Development of QSAR models for in silico screening of antibody solubility

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ABSTRACT

Although monoclonal antibodies (mAbs) have been shown to be extremely effective in treating a number of diseases, they often suffer from poor developability attributes, such as high viscosity and low solubility at elevated concentrations. Since experimental candidate screening is often materials and labor intensive, there is substantial interest in developing in silico tools for expediting mAb design. Here, we present a strategy using machine learning-based QSAR models for the a priori estimation of mAb solubility. The extrapolated protein solubilities of a set of 111 antibodies in a histidine buffer were determined using a high throughput PEG precipitation assay. 3D homology models of the antibodies were determined, and a large set of in house and commercially available molecular descriptors were then calculated. The resulting experimental and descriptor data were then used for the development of QSAR models of mAb solubilities. After feature selection and training with different machine learning algorithms, the models were evaluated with external test sets. The resulting regression models were able to estimate the solubility values of external test set data with R^2 of 0.81 and 0.85 for the two regression models developed. In addition, three class and binary classification models were developed and shown to be good estimators of mAb solubility behavior, with overall test set accuracies of 0.70 and 0.95, respectively. The analysis of the selected molecular descriptors in these models was also found to be informative and suggested that several charge-based descriptors and isotype may play important roles in mAb solubility. The combination of high throughput relative solubility experimental techniques in concert with efficient machine learning QSAR models offers an opportunity to rapidly screen potential mAb candidates and to design therapeutics with improved solubility characteristics.

Introduction

Monoclonal antibodies (mAbs) have clearly emerged as the dominant class of biotherapeutics and the recent approval by the Food and Drug Administration of a mAb for Alzheimer’s and antibody therapies for COVID-19 will likely dramatically increase the scale required for mAb biomanufacturing. Importantly, the development of platform processes have enabled significant reductions in the time required to proceed from discovery to production. Despite these advances, antibody candidates often suffer from ‘developability’ challenges, where ‘developability’ is a metric for evaluating the potential for successful development of a stable, safe, and efficacious product. This is often accessed by measuring colloidal stability correlated properties such as aggregation propensity, solubility, and viscosity. Experimental techniques such as cross-interaction chromatography (CIC), standup monolayer adsorption chromatography(SMAC), affinity-capture self-interaction nanoparticle spectroscopy (AC-SINS), clone self-interaction nanoparticle spectroscopy (CSI-BLI) or dynamic light scattering (DLS) have been used to directly or indirectly evaluate the colloidal stability of candidates. However, the direct measurements of properties such as solubility and viscosity often require elevated mAb concentrations and are still labor and material intensive, which makes them impractical in early development.

The development of in-silico approaches to estimate properties related to developability could substantially accelerate the process of screening potential lead candidates, reducing the amount of materials required. A number of approaches have been examined for developing in silico tools evaluating antibody developability. Coarse-grained simulations have been applied for predicting antibody viscosity. Single molecular parameters, such as charge distribution and hydrophobic index have been shown to be correlated with viscosity and chemical stability. Raybold et al. reported on the use of a small set of molecular descriptors to provide guidelines for developability. Sequence compositions and residue-based descriptors have also been applied for developing methods to access self-association or colloidal stability and to assess aggregation propensity and viscosity. In addition, multiple physiochemical descriptors obtained from either sequence or three-dimensional structures have been used in models of assessing antibody oxidation risk, overall hydrophobicity, and solubility.
Although this prior work was useful for projecting mAb solubility and viscosity, most of these methods were developed using a relatively small set of antibodies or variants with a relatively small number of preselected descriptors. It would be extremely useful to develop in silico tools that were able to model the developability behavior of a wider range of mAb candidates. Quantitative structure activity relationship (QSAR) models can provide a unique link between the solute activity being modeled (e.g., solubility) and the important molecular properties of the solutes. Widely used in the small molecule drug development space, QSAR models have also proven useful for evaluating the behavior of large biomolecules, particularly in chromatographic applications. Robust QSAR models have been developed for a wide range of proteins, in a myriad of chromatographic media. Recently, QSAR models have also been used to estimate protein diffusion coefficients in formulation applications.

