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Exploratory analysis of kinetic solubility measurements of a small molecule library

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ABSTRACT

Kinetic solubility measurements using prototypical assay buffer conditions are presented for a ~58,000 member library of small molecules. Analyses of the data based upon physical and calculated properties of each individual molecule were performed and resulting trends were considered in the context of commonly held opinions of how physicochemical properties influence aqueous solubility. We further analyze the data using a decision tree model for solubility prediction and via a multi-dimensional assessment of physicochemical relationships to solubility in the context of specific ‘rule-breakers’ relative to common dogma. The role of solubility as a determinant of assay outcome is also considered based upon each compound’s cross-assay activity score for a collection of publicly available screening results. Further, the role of solubility as a governing factor for colloidal aggregation formation within a specified assay setting is examined and considered as a possible cause of a high cross-assay activity score. The results of this solubility profile should aid chemists during library design and optimization efforts and represent a useful training set for computational solubility prediction.

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1. Introduction

Aqueous solubility is a governing property for how small molecules interact with biomolecules (proteins, nucleic acids, etc.) and living systems (cells, tissues and whole organisms). A broader appreciation of aqueous solubility is changing how researchers pursue the drug discovery process from library design to screening to hit optimization. The physicochemical properties of small molecules intended for primary screening and lead development changed greatly as drug discovery approaches evolved in response to the advent of combinatorial chemistry and high-throughput screening (HTS).^{1–5} The synthesis and screening of hundreds of thousands of small molecules made it difficult and, to some, unnecessary to pay attention to the physicochemical properties of all library members and, as a result, screening hits became increasingly lipophilic in nature. A growing contingent of researchers began to consider the increasing difficulty in transforming screening hits into clinical candidates by asking fundamental questions about the differences between failed lead compounds and approved drugs. Numerous lessons resulted from these analyses

including the set of parameters known as the rule of five first defined by Lipinski.⁶ In this analysis, Lipinski et al. scrutinized a set of >2000 orally bioavailable clinical agents for physical and calculated properties including molecular weight, H-bond donors, H-bond acceptors and *c Log P*. The results suggested that the 90th percentile of drug-like compounds with acceptable solubility and permeability possessed a MW under 500, fewer than five H-bond donors and 10 H-bond acceptors and a *c Log P* value less than five. Subsequent guidelines have been set forth that include other physicochemical descriptors including compound flexibility and polar surface area.²

An additional consequence of the combichem-HTS revolution was the requirement for a carrier solvent for small molecules entering HTS assays. The kinetics of these assays required the rapid aqueous/buffer solvation of each library member and dimethyl sulfoxide (DMSO) was widely adopted as an appropriate solvent for the storage and dispensing of compound libraries into assay wells. This was a sharp departure from experiments that allow an agent to reach aqueous solubility equilibrium over time. The former version of solubility (referred to as kinetic solubility) is considered more appropriate for discovery settings while the latter (referred to as thermodynamic solubility) often takes precedent in formulation and dosing studies.^{7,8} The differences associated with kinetic and thermodynamic solubilities are often ignored in discovery settings.

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Strategies for optimizing libraries for kinetic solubility have greatly aided both assay performance and the ability of screening leads to enter optimization efforts.^{9,10} Many of these strategies derive from a combination of chemist's intuition and the critical analyses of existing data sets leading to predictive models of solubility. Unfortunately, the ability to gather true aqueous solubility values at a defined pH on large compound collections has been limited by the technologies and cost of acquiring such data and most data sets are restricted to related analogues being evaluated as part of an optimization effort. These data sets are often compromised due to inadequate structural diversity of the compounds, inconsistencies in methods and bias toward expected outcomes. Despite imperfect training sets, numerous computational tools (commercial and noncommercial) for solubility prediction exist to help during library design and optimization efforts.^{11–15} Solubility measurements from a large compound collection achieved via a common methodology would be useful in terms of validating current dogma and as a relevant training set for advanced computational models. In 2009, Clark and co-workers reported the kinetic solubilities from a drug-like collection of >700 compounds and provided an analysis of the results in terms of selected physical and calculated descriptors of the library members.¹⁶ In 2010, Hill and Young reported an analysis of kinetic solubility of a large compound library (~100 K) and experimentally derived values of hydrophobicity ($\text{Log } D_{\text{pH } 7.4}$) for a subset of this library (~20 K).¹⁷ This report provided enlightening lessons on the relationship between calculated and experimentally determined $\text{Log } D/\text{Log } P$ values and also explored the impact that aromatic ring content had on solubility.

Here, we describe an exploratory analysis of kinetic solubility measurements for 57,857 compounds of the NIH Molecular Libraries Small Molecule Repository (MLSMR). We related the solubility of this library to specific compound physical characteristics and calculated properties. Further, we examine subsets of this data to help understand compounds that deviate from expected trends and specific 'rule-breakers' in order to better advise chemists hoping to optimize agents with undesirable physicochemical properties. We also examine the relationship between solubility and the frequency of reported activities from primary screens with a particular focus on agents that are putative aggregators within a reported β -lactamase screen.¹⁸ Importantly, the results from this study are publically available through the PubChem database to allow researchers access to this valuable data set (<http://pubchem.ncbi.nlm.nih.gov/>).

