all contain essentially the same fragment (a dimethylamine substituted phenylacrylamide group), but their experimental reactivity differs by ~2 log units. This variation in reactivity highlights that modifications to compounds distal from the site of reaction can have a significant impact on reactivity (conformational effects will also contribute).

The ΔEreact calculations also showed generally poor correlation with GSH reactivity. This is in contrast to previous work by Lonsdale and colleagues who reported a correlation of R²=0.69. Their study combined diverse molecules and warheads. ΔEreact values may offer reasonable reactivity prediction for compounds that differ greatly in reactivity, but will generally not predict reactivity within a chemical series, as shown by our results. It is also likely that reaction energy calculations are also more susceptible to conformational effects than ΔGPA, due to the greater variability in geometry optimization of methyl thiolate relative to the reactive warhead.

Although previous studies have reported success in using a fragment based approach to improve correlation between ligand-only reactivity metrics and experimental reactivity, we do not find it useful in the context of PA. Inspection of the experimental data and molecular structures casts further doubt on using a fragment approach to predict the reactivity of larger molecules. For instance, compounds 15 (data set 4), 23, 24 and 25 from data set 5 (Figure 8) all contain essentially the same fragment (a dimethylamine substituted phenylacrylamide group), but their experimental reactivity differs by ~2 log units. This variation in reactivity highlights that modifications to compounds distal from the site of reaction can have a significant impact on reactivity (conformational effects will also contribute).

For biological reactivity, PA calculations require careful consideration that input geometries represent local minima corresponding to the noncovalently or covalently bound conformation, as opposed to global energy minima adopted in solution. Reactivity of covalent fragments with GSH in solution is likely to be less constrained, but still dominated by a small number of conformations. Although conformational searches and low energy conformers were selected for our calculations, it is possible that these low energy conformers are not representative of the conformations that react with GSH in solution. Even Boltzmann weighted PA values, averaged over 4 low energy conformers that would be expected to dominate reactivity did not improve reactivity predictions (Supporting Information, Figure S1). In the protein environment it is easy to envisage that the reaction proceeds from the bound conformation of the ligand and placed optimally for reactivity with the cysteine in the protein pocket. Replicating these reactive conformations is difficult to achieve for PA calculations and in part explains the lack of predictive ability of ligand-only approaches and indicates the need for treatment of biological reactivity in situ in biological targets.

Figure 8. Four compounds that contain the same dimethylamine substituted phenylacrylamide fragment (highlighted in red) but have a wide range of experimental GSH reactivity, covering almost 2 log units.

All other factors being equal, the change in reactivity caused by a small structural modification may be captured by PA calculations. Indeed, there could be scenarios in lead optimization where analogues are so similar that this method delivers results that translate into useful design prioritization. PA and reaction energies seem to be attractive reactivity metrics due to their simplicity. For PA, only geometry optimizations of the covalent thiol adduct and its conjugate base are required, negating the need to control the position of methyl thiolate during transition state searches and calculations of reaction energies. However, we see here that, for many common modifications in the medicinal chemistry data sets, further effects are at play. The calculation of PA, reaction energies and reaction barrier heights all require careful consideration of conformational variations in each compound as it is important to ensure that differences in the energy calculation do not arise from competing conformational effects. Lonsdale et al. found that the calculation of reaction energy barriers through transition state optimizations to be particularly challenging for large ‘drug like’ compounds and opted to calculate reaction energies instead.

For biological reactivity, PA calculations require careful consideration that input geometries represent local minima corresponding to the noncovalently or covalently bound conformation, as opposed to global energy minima adopted in solution. Reactivity of covalent fragments with GSH in solution is likely to be less constrained, but still dominated by a small number of conformations. Although conformational searches and low energy conformers were selected for our calculations, it is possible that these low energy conformers are not representative of the conformations that react with GSH in solution. Even Boltzmann weighted PA values, averaged over 4 low energy conformers that would be expected to dominate reactivity did not improve reactivity predictions (Supporting Information, Figure S1). In the protein environment it is easy to envisage that the reaction proceeds from the bound conformation of the ligand and placed optimally for reactivity with the cysteine in the protein pocket. Replicating these reactive conformations is difficult to achieve for PA calculations and in part explains the lack of predictive ability of ligand-only approaches and indicates the need for treatment of biological reactivity in situ in biological targets.
Conclusions
In summary, calculated proton affinity values can serve as a useful reactivity metric for small molecule fragment compounds. However, we find proton affinity to be a poor reactivity metric for larger, 'drug-like' compounds. Calculated reaction energies are generally worse as a reactivity metric than proton affinities for both fragments and drug-like compounds. Our work shows that ligand-only approaches such as the calculation of reaction energies, barrier heights and proton affinity values are insufficient for reliable covalent reactivity prediction. These methods oversimplify covalent determinants of reactivity, and do not account for factors that will affect the reactivity of a covalent drug 'in-situ' (e.g. effects of the environment within a protein target). Calculated PA and reactivity are conformation dependent; experimental studies of ligand-only reactivity may also be confounded by conformational effects. More detailed methods are needed to predict reactivity of covalent drugs in a protein environment, which include the (e.g. electrostatic) effects of the environment, the bound conformation and solvation of the covalent inhibitor in the target site; such methods, capable of modeling reactivity within proteins, including combined QM/MM techniques.

ASSOCIATED CONTENT
Supporting Information
The Supporting Information is available free of charge on the ACS Publications website.

Molecular structures, calculated proton affinity values and reaction energies for all compounds studied (PDF)

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ABBREVIATIONS
TCI, targeted covalent inhibitor; PA, proton affinity; QM, quantum mechanical; GSH, glutathione; DFT, density functional theory; PES, potential energy surface; TS, transition state.

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