For this report, we developed a QSAR based in silico screening strategy for modeling relative mAb solubility. A previously developed experimental high-throughput mAb solubility screening assay was used to determine the solubilities of a relatively large set (111) of diverse antibodies in histidine buffer, pH 6.0. A broad range of in house and commercially available molecular descriptors were then calculated based on antigen-binding fragment (Fab) homology models and feature selection was carried out to determine the key descriptors for use in the models. Quantitative regression and qualitative classification models were then trained with different machine learning algorithms and the top models were shown to be effective in screening mAb on relative solubility. Finally, interpretation of the models was carried out to provide mechanistic insights into the mAb solubility behavior.

Results

Antibodies relative solubility distribution

A dataset of 111 antibodies composed of diverse molecules from various mAb discovery platforms, and different antigen targets was curated. The solubility of antibodies in 10 mM histidine buffer were determined by high-throughput PEG-induced precipitation. As described in the methods section, the PEG experiments were carried out and the percentage of PEG that resulted in an abrupt decrease in absorbance (i.e., the onset) was used as a surrogate for ranking solubility. The values of PEG percentages were then normalized on a zero to one scale using a Min-Max normalization based on the solubilities of two control molecules. As shown in the histogram of the normalized solubilities (Figure 1), while the solubility of the 111 molecules in the set were distributed across this scale, 34 of the mAbs had high solubility (≥1.0). In addition, 2 mAbs exhibited lower solubilities than the low control and 1 mAb had a higher solubility than the high control. The range of solubility behavior along with the diversity of this mAb set enabled us to develop models for a wider range of mAbs than has been previously reported.

Regression models for antibody solubility

Based on the solubility data of all 111 mAbs shown above, the first regression model was developed following the QSAR model development workflow described in the methods section. The randomly selected training set contained 91 molecules and the remaining 20 molecules were used as the test set. Multiple rounds of feature selection were first carried out, after which different algorithms were tested on the resulting feature set. Validation was then carried out using 10 times five-fold cross-validation and 50 rounds of Y-scrambling. This workflow of feature selection, algorithm selection and model validation were carried out in an iterative manner and the best performing model for this data set was found to be a five-component partial least square model using 10 molecular descriptors. A comparison between the experimentally determined solubilities and the projected values are presented in Figure 2a. The coefficient of determination R² for the training and test sets were 0.883 and 0.811, respectively. The average score of the 10 times fivefold cross-validation was 0.833, which was close to the training and test set performance, indicating decent model robustness. Finally, the root mean squared deviations (RMSD) were 0.114 and 0.168 for the training and test set (dotted line).

In Figure 2a, most of the outliers had experimentally determined solubilities of 1.0, which was the upper bound of the assay and includes mAbs with solubility values equal to or above this value. Further, we were primarily interested in identifying mAbs with lower solubilities, which would represent potentially problematic candidates. For both of these reasons, we developed a second QSAR regression model that excluded the highly soluble mAbs (Solubilities ≥ 1.0). The resulting dataset included 75 points with 63 of them used in the training set and 12 as the test set. Following a similar development procedure, the best performing model was found to be a four-component PLS model using 17 molecular descriptors (Figure 2b). The coefficients of determination for
the training and test sets were 0.881 and 0.847, respectively. In addition, the average score for the cross-validation was 0.772. Finally, the RMSD for the training and test sets (dotted line) were 0.083 and 0.114, respectively.

The descriptors selected for these two QSAR regression models are presented in Table S1. To further evaluate model robustness and to assure minimal overfitting of the data, 50 rounds of Y-scrambling were carried out for both models, as described in the methods section. As can be seen in Figure S1 and S2, models developed with the scrambled data were consistently unable to project the experimental values, suggesting that there was minimal overfitting in our QSAR models (Figure S1, S2). Furthermore, the original models clearly outperformed the “scrambled” models both with respect to the $R^2$ and cross-validation score values.

**Classification models for antibody solubility**

In addition to QSAR regression models, we explored classification models that could identify drug candidates with potential solubility challenges. In this study, we developed both three- and two-class classification models. As was done with the regression model developed with the entire data set, 91 mAbs were used for the training set and 20 mAbs were set aside for the external test set. Prior to training, the mAbs were divided into low, medium, and high solubility ranges for the three-class model and low and high solubility ranges for the binary model. To ensure that the training data was evenly distributed, cutoffs between the classes were determined to be 0.38 and 0.88 for the three class and 0.58 for the binary model. Support vector machine with a linear kernel was applied for both models. The confusion matrices for the training and test sets for the three- and two-class models are presented in Figure 3. In addition, the performance criteria of these models are summarized in Tables 1 and 2 and the descriptors selected for these models are presented in Table S1. As can be seen in the Table 1, for the three-class model, while the training set had high accuracy (0.88), the test set was less accurate (0.7) with 4 of the 5 misprojections occurred in the medium class. The accuracy of this model was also indicated by the cross-validation score, which was determined to be 0.85. The F1-score is a useful metric for evaluating model performance by aggregating the precision and recall scores. While the F1-scores for all three classes were over 0.8 for the training set, a low F1-score of 0.29 was observed for the medium class in the test set.