2. Method

Solubility measurements were accomplished from stock 10 mM DMSO solutions (6 μL) dispensed into PBS buffer (294 μL , pH 7.4) via Chemiluminescent Nitrogen Detection (CLND).¹⁹ The equimolar nitrogen response of the detector was calibrated using TRIZMA base at 28 concentrations spanning the dynamic range of the instrument from 0.08 to 4500 $\mu\text{g/mL}$ nitrogen and the measured solubility values were corrected for background nitrogen. On board performance indicating standards (Imipramine HCl, Sulfamethizole and Astemizole) were used to validate assay results. A detailed description of the CLND method is presented in the [Supplementary data](#). The entire data set is deposited in the PubChem database (AID 1996). In PubChem, each compound's individual solubility is listed and the data is additionally organized to reflect low solubility (<10 $\mu\text{g/mL}$), medium solubility (10–60 $\mu\text{g/mL}$) and high solubility (>60 $\mu\text{g/mL}$) with 3,060 structures being annotated as below the limit of quantification (<LOQ). We obtained the measured solubility data and associated chemical structures directly from Pubchem, using the export function (grouped by substance).

The exported structures were cleaned in MOE (v2008.10; Chemical Computing Group) using the default settings for the Wash command and a set of constitutional and topological descriptors were evaluated. In addition we evaluated physicochemical descriptors (polar surface area, H-bond acceptor and donor counts, $\text{Log } P$ and $\text{Log } D$) using ACDLabs PhysChem Batch (v12.01). Finally we also evaluated a number of custom descriptors that we describe in more detail in the following sections. Importantly, the results from this study are publically available via the internet to allow researchers access to this valuable data set (<http://assay.nih.gov/assay/solubility/>).

3. Results and discussion

Figure 1 summarizes the distribution of measured solubility values and the breakdown of the dataset by the solubility classes ranging from <5, 5–10, 10–15 $\mu\text{g/mL}$, and so on [all data groupings are closed to the lower number (i.e., 5 $\mu\text{g/mL}$ to the lowest value below 10 $\mu\text{g/mL}$)]. Based upon the PubChem definitions, the majority of compounds are moderately soluble (39,301 or 67.9%) followed by a large percentage of low soluble compounds (17,574 or 30.4%). The high soluble group constitutes just 1.7% of the entire dataset and only 75 compounds registered solubility greater than 75 $\mu\text{g/mL}$ (note: the upper confidence limit is affected by molecular weight and artifacts are common for highly soluble, low-molecular weight agents). To ascertain the diversity of the dataset, we considered the distribution of the compounds in a seven-dimensional physicochemical space, defined using molecular weight, $\text{Log } P$, fraction of rotatable bonds, H-bond donor and acceptor counts and the topological polar surface area (TPSA) [physical descriptors and calculated properties were determined using ACDLabs PhysChem Batch (v12.01) and the number of aromatic rings (our analysis considers fused aromatic rings as a single ring) using in-house code].²⁰ The 7-D descriptor space was reduced to two dimensions using principal components analysis and a density plot of the two principal components is shown in the [Supplementary data](#). The dataset is distributed in a relatively homogenous manner (i.e., no distinct clusters are visible) including moderate outliers at the 'edges' of the dataset. From this breakdown of the

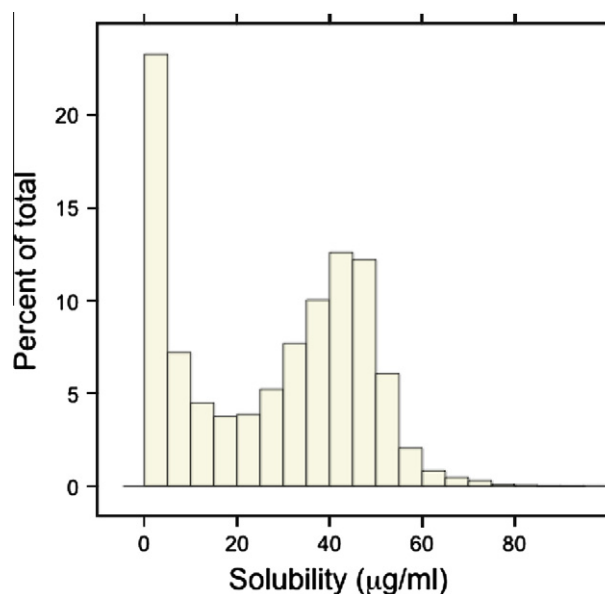


Figure 1. (A) Distribution of measured solubility values [outliers (solubilities >100 $\mu\text{g/mL}$) are not shown in this analysis].

data it was clear that more focused analyses would be required to fully appreciate this data's value.

An important aspect of the data and the subsequent analysis is the use of $\mu\text{g/mL}$ as the unit of solubility. An insightful review of our analysis highlighted that using $\mu\text{g/mL}$ inserts a dependency on molecular weight (MW) into the solubility outcome. To mitigate this we converted the $\mu\text{g/mL}$ values to μM and generated a fine-grained 8-class classification of these values. Overall, the analyses were not significantly different when changing between $\mu\text{g/mL}$ and μM units. The relationship between solubility and MW was the unsurprising exception and a distribution of MW by solubility class as judged using $\mu\text{g/mL}$ and μM provides insight into how units can influence solubility trends (Fig. 2). Importantly, when utilizing μM units there is a modest increase in solubility associated with lower MW compounds. This dogmatic trend is, to a degree, reversed in the analysis that utilizes $\mu\text{g/mL}$ units. All subsequent analyses are generated for both $\mu\text{g/mL}$ and μM units (corresponding analyses are presented in the [Supplementary data](#)).

