Brief Report

Use of Debye–Hückel–Henry charge measurements in early antibody development elucidates effects of non-specific association

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ABSTRACT

The diffusion interaction parameter (\(k_D\)) has been demonstrated to be a high-throughput technique for characterizing interactions between proteins in solution. \(k_D\) reflects both attractive and repulsive interactions, including long-ranged electrostatic repulsions. Here, we plot the mutual diffusion coefficient (\(D_m\)) as a function of the experimentally determined Debye–Hückel–Henry surface charge (\(Z_{DHH}\)) for seven human monoclonal antibodies (mAbs) in 15 mM histidine at pH 6. We find that graphs of \(D_m\) versus \(Z_{DHH}\) intersect at \(Z_{DHH} \approx 2.6\), independent of protein concentration. The same data plotted as \(k_D\) versus \(Z_{DHH}\) show a transition from net attractive to net repulsive interactions in the same region of the \(Z_{DHH}\) intersection point. These data suggest that there is a minimum surface charge necessary on these mAbs needed to overcome attractive interactions.

Statement of Significance: Within the protein electrostatics framework, a unique discussion is being carried out to depict the thermodynamic and hydrodynamic contributions to \(k_D\) via a novel model using the diffusion coefficient as a function of Debye–Hückel–Henry surface charge (\(Z_{DHH}\)). This work is a guide for protein engineering towards more developable mAbs.

KEYWORDS: antibody developability; surface charge; colloidal interactions; dynamic light scattering; diffusion interaction parameter

INTRODUCTION

Despite decades of research, the relationship between the colloidal stability of protein solutions at high concentrations and pairwise protein–protein interactions is still not well understood [1, 2]. Part of the problem is that the bulk of this research has been done using globular proteins such as bovine serum albumin [3], \(\alpha\)-chymotrypsinogen A [4], ovalbumin [5], and lysozyme [6]. However, further investigation on this relationship is needed for the more asymmetric, therapeutically relevant monoclonal antibodies (mAbs) [7].

The diffusion interaction parameter (\(k_D\)), measured using dynamic light scattering (DLS), is a high-throughput, plate-based technique for characterizing protein interactions. In various low ionic strength buffer conditions, good correlation is observed between \(k_D\) with both viscosity [8] and accelerated storage stability of mAbs [9].

\(k_D\) is calculated from Equation 1, where the mutual diffusion coefficient (\(D_m\)) is measured as a function of protein concentration, \(c\). The linear fit of the data yields the self-diffusion coefficient (\(D_s\)) from the intercept at \(c = 0\) and yields the \(k_D\) from the slope [10].

\[
D_m = D_s (1 + k_D c)
\]  

(1)

\(k_D\) depends on both the thermodynamics (\(k_T\)) of the protein–protein interactions and the diffusion hydrodynamics...
shows the dependence of  

\[ k_D = k_T - k_H \]  

(2)

\[ k_D = 2M_w B_{22} - (\xi_1 + v_{sp}) \]  

(3)

Equation 3 shows the explicit form for both \( k_T (2M_w B_{22}) \) and \( k_H (\xi_1 + v_{sp}) \), where \( M_w \) is the molecular weight, \( B_{22} \) is the second virial coefficient, \( v_{sp} \) is the protein partial specific volume, and \( \xi_1 \) is the first-order coefficient of the virial expansion of the friction factor with protein concentration [10]. \( B_{22} \) is made more positive with increasing excluded volume and increasing charge-charge repulsion. Any attractive interaction, including mass action association, will decrease \( B_{22} \); hence, decrease \( k_D \). \( \xi_1 \) depends directly on the solution viscosity and is sensitive to the size and shape of the protein. Any attractive interactions between molecules increases \( \xi_1 \), thereby decreasing \( k_D \). Notice then that \( k_D \) is always decreased by attractive energies between proteins.

At the protein concentrations used in this study, only bimolecular interactions will contribute significantly to \( k_D \). The charge–charge repulsion contribution to \( k_D \) will be greatest at low salt conditions. Thus, it is anticipated that \( k_D \) will be greater at low salt than at high salt concentrations. Electrostatic interactions that are attractive in nature (e.g. charge–dipole) are less affected by salt than interactions that are repulsive in nature (e.g. charge–charge), thus these attractive interactions become more evident at higher salt concentrations [11, 12]. Because both \( k_T \) and \( k_H \) are positive values, positive values of \( k_D \) necessarily indicate the repulsive contributions to \( k_T \) (charge and excluded volume) to dominate over any attractive energy.

