SAMPL6 pKa reference calculations using Epik and Jaguar

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2 NOTES: Objectives for this manuscript

track down artifacts that seem to be present in free energy values

- Provide a baseline for the expectation of performance of empirical models (Epik) and DFT models (Jaguar) on kinase inhibitor-like molecules and molecular fragments; draw comparisons to earlier benchmarks for these tools
- Describe basic concepts, along with strengths and weaknesses of each approach (Epik and Jaguar)
- · Lessons learned for participating in the challenge
 - How should we predict macroscopic pKas from microscopic pKas? Explain possibilities and justify choice
 - * Sequential titration (Epik)
 - * <n protons>/charge vs pH inflection points: Matches electrochemical titration, but not UV-metric
 - * Is there a way to predict what UV-metric would observe from populations vs pH?
- For microscopic pKa prediction: Epik may enumerate a different set of microstates than other tools (we proposed these as newly labeled states)
- Epik does not return state populations below a certain threshold, so we could not consider states proposed by SAMPL6 [Check this]
 - For worst predicted compounds, provide an explanation as to why the methods may perform poorly (descriptors?) [may not have enough data points to draw conclusion]

28 Figures and Tables

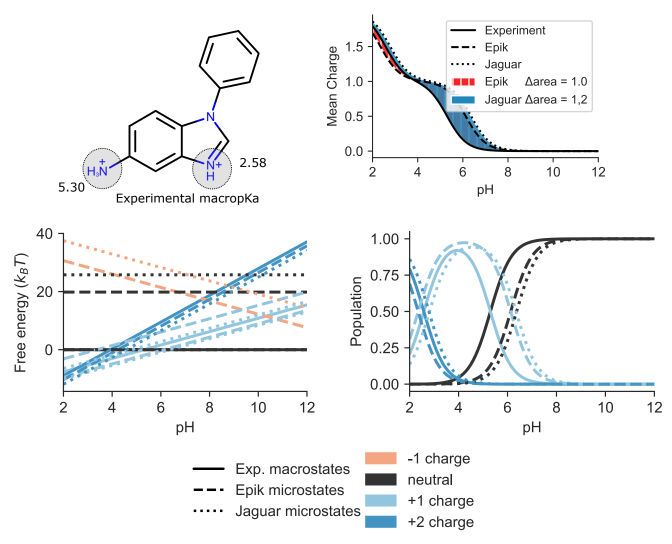


Figure 1. Epik and Jaguar microscopic pKas can be used to generate free energies (bottom left) or, populations (bottom right), and recapitulate the binding curve (top right) when the charge of each species is added up. Molecule SM14 pKa values (top left) were measured using UV (citation) for the purpose of the SAMPL6 challenge, and based on a subsequent NMR experiment the most likely sites of titration were established. We used Epik and Jaguar to predict microscopic pKas (denoted as Type I predictions in SAMPL6). The macrostate populations from experiment, and microstate populations from predictions can be compared directly to an experimental titration curve derived from UV. Since NMR data was available for this molecule, the charge state in the experiment is also known.