Although the three-class model performed relatively well, the low correlation observed for the medium class motivated the development of a binary model. As shown in Table 2, the binary model had overall improved accuracies of 0.92 and 0.95 for the training and test sets, respectively. Importantly, only seven of 91 instances were misprojected in the training set and only one misprojection was observed for the testing set. The cross-validation score for the binary classification model was 0.9 and the F1-scores were over 0.90 for all classes in both the training and test sets, indicating high model accuracy.

**Identification of key features related to mAb solubility**

After developing various QSAR models for estimating mAb solubilities, descriptor analysis was carried out to better understand the molecular basis. One metric for evaluating the significance of features is their coefficient in the linear regression models, where higher absolute values indicate their relative importance in determining solubilities and positive or negative characteristics suggest their correlation or inverse correlation with solubility. Another important metric for features selection in the PLS models is the variable influence on projections (VIP) score. This parameter aggregates projections of the feature weights onto latent hyper planes. Features that have VIP values over 1 are considered important in the model whereas features
with values less than 1 have a minor effect on the projections.\textsuperscript{38,39,41} A comparison of the feature coefficients and their VIP scores from the regression model based on all the mAb data is presented in Figure 4, which includes a horizontal dashed line for better identification of positive/negative characteristics of the coefficients. In Figure 4, the top four descriptors that had VIP values equal or greater than one were: 1) isotype identifier, 2) average electrical potential of the top 25% strongest negative clusters(EPL\_str), 3) protein isoelectric point (pI) based on three-dimensional structure(pro\_pl\_3D) and 4) the charge symmetry parameter of the variable region(FvCSP). The other descriptors in the left side of the figure all correspond

| Class    | Precision | Recall | F1-score | Support |
|----------|-----------|--------|----------|---------|
| Low      | 0.90      | 0.81   | 0.85     | 32      |
| Medium   | 0.79      | 0.82   | 0.81     | 28      |
| High     | 0.94      | 1.00   | 0.97     | 31      |
|          | overall accuracy |        | 0.88     | 91      |

| Class    | Precision | Recall | F1-score | Support |
|----------|-----------|--------|----------|---------|
| Low      | 0.75      | 0.86   | 0.80     | 7       |
| Medium   | 0.50      | 0.20   | 0.29     | 5       |
| High     | 0.70      | 0.88   | 0.78     | 8       |
|          | overall accuracy |        | 0.70     | 20      |

Figure 3. Confusion matrix for classification models: A) training set for three-class model; B) test set for three-class model; C) training set for binary model; D) test set for binary model.
to hydrophobic descriptors, which are described in the discussion section. The definitions and coefficients of all the selected descriptors were included in Table S1.

To further investigate the relationship of these descriptors to mAb solubility, we mapped the solubility data onto the key features identified. Figure 5 presents a scatter plot where each data point corresponds to a specific mAb and the color coding represents their normalized solubilities (described in the methods section). Figure 5a projects this data onto the pro_pI_3D and FvCSP scores and Figure 5b projects it onto the EPL_str and FvCSP values. Interestingly, clustering behavior was observed for the high solubility mAbs in both figures and dashed lines are included to facilitate this visualization. In Figure 5a, high solubility mAbs were observed to cluster in the top-right quadrant, whereas in Figure 5b these mAbs clustered in the bottom right. We explore the implications and applications of these findings in the discussion section below.

Figure 4. Variable influence on projection score (VIPs) versus descriptor coefficient in model. Labeled descriptors are: 1) ‘isotype’; 2) ‘EPL_str’; 3) ‘pro_pl_3D’; 4) ‘FvCSP’; 5) ‘aroores_nstr’; 6) ‘aroH_nstr’; 7) ‘aromatic_acid_num’; 8) ‘aromatic_num’; 9) ‘pro_patch_hyd’; 10) ‘Basic_max’.