We have measured the Debye–Hückel–Henry surface charge (\( Z_{DHH} \)) for a set of seven mAbs with diverse isoelectric points (\( pI \)s of 8.5 ± 2) and measured \( D_m \) to obtain \( k_D \) values using DLS. We find graphs of \( D_m \) versus \( Z_{DHH} \) for the mAbs intersect at a single value of \( Z_{DHH} \) of ~2.6. Correspondingly, values of \( k_D \) cross over from negative to positive over this same narrow range. We propose that the \( Z_{DHH} \) of 2.6 represents the minimum protein surface charge to overcome attractive interactions.

MATERIALS AND METHODS
Monoclonal antibodies
The seven monoclonal antibodies used in this study have been produced by AbbVie and were IgG1s and chosen to be included in this study based on their pre-formulation developability screening assessment and their differing calculated isoelectric points (\( pI \)s of 8.5 ± 2). All seven of these mAbs experienced different levels of Chemistry, Manufacturing and Controls (CMC) development. The heavy chain isotype was either IgG1 or IgG2 and was combined with either a \( \kappa \) or \( \lambda \) light chain.

DLS
Concentrated protein stocks were thawed benchtop from −80°C and buffer exchanged into 15 mM histidine, 100 mM KCl, pH 6.0 using overnight dialysis against 5 L of buffer. Protein solutions were made between 2 and 12 mg/mL and concentration checked on a Thermo Scientific Nanodrop One (Waltham, MA). After passing thorough a 0.22 µm sterile filtration using Corning Costar Spin-X centrifuge tube filters (Salt Lake City, UT) at 1000×g, the diffusion coefficients of the proteins in the solutions were then run on a Malvern Zetasizer Nano ZS (Malvern, Worcestershire, UK) in triplicate.

Membrane-confined electrophoresis
Proteins were buffer exchanged into 15 mM histidine, 100 mM KCl, pH 6 via overnight dialysis against 5 L of buffer. After concentration check and dilution to 2 mg/mL, the Debye–Hückel–Henry surface charge was measured in triplicate in a steady state method (50, 75, and 100 µA) using the membrane-confined electrophoresis (MCE) instrument (Spin Analytical, Berwick, ME). The absorbance traces as a function of time were analyzed in the provided software as instructed by Spin Analytical.

Data and structure modeling
Calculated surface charge values at pH 6 were generated using Chemical Computing Group’s MOE 2016.0802 software (Montreal, Canada). DLS diffusion data were analyzed and graphed in MATLAB (Mathworks, Natick, MA).

RESULTS AND DISCUSSION
The measured Debye–Hückel–Henry charge is different than the calculated surface charge
A growing body of literature has already described the dominant role that protein surface electrostatics has on protein–protein interactions, determined by measurements such as \( B_{22} \) or \( k_D \) [6, 10, 13, 14]. Here, we investigated the relationship between \( k_D \) and the Debye–Hückel–Henry surface charge (\( Z_{DHH} \)) [15] for a set of seven therapeutically relevant mAbs. These mAbs were chosen for their diverse isoelectric points and thus for their surface charges in 15 mM histidine buffer at pH 6, to reflect the suitability to be a development platform approach. For same pH 6 conditions, Figure 1 shows for each mAb the measured \( Z_{DHH} \) by using MCE compared with the theoretical surface charge calculated by MOE. As described in the Methods section, the MCE measurement of \( Z_{DHH} \) includes 100 mM KCl. \( Z_{DHH} \) is a system property that depends on pH and salt concentration [15]; however, the changes in \( Z_{DHH} \) over the salt concentration range used here (< 100 mM) are small (TL, personal observation). The seven mAbs have measured \( Z_{DHH} \) values between 0.9 and 11.1, whereas the calculated surface charges spans from 9 to 44. The substantial difference between the calculated charge and \( Z_{DHH} \) is primarily a consequence of anion binding [15–17].