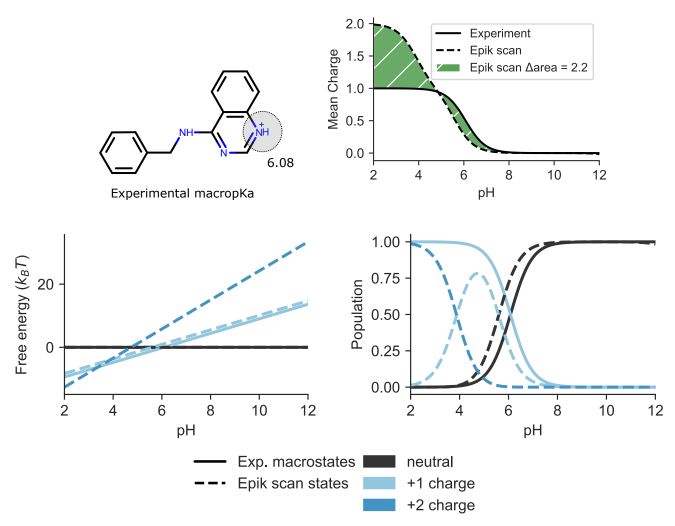


Figure 2. Epik scan provides sequential pKa values for adding single protons, which can approximate a macroscopic titration curve. Molecule SM07 pKa values were measured using UV (citation) for the purpose of the SAMPL6 challenge, and based on a subsequent NMR experiment the identity of the microstates were identified. We used Epik scan mode to predict most prevalent microstates from sequential pKas, which can approximate macroscopic pKas. The macrostate populations from experiment, and microstate populations from predictions can be compared directly to an experimental titration curve derived from UV pKa values. Since NMR data was available for this molecule, the charge identity in the experiment is also known.

Table 1. Overall performance of each method using either the closest pKa matching, Hungarian pKa matching, or sequentially aligned pKas, and the area between the experimental and predicted macroscopic charge titration curve

		рКа	
		RMSE	
	closest	hungarian	align
Epik micropka	0.5; [0.3, 0.7]	0.5; [0.3, 0.7]	
Epik scan	0.8; [0.6, 1.0]	0.8; [0.6, 1.0]	0.8; [0.6, 1.0]
Jaguar	0.3; [0.2, 0.5]	0.3; [0.2, 0.5]	_
		Pearson ρ	
	closest	hungarian	align
Epik micropka	0.96; [0.93, 0.99]	0.96; [0.92, 0.99]	
Epik scan	0.94; [0.88, 0.97]	0.94; [0.87, 0.97]	0.94; [0.87, 0.97]
Jaguar	0.986; [0.966, 0.995]	0.986; [0.969, 0.995]	_
	Charge vs pH		
	Δ area		
Epik micropka	1.8; [1.5, 2.1]		
Epik scan	1.7; [1.2, 2.2]		
Jaguar	1.5; [1.0, 2.1]		

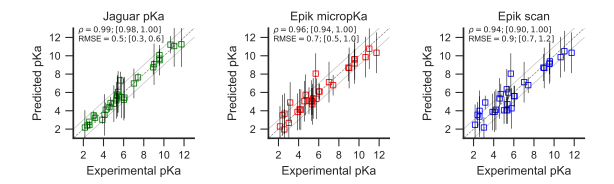


Figure 3. Comparison of computed pKa values to experimental macroscopic pKa values. *Left:* Microscopic pKa values for Jaguar were matched of the experimental dataset using algorithm 1. An alternative alignment of pKa values using the Hungarian algorithm (??) is available as a supplementary figure. *Middle:* Microscopic pKa values for Epik were matched of the experimental dataset using algorithm 1. An alternative alignment of pKa values using the Hungarian algorithm (??) is available as a supplementary figure. *Right:* Macroscopic pKa values for Jaguar were matched of the experimental dataset using algorithm 1. An alternative alignment of pKa values using the Hungarian algorithm (??) is available as a supplementary figure.

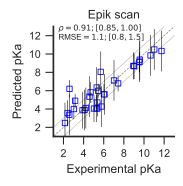


Figure 4. Epik macroscopic pKa values compared to macroscopic experimental pKa values, sequentially matching pKa values Comparison of the Epik sequential scan with entire experimental data set is shown, allowing only sequential pKa values to match experiment.

Table 3. Microscopic pKa per molecule for each method as matched by the "closest" algorithm (algorithm 1), compared to experiment. The uncertainty indicated is the standard error from the experiment, and the reported standard error by the prediction method.