Analysis of solubility in terms of physical characteristics including H-bond donor count and H-bond acceptor count was accomplished using ACD Labs PhysChem Batch (v12.01) rule settings (for instance, a H-bond donor is described as the sum of OHs and NHs while H-bond acceptors are the number of Ns and Os). We also examined how solubility was influenced by the fraction of rotatable bonds and aromatic fraction which were calculated using MOE. The results of these analyses are shown in [Figure 3](#). The general outcomes are consistent with current dogma on solubility trends; that is, increasing # of H-bond donors and acceptors leads toward a higher fraction of soluble compounds, an increasing fraction of rotatable bonds increases the fraction of soluble compounds, and a lower ratio of nonaromatic to aromatic carbons increases the fraction of soluble compounds. However, despite concurrence with accepted tenets, the data also highlights that within any one-dimensional assessment of the data a significant percentage of compounds with low solubility remains within the highest-value parameter bins. For instance, of the 3143 compounds with eight H-bond acceptors 804 (25.5%) of them have solubilities less than $10 \mu\text{g/mL}$. Examination of several commonly relied upon calculated properties, including Log *P*, Log *D*, PSA, and pK_a , for their association with solubility trends are presented in [Figure 3](#). [Note: ACD Labs software was utilized within these

analyses. A comparison of ACD values with ChemDraw Ultra (v10.0) is presented in the [Supplementary data](#).] From these analyses the relationship between solubility and calculated Log *P* and Log *D* (not calculated *c* Log *P* or *c* Log *D*) values underscored the value of these descriptors in predicting aqueous solubility. The relationship between polar surface area (PSA) and solubility also tracked with presumed dogma whereby a general increase in the sum of polar atoms/polar fragments of a small molecule resulted in a trend toward higher solubility. The results linking the calculated pK_a of an agent to its solubility were less obvious. It might be assumed that a pK_a measurement at the extremes might be indicative of moieties with enough acidic or basic character to be deprotonated or protonated, respectively, leading to formal charges that would confer an increase in aqueous solubility. This is not reflected in the data and only a modest trend toward higher solubility as pK_a values rise is found. It should be noted that, like the data with specific physical descriptors, there are significant percentages of highly-soluble and minimally-soluble compounds present in the ranges of specified calculated properties that are ultimately counterintuitive.

An additional analysis that this data set enabled was the investigation of solubility trends associated with compounds that fit the standard requirements for use within fragment-based screening efforts. A multitude of commercial vendors offer fragment libraries that satisfy so-named 'rule of 2.5' compliance by possessing MW < 250 daltons, <3 H-bond donors and <6 H-bond acceptors. Fortunately, nearly 10% of the ~58,000 member library analyzed in this study satisfied this criteria and [Figure 4](#) summarizes the distribution of the solubility values in both $\mu\text{g/mL}$ and μM units (probability density estimates rather than frequencies were used to allow visual comparisons of the two data sets). This analysis demonstrates that fragment-like agents possessed a greater ratio of soluble agents as compared to the nonfragment allotment. This analysis further demonstrated the role that the units play in these analyses as the μM -based analysis realigned the solubility distributions for the two data sets.

While confirming many of the established solubility trends associated with physical and calculated descriptors these analyses represent highly one-dimensional views of this data set. While a single factor like the number of H-bond donors or the calculated Log *D* will play a key role in a chemist's belief that any given

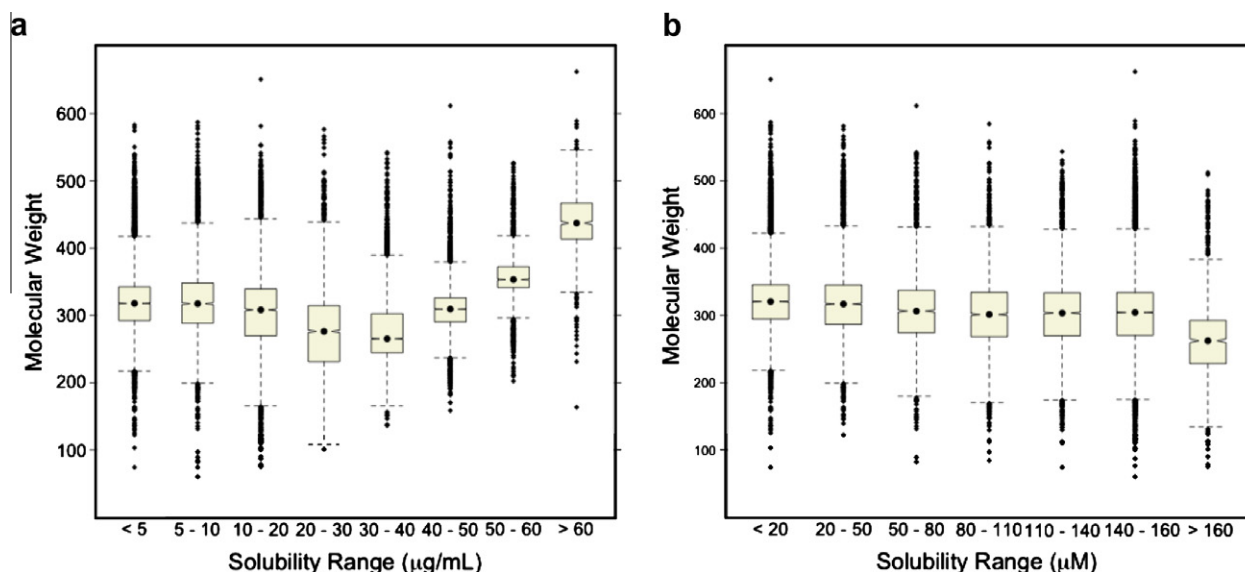


Figure 2. (a) Distribution of MW by solubility class as judged in $\mu\text{g/mL}$ units. (b) Distribution of MW by solubility class as judged in μM units. [To improve visualization we discarded outlying data points with molecular weight >800 daltons.]