Figure 5. Key Descriptor 1 value versus key descriptor 2 value scatter plot: A) pro pl 3D vs FvCSP; B) EPL_str vs FvCSP. Color bar represents solubility value of each molecule. Green represents high solubility, rose represents low solubility.

Table 2. Summary of performance metrics for binary model.

| Class | Precision | Recall | F1-score | Support |
|-------|-----------|--------|----------|---------|
| Low   | 0.93      | 0.91   | 0.92     | 46      |
| High  | 0.91      | 0.93   | 0.92     | 45      |
| Train | overall accuracy | 0.92 | 91 |
| Test  | Class | Precision | Recall | F1-score | Support |
| Low   | 1.00 | 0.89 | 0.94 | 9 |
| High  | 0.92 | 1.00 | 0.96 | 11 |
| Test  | overall accuracy | 0.95 | 20 |
**Impact of mAb isotypes**

Another interesting finding from this study was that ‘isotype identifier’ was found to be an important factor impacting mAb solubilities at pH 6.0. Different isotypes were included in this investigation; the 111 mAbs included 66 IgG4s, 42 IgG1s, and 5 IgG2s. We carried out a comparative analysis of the solubility behavior of the IgG1 and IgG4 mAbs, the dominant isotypes, and their correlations to key molecular descriptors. As can be seen in Figure 6a, the isotypes exhibited significant differences in their solubilities. The IgG1 mAbs (Blue) had many molecules with high solubilities along with a relatively small number of IgG1s with a range of lower solubilities, reaching into the low soluble region. On the other hand, the IgG4s exhibited moderate solubilities (0.2–0.4) with only a small number achieving high solubility. Interestingly, the solubility data for these two isotypes had very different correlations to the key molecular descriptors EPL_str and pro_pI_3D of the Fabs (Figures 6b-c). While the solubility of IgG1s had a strong linear correlation (Pearson coefficient $R^2 = 0.85$) with the pro_pI_3D of the Fabs, the solubility data of IgG4s had a much weaker correlation (Pearson coefficient $R^2 = 0.62$). The results with the EPL_str (negative electrical potential) descriptor were even more striking. The solubility of the IgG1s exhibited a very high inverse correlation with EPL_str ($R^2 = 0.90$). On the other hand, the IgG4 solubility data was moderately correlated with a $R^2$ of 0.65. The electrostatic potential maps for the IgG1 and IgG4 Fcs at pH 6 were quite different, as shown in Figure 7. These differences in solubility behavior, correlations, and surface properties for the two isotypes are discussed below.

**Discussion**

In this study, we developed both regression and classification models using mAbs exhibiting a range of solubility behavior obtained from discovery campaigns at Lilly. Compared to previously described methods in the literature, our study is more expansive than previous work with respect to both the range of mAbs and the molecular descriptors used for model generation. Importantly, the QSAR regression and classification

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**Figure 6.** Behavior differences between IgG1s and IgG4s: A) Histogram of solubility distribution; B) scatter plot of pro_pI_3D vs solubility; C) scatter plot of EPL_str vs solubility. In all three plots, red represent IgG1, and indigo represent IgG4.
models for mAb solubilities developed in this work were shown to have both high accuracy and robustness, which may provide useful in silico screening tools for evaluating biomanufacturability. Further, the descriptor analyses carried out in this work have provided new information about molecular properties that may be important to consider when developing mAbs with improved solubility.

Regression models were first developed using experimental mAb solubility data obtained in histidine buffer at pH 6. The solubilities of 111 mAbs were determined and the data were normalized based on two control molecules as described in the methods section. It should be noted that the PEG precipitation assay used in this study was designed for high-through screening of mAbs during drug development. While it serves as a surrogate for relative solubility, it is not a direct measure of mAb solubility. Two regression models were developed, model A (shown in Figure 2a), which included all 111 mAbs, and model B (shown in Figure 2b) with a smaller set of 75 mAbs that exhibited low to moderate solubilities. Model B was developed to focus on mAbs that may have solubility issues. Both models performed in a similar manner with high R² values for the training, cross-validation, and test sets. In addition to regression models, the full solubility data set was used to generate three-class and binary classification models. As can be seen in Figures 3 A and 3b and Table 1, the three-class model resulted in satisfactory performance for the training set with an overall accuracy of 0.88 and an averaged cross-validation score of 0.85. For the high solubility class, all 31 values were correctly projected and a F1 score of 0.97 was achieved. The results with the test set were less accurate. Although only 1 misprojection occurred for the low and high solubility classes, 4 of 5 solubility values were misprojected for the medium test set. This resulted in a low F1-score of 0.29 for the medium class and a 0.70 overall accuracy for the test set. One explanation for the results with the medium test set could be the lack of balance of the solubility data distribution. In order to generate the three-class model, we distributed the data such that there were comparable numbers of mAbs in the training sets for these three classes. This necessitated less well-defined cutoffs for the medium class, with a bias toward higher solubility values. While a variety of cutoffs and distributions of data were examined for the three class models, they consistently underperformed for the medium class projections.