Molecule	Experimental	Epik-micropka	Jaguar	Epik-scan
SM01	9.53 ± 0.01	10 ± 1	9.7 ± 0.8	9 ±1
SM02	5.03 ± 0.01	5 ±1	5 ± 2	4 <u>±</u> 1
SM03	7.02 ± 0.01	7 ± 2	7 ± 2	7 ± 2
SM04	6.02 ± 0.01	6.1 ± 0.9	5 ± 2	6 <u>±</u> 1
SM05	4.59 ± 0.01	5 ± 2	4.4 ± 0.5	5 ± 2
SM06	3.03 ± 0.04	3 ± 2	3.5 ± 0.5	2.2 ± 0.9
SM06	11.74 ± 0.01	10 ± 2	11 ± 2	10 ± 2
SM07	6.08 ± 0.01	6.1 ± 0.9	5 ± 2	6 ± 2
SM08	4.22 ± 0.01	4 ± 2	4.1 ± 0.8	4 ± 2
SM09	5.37 ± 0.01	5 ±1	6 ± 2	4.0 ± 0.9
SM10	9.02 ± 0.01	9 ± 2	9 ± 2	9 <u>+</u> 1
SM11	3.89 ± 0.01	4 ± 2	3.0 ± 0.4	4 ± 2
SM12	5.28 ± 0.01	5 ± 1	5 ± 2	4 ± 1
SM13	5.77 ± 0.01	5 ± 2	5 ± 2	4 ± 2
SM14	5.30 ± 0.01	5 ± 2	5.3 ± 0.7	6 ± 2
SM14	2.58 ± 0.01	3 ± 2	2.8 ± 0.1	3 ± 2
SM15	8.94 ± 0.01	9 <u>±</u> 1	8.7 ± 0.8	9 ±1
SM15	4.70 ± 0.01	6 ±1	4.8 ± 0.7	6 ±1
SM16	5.37 ± 0.01	5 ± 2	6 ± 2	4.7 ± 0.9
SM16	10.65 ± 0.01	10 ± 2	11.2 ± 0.9	10 ± 2
SM17	3.16 ± 0.01	4.9 ± 0.8	3.2 ± 0.6	4.9 ± 0.7
SM18	9.58 ± 0.03	10 ± 2	10 ± 2	9 ± 2
SM18	2.15 ± 0.02	2 ± 2	2 ± 2	2 ± 2
SM18	11.02 ± 0.04	11 ±2	11 ± 2	10 ± 2
SM19	9.56 ± 0.02	9 ± 2	10.1 ± 0.7	9 ± 2
SM20	5.70 ± 0.03	8 ± 2	7.3 ± 0.9	8 ± 2
SM21	4.10 ± 0.01	4 ± 2	4 ± 2	4 ± 2
SM22	7.43 ± 0.01	6.8 ± 0.9	7.6 ± 0.5	6.8 ± 0.9
SM22	2.40 ± 0.02	4 ± 2	2.4 ± 0.5	4 <u>±</u> 1
SM23	5.45 ± 0.01	5 ± 2	6 ± 2	6.0 ± 0.8
SM24	2.60 ± 0.01	4 ± 2	2.5 ± 0.3	4 ±1

Table 2. Macroscopic pKas computed via Epik's sequential scan procedure. Experimental and computed pKa values matched using sequential alignment of pKa values.

Molecule	Experimental	Epik-scan
SM01	9.53 ± 0.01	9 ± 1
SM02	5.03 ± 0.01	4 ± 1
SM03	7.02 ± 0.01	7 ± 2
SM04	6.02 ± 0.01	6 ±1
SM05	4.59 ± 0.01	5 ± 2
SM06	3.03 ± 0.04	4 ± 1
SM06	11.74 ± 0.01	10 ± 2
SM07	6.08 ± 0.01	6 ± 2
SM08	4.22 ± 0.01	4 ± 2
SM09	5.37 ± 0.01	4.0 ± 0.9
SM10	9.02 ± 0.01	9 ± 1
SM11	3.89 ± 0.01	4 ± 2
SM12	5.28 ± 0.01	4 ± 1
SM13	5.77 ± 0.01	4 ± 2
SM14	2.58 ± 0.01	3 ± 2
SM14	5.30 ± 0.01	6 ± 2
SM15	4.70 ± 0.01	6 ±1
SM15	8.94 ± 0.01	9 ±1
SM16	5.37 ± 0.01	4.7 ± 0.9
SM16	10.65 ± 0.01	10 ± 2
SM17	3.16 ± 0.01	4.9 ± 0.7
SM18	2.15 ± 0.02	2 ± 2
SM18	9.58 ± 0.03	9 ± 2
SM18	11.02 ± 0.04	10 ± 2
SM19	9.56 ± 0.02	9 ± 2
SM20	5.70 ± 0.03	8 ± 2
SM21	4.10 ± 0.01	4 ± 2
SM22	2.40 ± 0.02	4 ± 1
SM22	7.43 ± 0.01	6.8 ± 0.9
SM23	5.45 ± 0.01	6.0 ± 0.8
SM24	2.60 ± 0.01	4 ±1