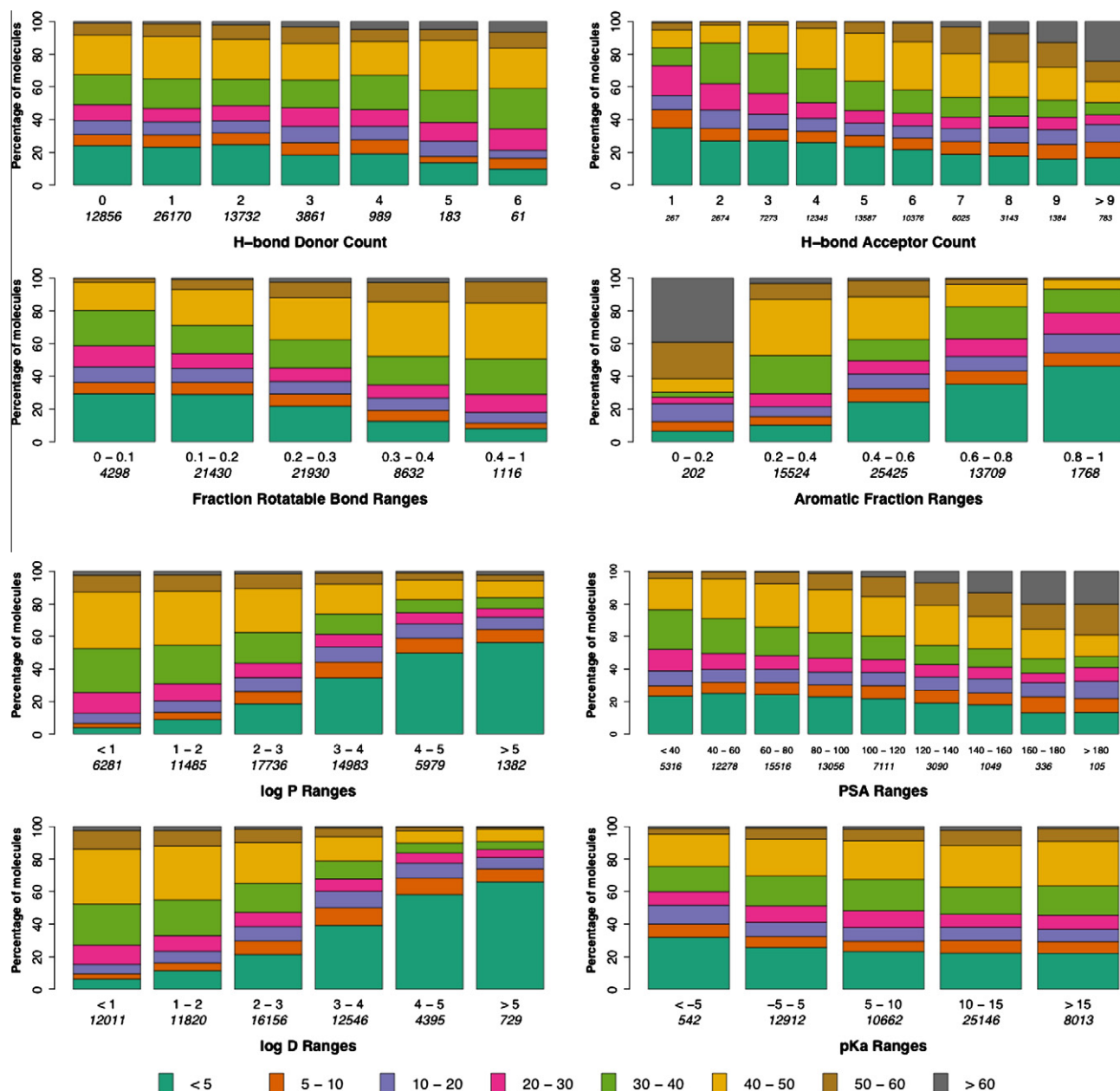


Figure 3. Relationship of solubility ranges as related to specific physical descriptors and specific calculated descriptors. Below each column are the physical descriptor ranges and the number of compounds that fit that descriptor range. The key at the bottom maps the colors to the solubility ranges (specified in $\mu\text{g/mL}$) (a comparative analysis utilizing μM units is presented in the Supplementary data).

compound will be soluble in aqueous media it was important to consider these data more multi-dimensionally. Computational models of solubility provide these types of analyses. Recently, Cheng et al used this dataset to develop a binary classifier using a support vector machine.²¹ This model, like others, involves a relatively large number of structural descriptors and does not necessarily provide a simple rule of thumb that a bench chemist might use to judge solubility. Hill and Young applied a Solubility Forecast Index (SFI) to a similar dataset and found that a molecule with a combined Log *D* and aromatic fraction value less than five will likely possess acceptable aqueous solubility.¹⁷ Using the SFI within our dataset we found accurate classification of the medium and high soluble classes (83.0% and 84.9%, respectively) while the performance declined for the low solubility class (55.6% correct classification). Motivated by the simplicity of this analysis we

considered a variety of combinations of physical and calculated descriptors to obtain simple rules that could be used to guide solubility assessments. With the goal of identifying simple guidelines, we evaluated decision trees for 18,721 2-descriptor combinations from a pool of 194 topological and constitutional descriptors. While a variety of descriptor combinations (such as TPSA and Log *P*) correctly predicted the majority (>90%) of the soluble class, they invariably predicted the bulk of the insoluble compounds as soluble, as well. Given the overlapping distributions of many of these properties for the soluble and insoluble classes for this dataset, this observation is not unexpected. The best performing model that we obtained used a combination of Log *D* and the GCUT_SMR_0 descriptor (which characterizes the distribution of molar refractivity over the molecular structure). While this model exhibited good numerical properties (83% sensitivity and 53%