In order to create a more meaningful distribution and cutoff between the solubility classes, we developed a binary model. This enabled us to have similar numbers of data in the classes as well as a solubility transition, 0.5, that reflected the distribution of the data. As shown in Figure 3 c and d and Table 2, the binary classification model achieved excellent projections for the training, cross-validation, and test sets, with overall accuracies of 0.92, 0.89, and 0.95, respectively. Notably, only 1 of 20 data points in the test set was found to be misprojected. In fact, the binary model had superior performance as compared to both the three-class classification model and the two QSAR regression models.

It was also of interest to examine the descriptors selected in the models to provide some insights into the underlying protein–protein interactions influencing the mAb solubilities. Accordingly, we first examined the descriptors selected for regression model A by comparing the VIP scores and the descriptor coefficients (Figure 4). As can be seen, from the 10 features selected for this regression model, four had VIP scores higher than 1.0, reflecting their importance in the model. Interestingly, three of the top four descriptors were related to the charge properties of the Fab region. In agreement with previous work in the literature, the charge symmetry parameter, FvCSP, was found to have a positive coefficient, indicating its correlation with mAb solubility. In addition, the pI determined from the 3D Fab structure (pro_pl_3D) and the average of the top 25% of the negatively charged clusters (EPL_str) were found to be correlated and inversely correlated, respectively, with the mAb solubilities. While ‘FvCSP’ and ‘pro_pl_3D’ were global charge descriptors describing overall charge distribution on the Fab surface, ‘EPL_str’ was a cluster
descriptor describing the strength of negatively charged patches in the Fab region. Since the inclusion of all three of these charge-based descriptors were shown to be very important for model performance, we believe that they worked synergistically to enable the solubility projections. These results are supported by previous observations\textsuperscript{17,18,42} that specific charge–charge interactions involving the Fab can significantly affect the colloidal behavior of mAb solutions and that negatively charged patches on the Fab can reduce mAb solubility.

Previous studies have indicated that hydrophobicity, π–π and π-cation mAb interactions may contribute to mAb viscosity.\textsuperscript{12} In addition to the three charge-related and isotype descriptors, descriptors related to Fab hydrophobicity were also selected for our regression model (Table S1). These included descriptors based on aromatic clusters (5, 6, 8), hydrophobic patches (9), and overlapping hydrophobic and charge cluster-based descriptors (7), where the indicated numbers correlate to the labels in Figure 4. As can be seen in Figure 4, although none of these had VIP scores over 1.0, the absolute values of their coefficients (see the left side of the plot) were comparable to the charge-based descriptors, which made them indispensable contributors to the models. The results indicate that hydrophobic interactions, in concert with electrostatic interactions, are likely playing a role in mAb solubility for the histidine buffer system.

To further understand the relationship between solubility and key descriptors, the mAb solubilities were mapped onto the key charge features described above (Figure 5). As can be seen, some clear clustering behavior was observed for the highly soluble mAbs. Figure 5a which plots the pro pI 3Ds versus the FvCSP values, illustrates that mAbs with high solubilities (green) were clustered in the upper right quadrant with only 7 of the 77 molecules in this region having solubilities less than 0.38 (the cutoff used in the three-class model for low solubility). This suggests that highly soluble mAbs tend to have high Fab pIs (above 7.5) as well as charge symmetry in the variable region (FvCSP > 0). In addition, Figure 5b indicates that high solubility mAbs tended to cluster in the lower right quadrant, where only 6 of 74 mAbs in this region had low solubilities. This indicates that in addition to having charge symmetry in the Fv region, high solubility mAbs also tended to have weaker negatively charged clusters on the Fab surface.