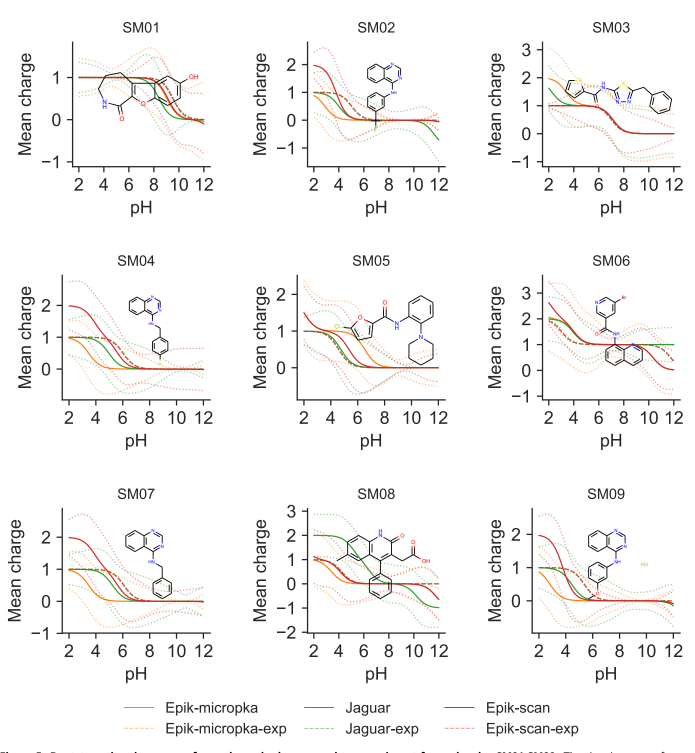


Figure 5. Bootstrap titration curves for each method compared to experiment for molecules SM01-SM09. The titration curve for each method is shown as a solid line, and 96 % confidence intervals from bootstrap have been shown as dotted lines. A numerical comparison to experiment is presented in ??.

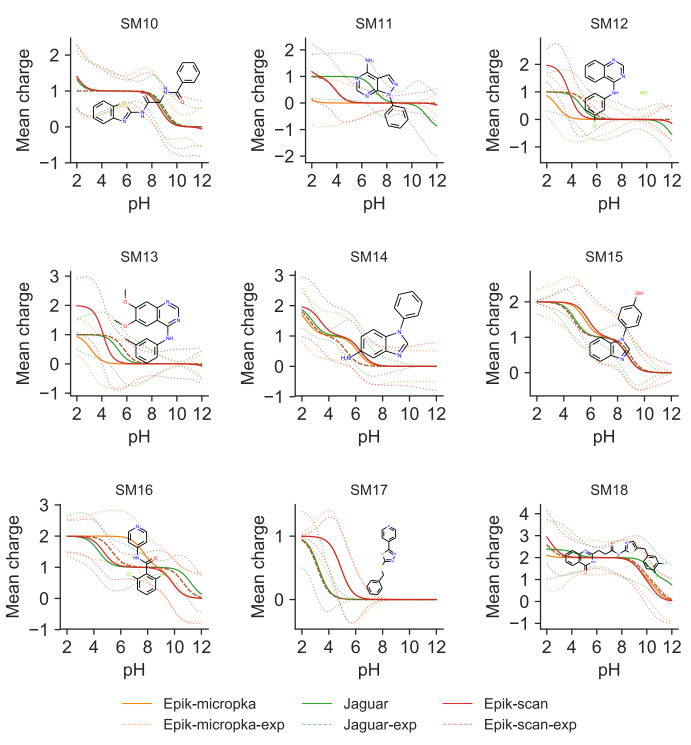


Figure 6. Bootstrap titration curves for each method compared to experiment for molecules SM10-SM18. The titration curve for each method is shown as a solid line, and 96 % confidence intervals from bootstrap have been shown as dotted lines. A numerical comparison to experiment is presented in ??.