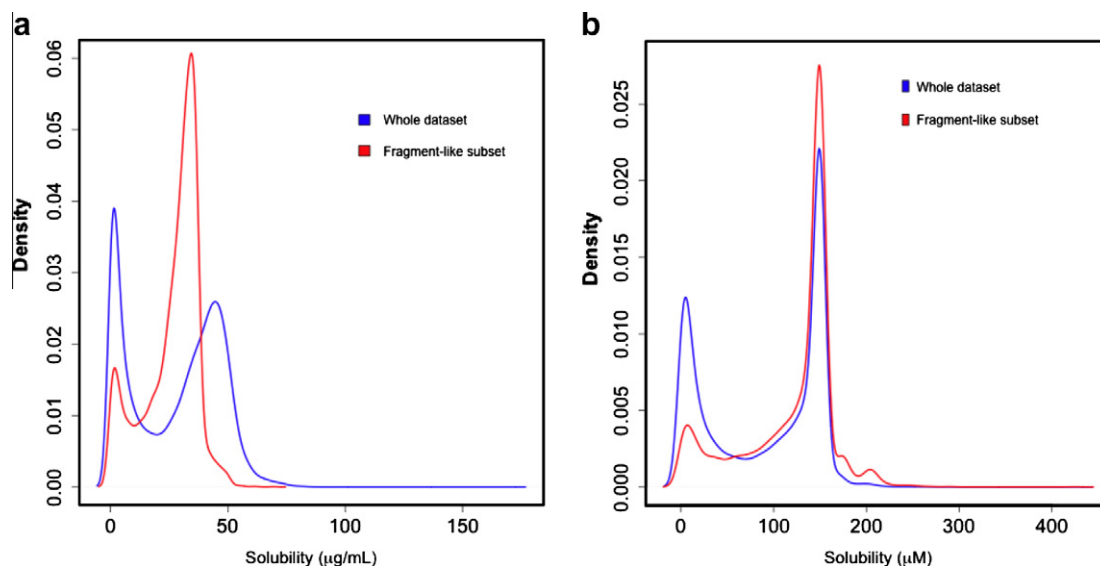


Figure 4. Probability density estimates of the solubility distribution of the subset of compounds satisfying the ‘Rule of 2.5’ in the context of the entire dataset based upon (a) µg/mL units and (b) µM units.

specificity) the use of the GCUT_SMR_0 descriptor is neither intuitive nor easy to calculate. A number of more usable rule sets were found with only slightly poorer performance. Figure 5 displays the top two decision tree models based on a combination of numerical performance and simplicity. A combination of Log *D* and the fraction of sp³ carbons (Fsp3) exhibited a sensitivity of 87.3%, a false positive rate of 12.7% and a specificity of 50.1%. The second utilizes a combination of Log *D* and the number of aromatic atoms (Naro). This decision tree resulted in a model with a sensitivity of 88% and a specificity of 45%; however, this model performed poorly within the insoluble class. The best performing models, in general, made use of Log *P* or Log *D* followed by carbon hybridization. It is important to note that the rules summarized in Figure 5 were developed utilizing the ACD Labs implementation of Log *D*.

Among the most interesting aspects of this profile were the small molecule clusters that appear to be ‘rule-breakers’ in terms of their solubility measurements. For the purposes of this analysis, the ‘rules’ are associated to current dogma on solubility: (1) higher solubility is associated with lower Log *P*/Log *D* values; (2) higher solubility is associated with increasing number of HBD and HBA; (3) higher solubility is associated with lower AF; (4) higher solubility is associated with higher PSA values. An analysis of these agents presented several fascinating findings (Table 1 and Table 2).

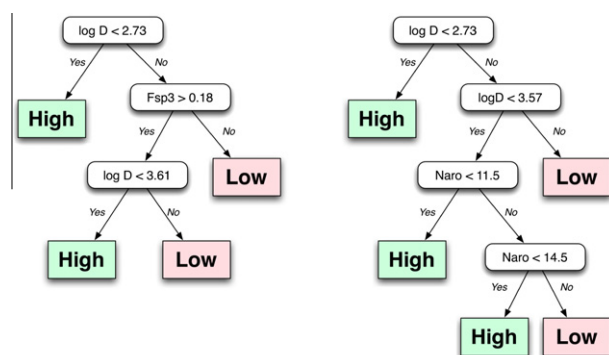


Figure 5. The decision tree model to classify small molecules as having high or low aqueous solubility based on calculated Log *P* values and the ratio of sp³ hybridized carbons.

Importantly, the differences associated with µg/mL (Table 1) versus µM (Table 2) values played a major role in this analysis. Foremost, when examining the role of MW in determining solubility using µg/mL values there is a trend that suggests an increase in solubility accompanies an increase in MW. This finding represents a major divergence from the accepted dogma on trends in solubility. However, this trend is reversed when analyzing the data utilizing µM values.