This clear clustering behavior related to this relatively small set of charge-based descriptors can potentially be used to help guide the design of mAbs with higher solubilities. Currently, the cutoffs between quadrants represented by the dashed lines in Figure 5 were determined such that at least 90% of the mAbs within the clustering quadrant had either high or moderate solubilities. These cutoffs can be further tuned based on the accuracy needed for the screening. The clustering behavior shown in Figure 5, indicated that high solubility mAbs tended to have similar charge characteristic in the Fab or variable region.

In contrast to the high solubility data, the mapping of the low solubility mAbs was observed to be more scattered in these plots. Nevertheless, certain patterns could be determined, such as the tendency for low solubility mAbs to have lower Fab pIs, stronger negative charged patches on the Fab surface and less charge symmetry of the Fv region. In order to clearly establish these patterns, we would need to include more low solubility mAbs in our analysis. In addition, the dispersion of the data in this plot could also be due to other factors playing an important role in this low solubility regime beyond these simple charge-based descriptors.

Another important finding of this study was that 'isotype' was identified as the descriptor having the highest VIP score and absolute value of the coefficient (Figure 4), which made it the most influential descriptor in the model. In initial exploratory studies, the Fc features were included in the descriptor pool; however, after feature selection, we consistently obtained one Fc descriptor for each model which was selected as a surrogate for 'isotype identifier'. When more Fc-based descriptors were examined in these models, no obvious improvements were obtained. Accordingly, we used 'isotype identifier' as one of the features and developed Fab homology models that were then used to determine the Fab-based descriptor sets. Interestingly, drastically different solubility behaviors were observed for the different isotypes. As can be seen in Figure 6a, while more than half of the IgG1s had high solubilities (≥1.0) with much small numbers in the lower solubility regimes, the IgG4s had a broad distribution with the peak centered in the moderate solubility region (0.25–0.50).

We also examined the relationship between mAb solubilities and some of the key descriptors identified for the two isotypes. As can be seen in Figure 6b, a high correlation ($R^2 = 0.85$) between solubility and Fab pI was observed for the IgG1s, whereas only a moderate correlation ($R^2 = 0.62$) was found for the IgG4s. A similar result was obtained with the strength of the negatively charged clusters (EPL_str) shown in Figure 6c where IgG1 solubility had a very high inverse correlation ($R^2 = 0.90$) with solubility, whereas the IgG4 solubility was less well correlated ($R^2 = 0.65$). This strong linear correlation between these single charge-based descriptors and mAb solubilities for the IgG1s was quite striking. We hypothesized that the difference in the solubility behavior of the two isotypes may be due to differences in the charge profiles of the Fc regions. Accordingly, we generated electrostatic potential (EP) maps at pH 6 for the Fc regions of the IgG1s and IgG4s used in this study (Figure 7). In these plots, red and blue represent negative and positive charged regions on the protein surface. As can be seen in the figure, the IgG1 Fc was significantly more positively charged than the IgG4 Fc. The IgG1 Fc (ABC) surfaces were mostly covered by contiguous positively charged patches with a relatively small number of discontinuous negatively charged patches. In contrast, IgG4 Fc had a large contiguous negative patch on both the bottom and the back regions of the molecule. These observations are in agreement with the theoretical 3D pls of the IgG1 and IgG4 Fcs, which are 7.59 and 6.61, respectively.

The EP maps also help to explain the strong correlation between Fab pI and mAb solubilities for the IgG1s. The higher the Fab pI, the more positively charged patches will occur on the Fab surface at pH 6. This will produce stronger electrostatic repulsions between the positively charged Fc and Fab surfaces, resulting in higher mAb solubilities. In contrast, the large negative patch on the IgG4 Fc could result in electrostatic attractions between the positively charged FabS and the Fc. This could in turn result in additional interactions occurring
between the regions, which would reduce mAb solubilities and weaken the linear correlations between the charge-based descriptors and the IgG4 solubility. These results are also consistent with a previous observation that the charge profile of IgG1s and IgG4s modulate the differential developability properties of the two isotypes. This analysis of the correlations of the mAb solubilities with charge-based descriptors, in concert with the protein surface plots for the two isotypes, provides potentially useful information for engineering antibody candidates with improved solubility behavior and for selecting appropriate scaffolds for development.