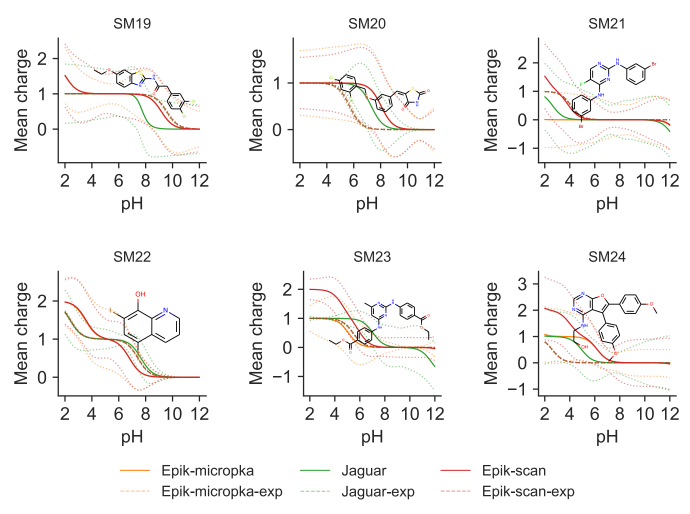
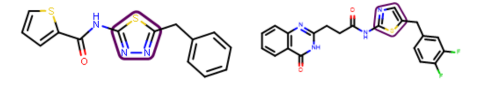


Figure 7. Bootstrap titration curves for each method compared to experiment for molecules SM19-SM24. The titration curve for each method is shown as a solid line, and 96 % confidence intervals from bootstrap have been shown as dotted lines. A numerical comparison to experiment is presented in ??.

Table 4. Mean charge titration curve comparison with respect to experiment (Δ area) with confidence intervals. Confidence intervals were obtained by gaussian bootstrapping over pKa values from the standard errors reported by the software.

	Epik micropka	Jaguar	Epik scan
Molecule			
SM01	0.5; [0.2, 4.9]	1.1; [0.0, 3.5]	0.5; [0.1, 4.0]
SM02	2.1; [0.2, 3.1]	1.6; [0.3, 5.3]	1.9; [0.7, 4.2]
SM03	1.5; [1.0, 4.9]	0.5; [0.8, 4.8]	0.1; [0.2, 4.7]
SM04	2.6; [0.2, 4.0]	1.1; [0.1, 4.0]	2.2; [0.3, 5.0]
SM05	2.2; [0.6, 4.7]	0.2; [0.1, 4.5]	1.1; [0.5, 4.6]
SM06	2.2; [0.7, 9.5]	1.3; [0.6, 4.7]	2.6; [0.6, 6.8]
SM07	2.6; [0.1, 4.1]	0.8; [0.1, 4.1]	2.2; [0.7, 4.8]
SM08	1.2; [0.5, 6.2]	7.1; [3.3, 9.2]	0.6; [0.1, 5.8]
SM09	2.5; [0.3, 3.4]	1.0; [0.1, 4.3]	2.3; [0.6, 3.3]
SM10	0.6; [0.2, 5.1]	0.3; [0.2, 5.3]	0.6; [0.1, 5.6]
SM11	1.9; [0.2, 4.1]	3.9; [0.5, 6.8]	0.1; [0.2, 5.0]
SM12	2.5; [0.2, 3.4]	0.9; [0.4, 4.7]	2.2; [1.1, 3.6]
SM13	2.6; [0.2, 3.8]	0.5; [0.1, 4.0]	2.8; [0.3, 5.0]
SM14	1.0; [0.8, 5.4]	1.2; [0.1, 4.1]	1.8; [0.6, 5.2]
SM15	1.1; [0.4, 4.6]	0.3; [0.2, 2.4]	1.3; [0.4, 4.2]
SM16	2.9; [0.7, 4.7]	1.5; [0.5, 4.4]	1.4; [0.6, 4.7]
SM17	1.7; [0.1, 3.2]	0.1; [0.0, 1.6]	1.7; [0.2, 3.1]
SM18	0.8; [1.2, 9.7]	2.2; [1.5, 7.9]	1.0; [1.1, 8.2]
SM19	0.7; [0.2, 5.2]	1.8; [0.6, 6.7]	0.7; [0.2, 5.9]
SM20	2.3; [0.1, 4.7]	1.6; [0.2, 3.4]	2.3; [0.1, 4.8]
SM21	2.2; [0.8, 5.5]	1.6; [0.5, 7.0]	0.5; [0.4, 6.7]
SM22	1.6; [0.5, 4.8]	0.2; [0.1, 3.4]	1.6; [0.3, 3.9]
SM23	0.4; [0.2, 3.5]	2.1; [0.7, 5.7]	3.1; [1.9, 4.8]
SM24	3.6; [0.1, 5.0]	2.3; [0.2, 4.9]	5.6; [3.0, 8.6]