Other trends included the role that Log *P* and Log *D* play in defining the solubility associated with molecules with either high or low aromatic fractions (a good surrogate of the ratio of sp² and sp³ hybridized carbons). The trend associating a lower Log *P* and Log *D* value was a major determinant in correcting for ‘rule-breaking’ values in the HBD, HBA and PSA analyses as well. A high value for aromatic fraction played a significant role in the ‘rule-breaker’ result associated with low-soluble compounds that maintained a HBD value >5.0. The relative sample sizes associated with HBD’s underscores a major bias in this compound collection toward compounds with few HBD’s. Another interesting result from this analysis was the low values for PSA associated with compounds with a HBA value below 1.0. Agents with a Log *P* value >5.5 that had high solubility possessed a significantly lower Log *D* value and an increase in PSA value. Holding PSA values the same (either high or low) offered a glimpse into this property’s role in governing solubility. In these analyses, Log *P* and Log *D* values were shown to play a significant role in defining the soluble versus insoluble trends while other descriptors maintained relatively close values.

As this library represents a subset of the NIH MLSMR this data set offers a unique opportunity to relate solubility to the results of numerous assays that are published in the PubChem database. Frequent reports and studies decry certain molecular scaffolds as being inappropriate for HTS settings and suggest a general set of rules for small molecules that are likely to be ‘frequent hitters’ within a multitude of assays. Assay promiscuity has been linked to compounds containing reactive ‘warhead’ moieties, agents that are autofluorescent, inhibitors of reporter constructs and even cytotoxic agents.^{22–24} Another major source of compound oriented assay interference is the colloidal aggregation phenomenon.²⁵ We have previously examined the aggregation results for a large set of screening results in a thiol protease assay and a β-lactamase

Table 1

Comparative analysis of putative 'rule-breakers' as judged by selected physicochemical properties.

Descriptor	Sample size	Solubility $\mu\text{g/mL}^a$	Molecular weight	Aromatic fraction	Log P	Log D	H-bond donor	H-bond acceptor	PSA
AF >0.7 Low Solubility	1572	0.42	305.32	0.78	3.85	3.59	1.04	4.18	66.15
AF >0.7 High Solubility	134	52.96	334.84	0.75	2.68	1.19	1.07	4.39	72.16
AF <0.2 Low Solubility	17	0.44	262.55	0.01	1.91	0.95	1.47	4.94	75.51
AF <0.2 High Solubility	211	59.67	395	0.11	2	1.38	1.49	7.65	105.8
HBD >5.0 Low Solubility	4	0.38	359.98	0.73	2.41	2.36	5	7.25	137.2
HBD >5.0 High Solubility	35	60.93	399.19	0.4	1.35	−0.2	5.57	9.26	164.4
HBD <1.0 Low Solubility	3949	0.48	315.17	0.61	3.71	3.46	0.63	4.37	67.13
HBD <1.0 High Solubility	3578	54.97	365.52	0.41	2.32	1.79	0.69	6.02	82.71
HBA >10.0 Low Solubility	32	0.42	377.11	0.52	2.77	2.56	2.28	10.28	144.9
HBA >10.0 High Solubility	292	64.58	441.45	0.36	1.66	0.77	2.33	10.46	151.4
HBA <1.0 Low Solubility	44	0.59	241.08	0.69	4.25	4.08	0.27	1	24.03
HBA <1.0 High Solubility	14	53.79	277.47	0.62	2.41	1.47	0.07	1	17.12
Log P >5.5 Low Solubility	169	0.46	337.75	0.59	5.94	5.03	0.78	3.85	63.14
Log P >5.5 High Solubility	31	58.19	405.32	0.45	5.84	3.05	1.55	6.03	99.95
Log P <−0.5 Low Solubility	3	0.57	248.04	0.41	− 1.31	−2.14	1.33	5.33	79.24
Log P <−0.5 High Solubility	117	55.47	343.16	0.38	− 1.41	−2.83	1.62	6.86	97.1
PSA >115 Low Solubility	342	0.44	345.85	0.55	3.04	2.58	1.87	7.32	129.4
PSA >115 High Solubility	1335	59.13	402.54	0.41	1.92	0.77	2.09	8.26	137
PSA <41 Low Solubility	584	0.55	292.78	0.63	4.2	4.13	0.45	2.68	30.58
PSA <41 High Solubility	253	54.03	313.44	0.49	2.74	1.71	0.48	3.06	29.66

^a Values are listed as the mean. AF = aromatic fraction, HBD = hydrogen bond donor, HBA = hydrogen bond acceptor, PSA = polar surface area ($1.0 \times 10^{-20} \text{ m}^2$).**Table 2**

Comparative analysis of putative 'rule-breakers' as judged by selected physicochemical properties.