The aim of this study was to develop in silico screening tools of mAb solubility. Using a QSAR modeling approach, regression, and classification models were developed to provide quantitative or qualitative projections of mAb solubilities, and our top models have the potential to be used for screening mAb relative solubility at early discovery. Finally, interpretation of the models was carried out to provide mechanistic insights into the mAb solubility behavior and our results indicated that isotype and Fab charge-based descriptors were important, particularly for the IgG1 mAbs. While this work was successful in generating QSAR regression and classification models for a given set of conditions (normalized solubility data based on a PEG precipitation assay for a given set of mAbs), it would be difficult to extend these particular models to data outside of this set due to the effect of differences in: 1) the biomolecules, 2) the buffer conditions, and 3) the particular biophysical techniques used as a surrogate for solubility. To further improve model performance and applicability to a wider set of molecules and conditions, future work will focus on constantly updating the model as more experimental data becomes available with a wider range of mAbs. In addition, we will extend these efforts to alternative formulation conditions (buffers, pHs, additives) and also develop models for other important biophysical properties related to developability.

Materials and methods

Antibody preparation

Antibodies used in this study were from various discovery campaigns and included IgG1, IgG2, and IgG4 subclasses. Samples were produced internally at Eli Lilly and Company. mAbs were expressed in either transient 293 F or stable Chinese hamster ovary cells. Purification followed typical antibody purification procedures (protein A capture followed by polishing steps). All reagents and excipients were commercially available from Hampton Research, EM Chemicals, JT Baker, Sigma-Aldrich, and/or Mallinckrodt, and were of high grade (>98% purity).

Solubility determination by PEG-induced precipitation with automation

The experimental protocol was adapted from the method reported by Chai et al. and Oeller et al. Solutions of 10 mM histidine, pH 6.0, containing varying polyethylene glycol 3350 (PEG 3350) concentrations (v/v) (from 4 to 36%) were prepared using a Formulator (Formulatrix, Bedford, MA). All the mAb samples were buffer-exchanged and diluted to 1 mg/mL and plated (50 µL/well) onto the wells of 96-well polystyrene, V-bottomed assay plates (Greiner). A Biomek i7 (Beckman Coulter, Indianapolis, IN) liquid handler was used to add 50 µL/well of the PEG stock solutions from the Master Block plate into 96-well polystyrene, V-bottomed assay plates containing the mAb samples. The final composition of each well on the PS plate (in a total volume of 100 µL) contained 50 µg of sample per well, and 10 mM histidine with PEG 3350 concentration ranging from 2% to 18% (v/v) across each column of the plate in 1.5% increment. The assay plates were sealed with clear sealing film (Hampton Research, Aliso Viejo, CA) and then incubated at room temperature on a titer plate shaker (Lab-Line Instruments, Melrose Park, IL) for 24 hours. After incubation, the assay plate was centrifuged at 3500 RPM for 15 minutes at 25°C on an Allegra X12R benchtop centrifuge (Beckman Coulter, Brea, CA) to remove the precipitate. Then, 50 µL/well liquid was transferred from each well into a UV Star 384-well plate using a Biomek i7. After sealing the UV Star plate with clear sealing, the plate was centrifuged for two minutes at 3000RPM to remove air bubbles. Finally, the plate was read on a Tecan Infinite M1000 Pro UV/Vis Spectrophotometer (Männedorf, Switzerland) at 280 nm (with background subtraction at 320 nm). The absorbance data was de-convoluted and plotted in Excel, where the point of abrupt decrease in absorbance was determined. The nearest PEG 3350 concentration (%) that corresponded to the onset of precipitation was defined as Peg onset and used as a surrogate for mAb solubility. The resulted Peg onset was then normalized based on the values obtained with the two internal control molecules as follows:

\[
\text{Sol}_{\text{norm}} = \frac{\text{PEG}_{\text{test}} - \text{PEG}_{\text{low}}}{\text{PEG}_{\text{high}} - \text{PEG}_{\text{low}}} \tag{1}
\]

PEGlow and PEGhigh were the Pegonset for the two control molecules used for each measurement. Of note, for cases where the high solubility control molecule does not have a precipitation onset during the assay, we use the maximum PEG concentration used in the assay. The normalized PEG data of all the mAbs in this study are summarized in Table S3. The representative data of PEG-induced precipitation assay is presented in Figure S3. A four-parameter log-logistic function was applied for curve fitting and is indicated by the solid curves. Pegonset (indicated by the triangles in Figure S3) represents the PEG concentration corresponding to a 5% decrease in the soluble mAb concentration from the initial amount. For mAbs that did not precipitate at the highest PEG3350 concentration employed (36%), we assigned a Pegonset value of 36%. We choose Pegonset over Peg1/2 due to an improved sensitivity in ranking the molecules.