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Figure 8. Problematic lewis structure generated by Epik. Epik generated protonation states with invalid Lewis structure for SM03 and SM18, with a pentavalent nitrogen present in the thiazole/thiadiazole (the protonation pattern displayed on the right). One of these was also present in the set provided as part of SAMPL6 microstates, namely SM03_micro021. A valid Lewis structure could not be figured out for this protonation state.

See Images folder for titration curves and free energy images for each category. Will be added to supplementary info later.

Abstract The goal of the SAMPL6 pKa Challenge was to evaluate the performance of small molecule pKa prediction methods from challenge participants in a blinded fashion on a set of small, drug-like molecules resembling kinase inhibitor fragments. To

provide a useful point of comparison for blind participant predictions that may use experimental methods still under development,
we performed reference benchmark calculations using a popular empirical model (Epik) and quantum chemical approach (Jaguar)
from Schrödinger. Epik predicts microstate populations and pKas using a Hammett-Taft type model, while Jaguar is a fast
DFT quantum chemical method. In this work, we discuss how these reference calculations were performed, provide a broad
assessment of the performance of the method, and highlight challenges and considerations in predicting pKas to benchmark
against experiment macroscopic pKa measurements.

40 Introduction

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Titratable sites are ubiquitous in druglike small molecules.

JDC: It would be very useful if we could cite some sort of reference discussing how important protonation states are in small molecule drugs or druglike molecules, but I'm failing to find useful references right now. Alternatively, we could pull pKas from DrugBank https://www.drugbank.ca/releases/latest or use Epik to make our own plots, like the kinase inhibitor pKa plot we had previously generated.

Large-scale computational surveys suggest that 60% of all protein-ligand complexes undergo a change in ionization state upon binding [1], either due to protonation state changes of the small molecule or the protein (where roughly a third of all protein residues are ionizable [2]). More generally, protonation state effects—in which the dominant protonation, charge, or tautomer state shifts upon binding, or a mixture of protonation states are significantly populated in complex or solution—has the potential to cause large modeling errors if these effects are neglected. In the SAMPL5 distribution coefficient (logD) challenge, for example, protonation state effects were determined to be a major contributor to loss in accuracy for the otherwise mundane task of predicting a transfer free energy between aqueous and cyclohexane phases [3].

To isolate the question of how well pKa effects could be modeled—and therefore how accurately the community could address these effects—the SAMPL6 challenge featured a blind pKa prediction component, as an intermediate step to logD predictions in which we provide participants with pKas and later have them predict both pKa and logD [?]. The SAMPL6 pKa challenge consisted of predicting macroscopic pKas measured by UV-metric titration for a set of small moelcules that resembled kinase inhibitors and their fragments [?]. As participants in the SAMPL6 pKa challenge were expected to utilize a wide variety of methods still under development, we endeavored to provide a useful baseline reference set of predictions using well-established, widely-deployed, commercially available methods. We selected both an empirical method (Epik [4]) and quantum chemical method (Jaguar [5]) from the Schrödinger Suite of computational chemistry software, version 20XX-X.