Descriptor	Sample size	Solubility μM^a	Molecular weight	Aromatic fraction	Log P	Log D	H-bond donor	H-bond acceptor	PSA
AF >0.7 Low Solubility	1574	1.41	307.72	0.78	3.86	3.59	1.05	4.21	66.83
AF >0.7 High Solubility	422	162.59	267.61	0.77	2.31	0.77	1.19	3.74	61.81
AF <0.2 Low Solubility	20	1.88	305.2	0.04	2.38	1.69	1.35	5.4	87.5
AF <0.2 High Solubility	317	166.7	245.63	0.01	1.05	−0.69	1.48	4.38	67.29
HBD >5.0 Low Solubility	4	1.07	358.98	0.73	2.41	2.36	5	7.25	137.2
HBD >5.0 High Solubility	41	166.62	296.11	0.35	0.61	−1.67	5.56	7.73	140.5
HBD <1.0 Low Solubility	4006	1.55	318.43	0.6	3.71	3.45	0.63	4.44	68.05
HBD <1.0 High Solubility	3650	157.91	290.11	0.44	2.16	1.06	0.72	4.46	64.89
HBA >10.0 Low Solubility	39	1.42	379.06	0.52	2.76	2.41	2.31	10.23	143.9
HBA >10.0 High Solubility	42	155.39	404.7	0.41	1.99	−1.27	3.29	10.52	164.7
HBA <1.0 Low Solubility	36	2.25	247.36	0.68	4.34	4.14	0.25	1	23.65
HBA <1.0 High Solubility	64	189.94	222.41	0.49	2.02	0.79	0.55	1	15.97
Log P >5.5 Low Solubility	176	1.4	343.63	0.59	5.98	5.05	0.79	3.98	65.67
Log P >5.5 High Solubility	25	151.16	349.51	0.49	5.89	2.41	1.4	5.08	90.1
Log P <−0.5 Low Solubility	3	2.27	248.04	0.41	− 1.31	−2.14	1.33	5.33	79.24
Log P <−0.5 High Solubility	246	177.95	239.47	0.39	− 1.54	−3.49	1.64	4.72	68.35
PSA >115 Low Solubility	379	1.43	350.84	0.54	3.06	2.58	1.87	7.35	129.7
PSA >115 High Solubility	619	151.56	335.06	0.43	1.75	−1.16	2.32	7.16	132.9
PSA <41 Low Solubility	562	1.82	296.08	0.63	4.21	4.13	0.45	2.71	30.75
PSA <41 High Solubility	796	175.43	257.22	0.45	2.25	1.25	0.52	2.71	28.39

^a Values are listed as the mean. AF = aromatic fraction, HBD = hydrogen bond donor, HBA = hydrogen bond acceptor, PSA = polar surface area ($1.0 \times 10^{-20} \text{ m}^2$).

assay.^{26,27} While individual compound solubility has long been thought to play a role in the aggregation phenomenon it has yet to be directly correlated to assay results in large scale profiles. In order to investigate the role that solubility has on the frequency of putative actives across a large range of assays and what role solubility plays in the aggregation phenomenon we examined our results in the context of numerous assays reported in PubChem including an assay designed to identify aggregation events.

To identify cross-assay activity, we downloaded the data for all PubChem bioassays (462,437 assays in PubChem as of October 2010) and defined the cross-assay activity score (CAAS) as the number of times a compound was marked as 'active' divided by the number of assays in which it was tested. In this analysis a completely promiscuous compound (active in all assays it was tested in) would have a CAAS of 1.0 while a completely inactive compound would have a CAAS of 0.0. The results are shown in Figure 6. While there are a few molecules that appear to be quite promiscuous (CAAS greater than 0.25), the mean score for each group is generally less than 0.02. For the most soluble group (>60 $\mu\text{g/mL}$), the

bulk of the molecules do not exhibit significant cross-assay activity. Unsurprisingly, the least soluble group (<10 $\mu\text{g/mL}$) tends to exhibit a larger fraction of molecules with higher scores (mean = 0.012 and 3rd quartile = 0.019). While this result highly suggests that low solubility contributes to promiscuous activity it clearly shows that other factors play a role in defining a compound as a 'frequent hitter'.

To determine the agents within this profile that are putative aggregators we performed a quantitative high-throughput screen (qHTS)²⁸ of a recent version of the MLSMR (320,098 compounds) using the aforementioned β -lactamase assay (assay details can be found in Refs. 18,27). To identify aggregation-prone library members the assay is run in the presence and absence of detergent and of the 320,098 compounds screened against β -lactamase under both conditions 50,423 compounds were also measured in the current solubility assay. It should be noted that this assay does not detect the physical presence of aggregates but is a proxy for aggregation-based enzyme inhibition. To identify putative aggregators we considered compounds that exhibited complete

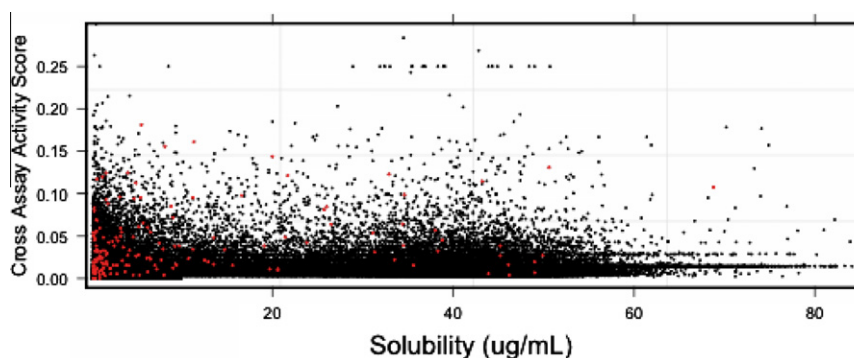


Figure 6. A plot of cross assay activity score versus solubility. 165 molecules are highlighted in red corresponding to those identified as putative aggregators in a previously validated β -lactamase screen.