Antibody 3D structure preparations

The Fab structures were determined using the antibody modeler module in Molecular Operating Environment 2019 (MOE) (Chemical Computing Group, Montreal, Canada). Settings used were the “Fine” refinement level, Amber10: EHT force field, and the R-solvation field. To compensate for errors due to protein flexibility, five repeated homology models were developed to create conformation ensembles for each Fab structure.
Calculation of molecular descriptors

Protein molecular descriptors from multiple sources were calculated for all the Fab homology models developed in this work. Global protein descriptors and protein surface patch descriptors were calculated using MOE. FvCSP was calculated according to methods developed by Sharma et al. 16 SCM score was calculated according to Agrawal et al. 26 Custom-designed residue cluster-based descriptors and overlapping clusters descriptors were calculated using the procedure published by Woo et al. 36 PDB2PQR software was used for structure corrections and protonation. 35 Poisson-Boltzmann electrostatic calculations were performed by Adaptive Poisson-Boltzmann Solver (APBS). 46 All the descriptors described above were calculated based on five repeated Fab homology models and the average values of five runs were used for model development, resulting in 310 molecular descriptors for each mAb prior to feature selection.

QSAR model development workflow

The QSAR model development process was implemented using python 3.7.1 on JupyterLab 2.1.5. The total dataset was composed of 111 mAbs solubilities in 10 mM histidine buffer, at pH 6, which were normalized into a 0 to 1 scale using ‘MinMax normalization’ according to the solubilities of two control molecules (high/low). Computational molecular descriptors were standardized such that each had a mean of 0 and a standard deviation of 1 for the entire mAb data set. After data preparations, the dataset was randomly split into a training set (82%) and a test set (18%). Multiple rounds of feature selection were then applied to filter out unrelated features and to remove redundancy within the feature set. Briefly, the initial selection removed features with low variance (p >.85). Secondly, features with high mutual correlations (>85%) was then removed. Finally, L1-based recursive feature selection was applied to remove as many unimportant features as possible while maintaining the model performance. 47 Different algorithms, including partial least square (PLS), support vector machine (SVM), tree-based algorithm, and multilayer perceptron, were evaluated. The average score of 10 five-fold cross-validation and 50 rounds of Y-scrambling 48 were used for model validation and to assure there was no overfitting of the models. Y-scrambling was implemented by keeping the x-dataset (the descriptor set) intact and shuffling the y-dataset (normalized solubility) 50 times. For each ‘scrambled’ dataset, a ‘scrambled’ model was trained using the same algorithm as the original model. The coefficient of determination R² and cross validation square correlation coefficient Q² were then used to access the performance of these resulting models of the scrambled data. The development process of feature selection, algorithm selection, and model validation were carried out in an iterative manner to select the best performing model, which was then evaluated using the external test set. All QSAR regression models developed in this study were developed following this workflow. The definitions and coefficients of descriptors included for all models are summarized in Table S1. The selected descriptor values of all mAbs included in this study are presented in Table S2.

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List of Abbreviations

| Abbreviation | Definition                                                   |
|--------------|--------------------------------------------------------------|
| ACSINS       | Affinity Capture Self-Interaction Nanoparticle Spectroscopy |
| CIC          | Cross-Interaction Chromatography                             |
| CSI-BLI      | Clone Self-Interaction Nanoparticle Spectroscopy            |
| DLS          | Dynamic Light Scattering                                    |
| EP           | Electric Potential                                          |
| EPL_str      | Potential Strength of Negative Charged Patch                |
| Fab          | Antigen-Binding Fragment                                    |
| Fc           | Fragment Crystallizable region                              |
| FvCSP        | Charge Symmetry Parameters of Fv                            |
| PEG          | Polyethylene Glycol                                         |
| PLS          | Partial Least Square                                        |
| pro_pl_3D    | Protein pI determined based on 3D structure                 |
| QSAR         | Quantitative Structure Activity Relationship                |
| RMSD         | Root Mean Squared Deviation                                 |
| SMAC         | Standup Monolayer Adsorption Chromatography                 |
| SVM          | Support Vector Machine                                      |

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