Reference calculations—which were not fully blinded—were performed in a manner that attempted to mimic standard use, using recommended settings for each program, without significantly modifying the input parameters. As the computation of UV-metric macroscopic pKas from the microscopic pKas predicted by the tools is not necessarily completely straightforward, we considered several alternative possibilities, which we discuss in more detail. We provide an analysis and broad assessment of the performance of the two methods, and highlight challenges and considerations in predicting pKas to benchmark against experiment. All analysis tools used to perform this study are available via GitHub at https://github.com/choderalab/SAMPL6-Reference-pKa-Calculations

Methods

- 66 Epik
- 67 Summary of how Epik works
 - Fill in snippet from john shelley
- 69 Jaguar

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- Summary of how Jaguar computes pKas
 - Could be filled in by Art Bochevarov?

Microstate predictions

- ₇₃ Epik: Fast empirical pKa predictions
 - were performed between pH 2-12

- Reported states with a minimum population of $e^{-10/RT}$
 - States proposed by SAMPL6 but not predicted by Epik were not considered in analysis.

Jaguar: Ab initio quantum chemical pKa predictions

- Ran using default settings (5 conformations) for each of the pre-enumerated specified microstate pairs.
- Input structures were first minimized using MMFF.
 - · Additional minimization was performed for structures for which scf/geopt did not converge

81 Predicting macroscopic pKas

- Epik and Jaguar predict microscopic pKas or microstate energies, which must be translated into macroscopic pKas for comparison
- to experiment. Several reasonable choices are possible for translating these microscopic properties into macroscopic pKas.

84 Sequential titration of dominant species

- Sequential titration avoids the need to compute the energies of all protonation states
- Epik can automatically perform a sequential scan (which can also be used in Jaguar with some automation)
- For this experiment, we started from the predicted highest-occupancy microstate at pH 7 and went in both directions, returning pKas between 2-12

89 Virtual electrochemical titration

 Average charge vs pH to match electrochemical titrations, look at the mean signed deviation (MSD) to see whether the charge curve behavior matches

92 Predicting UV-metric titration

- Actual experiments were UV-metric, where only UV-active transitions are observable
- There is potentially a route to predicting UV-active transitions via identifying of microstates that would have different UV spectra, perhaps using SMARTS matches or simple QM absorption spectra calculations?

96 Results

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- · How much conformation-dependence is there in Jaguar-derived pKas (or state energies)?
- Does sequential scan or mean molecular charge provide better agreement with experimental macroscopic pKas? Would it be worthwhile to develop predictive UV-metric models?
- Comparison of observed accuracies to previously reported/expected accuracies; expected accuracy on kinase inhibitors derived from this study
- How does Epik compare to Jaguar in terms of accuracy and computational cost?
- Discussion of outliers

104 Discussion

105 Overall performance

- Some low probability structures produced by Epik included questionable protonation states of a heterycyclic moiety, present in SM03 and SM18 (Figure 8).
 - Overall performance of Epik and Jaguar based on various metrics (figures/tables)

109 Matching of experimental and calculated pKas

- The experiments do not provide any microscopic information on what atom, or microstate a pKa belongs to. Therefore, it is necessary to perform a matching of pKa values between experiment and predicted pKa values. There are several ways one could go about this, each strategy can prioritize a different aspect to match. We consider three algorithms:
- 113 Closest pKa matching
- 114 Hungarian pKa matching
- This algorithm, also known as linear sum assignment. It finds the combination of pKa that minimizes the overall cost by picking
- rows and columns in a matrix, also considering the cost of not mapping certain pKa values in the case of different numbers of predictions and experimental values.