concentration-response curves [curve classes -1.1 , -1.2 , -2.1 , -2.2 in the qHTS curve class terminology (see Ref. 28)] with efficacies greater than 50% in the (–) detergent screen and no concentration dependent behavior (curve class 4) in the (+) detergent screen. This selection is more stringent than that originally published by combining the curve class and maximum inhibition restriction into a single constraint. This identified 1592 compounds as exhibiting aggregation-dependent inhibition from the β -lactamase screens, of which 165 had also been measured in the solubility screen (the remaining 1427 aggregators were not tested in the solubility screen). Of the tested samples, 121 compounds had solubilities less than 10 $\mu\text{g/mL}$, 43 exhibited solubilities between 10 and 60 $\mu\text{g/mL}$ and a single compound exhibited solubility greater than 60 $\mu\text{g/mL}$ (aggregating agents are designated by red points in Fig. 6). While these results support the assumption that aggregation is highly related to poor aqueous solubility (Figs. 6 and 7) it does not imply that poorly soluble compounds aggregate. Interestingly, the molecules identified as aggregators are not uniformly promiscuous. This supports the notion that aggregation events are highly assay-condition specific and agents identified as aggregator in one assay may not aggregate in another assay format.

A hopeful outcome from this study is wider use of solubility measurements to guide decision making within specific optimization efforts. Certainly this data will assist computational chemists in deriving new models for solubility prediction. Beyond this, we were interested in asking how this data can be informative for chemists aiming for specific pharmacological and physiological outcomes during particular compound optimizations. As a case study, we examined how the solubility data reported here may (or may not) be utilized in a theoretical optimization effort aiming for an agent that penetrates the blood–brain barrier. Recently, Hitchcock and Pennington published a wide-ranging perspective on structure–brain exposure relationships.²⁹ This review offered several suggestions for ranges of specified physicochemical

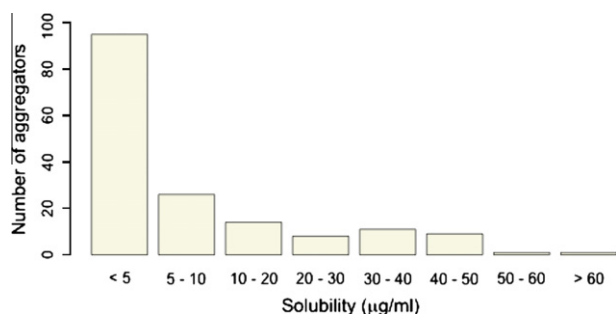


Figure 7. Distribution of the number of aggregators by solubility ranges.

properties to increase the probability of blood–brain barrier (BBB) penetration. Mean values of the top 25 CNS related drugs for PSA (47), H-bond donors (0.8), $c\text{Log } P$ (2.8), $c\text{Log } D_{\text{pH } 7.4}$ (2.1), and MW (293) were also provided. We examined the solubility values of the subset of our collection that matched the above five parameters within a narrow range. There were 667 compounds that satisfied all criteria and the results are shown in Figure 8. Interestingly, a large proportion of these agents had kinetic solubility values between 40 and 50 $\mu\text{g/mL}$. The relevancy of these results are unknown as kinetic solubility may play only a small role in passive diffusion across the BBB. Nonetheless, these results certainly challenge the common misconception that to attain BBB penetration a compound should possess modestly low solubility.

4. Conclusion

In this study we have examined the kinetic solubility for a large chemical library and examined the data in a variety of ways. Foremost, commonly used physicochemical properties were correlated to solubility outcomes and generally confirmed the existing dogma in terms of solubility relationships. However, the data also made it clear that chemists cannot rely solely on the predictive capacity of physicochemical trends to assure that their libraries and optimization efforts result in soluble compounds. The role that solubility plays in HTS settings also highlighted that a lack of aqueous solubility contributes to the phenomenon of ‘frequent hitters’ but is not the only determining factor. Further, a lack of kinetic solubility was

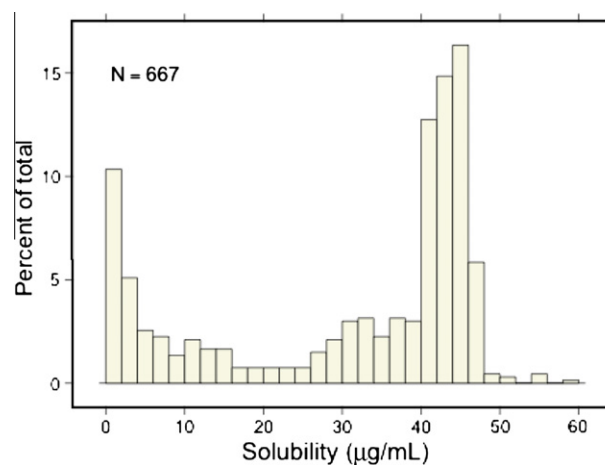


Figure 8. The solubility results for 667 compounds that retain PSA values between 42 and 52, between 0 and 2 H-bond donors, a $\text{Log } P$ value between 1.8 and 3.8, a $\text{Log } D$ value between 1.1 and 3.1 and a molecular weight between 273 and 313.

found to be a primary contributor to the colloidal aggregation phenomenon. The public availability of this data set is intended to allow researchers access to a uniform training set in order to create new computational tools for solubility prediction. In addition, it is hoped that the lessons from this study will add to chemist's already substantial intuition regarding structure property relationships. These results certainly concur with other reports in highlighting the importance of solubility in all facets of the drug discovery process.

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.bmc.2011.05.005](https://doi.org/10.1016/j.bmc.2011.05.005).

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