118 Sequential pKa matching

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For macroscopic only, because it doesn't make sense to sequentially align microscopic pKas to a set of macroscopic pKas algorithm 2

Macroscopic titration curves

Compare macroscopic titration results

Population curves from pKa

• Description of the population curve generation from pKa

$$g_i(pH) = \beta \left(n_i * pH - \sum_j pKa_j \right)$$
 (1)

$$\pi_i(\text{pH}) = \frac{e^{-g_i(\text{pH})}}{\sum_i e^{-g_i(\text{pH})}} \tag{2}$$

Virtual electrochemical titration

• Description of the calculation of the mean charge curve

$$\langle q_{\text{total}}\rangle(\text{pH}) = \sum_{i} q_{i} \times \pi_{i}(\text{pH})$$
 (3)

129 Microscopic pKa values

Compare microscopic pKa prediction results

131 Descriptor analysis

- which moieties are harder to predict?
- Do certain descriptors correlate with variance/absolute errors?
- Do number of rotatable bonds affect epik/jaguar results (conformations missing from prediction)?

135 Comparison with other methods

This subsection depends on how much information we already have in the overview paper

- · Compared to other Hammet-Taft methods, does Epik stand out? [based on early overview results?]
- · Compared to other QM methods, does Jaguar stand out? [based on early overview results?]

139 Suggestions for future challenges

- Future challenges could benefit from NMR experiments for microstates
- Potentiometric titrations could capture states that may have been left out
- Probabilistic models (i.e. bayesian hierarchical models) could be constructed for the analysis of experiments.

143 Conclusions

- Epik and Jaguar can be useful as baselines
- We can use various ways of judging each method.
- A conclusion about the accuracy
- Some take away points about difficulties about comparing microscopic to macroscopic
 - How a future challenge could make this easier

Code and data availability

Input files and analysis scripts are available at https://github.com/choderalab/SAMPL6-reference-pka-calculations

151 Author Contributions

(Follow the CRediT Taxonomy)

153 Acknowledgments

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156 Disclosures

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```
Result: Mapping of each experimental pKa i to predicted pKa j C is constructed, where every row i is an experiment, and every column j a prediction; C_{ij} = \text{cost}(\text{pKa}_{\text{exp},i}, \text{pKa}_{\text{pred},j}); while C.\text{size} > 0 do
\begin{array}{c|c} k, l = \text{arg min}(C_{ij});\\ \text{assign experimental value } k \text{ to prediction } l;\\ \text{remove row } k, \text{ column } l \text{ from } C;\\ \text{end} \end{array}
```

remaining values are unmatched;

Algorithm 1: This algorithm matches experiment with prediction based on how close each value is, one pKa value at a time. Unless the matrix *C* is square, some values will be unmatched. Those leftover pKas are returned at the end. It uses a cost function, such as root mean square deviation, to assess how close two values are.

```
Result: Mapping of each experimental pKa i to predicted pKa j
I = sorted experimental pKas;
J = sorted predicted pKas;
length = max(I.size, J.size);
Append placeholders such that I.size = length;
Append placeholders such that J.size = length;
min = \infty;
solution = J rolled 0 times;
for n in 0..length do
   S = | rolled n times ;
   total = 0.0;
   for m in 0..length do
       if I_m is placeholder then unmatched experiment
          if J_m \leq 7.0 then
           I_m = 0.0
          else
           I_m = 14.0
       else if J_m is placeholder then unmatched prediction
          if I_m \leq 7.0 then
           J_m = 0.0
          else
           J_m = 14.0
          end
       end
       total = total + cost(I_m, J_m);
   if total < min then solution is better
       min = total;
       solution = S;
   end
```

Algorithm 2: Sequential pKa mapping. It uses a cost function to measure cost, and it rolls (shifts by one, and reintroduces last element as first). Any unmatched pKas are represented by matching with a placeholder value. To calculate the cost, the placeholder is replaced by either 0, or 14, depending on whether the unmatched value is above or below 7.0.

191 Supplementary Information

- Titration curves and free energy plots for each compound, by each method
- The mean charge/deviation curves for each compound and each method
- .mae and .sdf files with results
- scripts and or jupyter notebooks for analysis
- csv version of tables

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