Concentration dependence of \( D_m \) reveals protein and ionic strength differences in \( k_D \)
Figure 2 shows the dependence of \( D_m \) as a function of protein concentration for each of the mAbs. In Figure 2A
it can be seen that each mAb exhibits a unique slope, where increased slopes generally correlate with increased values of ZDHH (Fig. 1). The slope of these lines in Figure 2 is \( k_D \). Although there are considerable differences in the slopes at low ionic strength (Fig. 2A), the curves collapse to a much narrower range of slopes at high ionic strength (Fig. 2B). The decreased slopes at higher ionic strength are anticipated by the decreased charge–charge repulsion.\(^{[11]}\) Indeed, the slope for all mAbs in Figure 2B is indistinguishable from that anticipated for mAbs exhibiting only excluded volume repulsion, \(-5.34 \text{ mL/g}^{[7]}\).

The near identity of \( k_D \) in high salt suggests that, for these mAbs, bimolecular interactions are dominated by charge–charge repulsion in low salt, which is overcome by Debye–Hückel screening at higher salt. This dominance of electrostatics at low mAb concentrations is in good agreement with previous work\(^{[8,13]}\). Very weak self-association has been observed for mAbs which is hypothesized to be important for effector function\(^{[18,19]}\). However, the effects of self-association only become apparent at concentrations far higher than those used here. Although there are examples of proteins that undergo significant changes in self-interaction properties as a function of pH, these proteins undergo pH-dependent enzymatic cleavage\(^{[20]}\).

**D\(_m\) and \( k_D \) dependence on ZDHH**

With an accurate value of \( Z_{DHH} \) and \( k_D \) for the different mAbs, the relationship between the diffusion parameters \( (D_m \text{ and } k_D) \) and \( Z_{DHH} \) was investigated. For this analysis, values of \( Z_{DHH} \) act as stand-ins for the different mAbs. Figure 3A shows the correlation of \( D_m \) with \( Z_{DHH} \) in 15 mM histidine buffer at \( pH \) 6. For each mAb (i.e. each value of \( Z_{DHH} \)), the concentration dependence of \( D_m \) for that mAb is graphed at the \( Z_{DHH} \). For each protein concentration, values of \( D_m \) vs. \( Z_{DHH} \) fit well to a general logarithmic function \( (D_m = a \cdot \ln(b \cdot Z_{DHH})) \), where \( a \) and \( b \) are the constants) with all curves having \( R^2 > 0.97 \). Strikingly, independent of the protein concentration, all fits intersected at the same \( Z_{DHH} \) of \( \sim 2.6 \).

Furthermore, \( D_m \) at this intersection (\( \sim 4.4 \times 10^{-7} \text{ cm}^2/\text{s} \)) corresponds to the self-diffusion, \( D_s \), value observed in Figure 2A and B and agrees with previous determinations of \( D_s \)\(^{[7]}\). Figure 3B highlights the explicit relationship between the \( k_D \) and \( Z_{DHH} \). It is striking that moving left to right across the \( x \)-axis, \( k_D \) switches from net attraction (\( k_D < -5.34 \text{ mL/g} \)) to net repulsion (\( k_D > 0 \)) at \( Z_{DHH} \). The blue dashed line in Figure 3B identifies that the \( k_T \) and \( k_H \) contributions to \( k_D \) are equivalent (\( k_D = 0 \text{ mL/g} \)). The red dashed line indicates the \( k_D \)
predicted by Yadav et al. where there are no net attractive or repulsive protein–protein interactions \( (k_D < -5.34 \text{ mL/g}) \) [7]. Thus, the pink shaded area in Figure 3A and B shows the \( Z_{DHH} \) region where protein–protein interactions switch from attractive (below the red dashed line) to repulsive (above the red dashed line). Though the critical \( Z_{DHH} \) range might be better defined using more mAbs, the fact that IgGs exhibit weak attractive interactions of varying strength [19] \( (K_D \text{-apparent} > 100 \mu M) \) makes it difficult to say whether this range would be broader or narrower. In any case, the implications of this result are intriguing.

DISCUSSION

For the first time we use measurements to identify a \( Z_{DHH} \) range where the thermodynamic and hydrodynamic contributions to \( k_D \) balance. In the absence of thermodynamic interactions, \( k_T = B_{22} = 0 \) (indicating no net attractive or repulsive interactions). Under these conditions only \( k_H \), the hydrodynamic contribution to \( k_D \), remains. It has been estimated that IgGs should exhibit \( k_D \) values of either \(-5.34 \text{ mL/g}\) or \(-8.0 \text{ mL/g} \) [21] under these conditions. Kholodenko and Douglas predict that \( k_D \) should equal the negative intrinsic viscosity \( [\eta_i] \), or \(-2.5 \text{ mL/g} \) for non-interacting spheres [22, 23]. However, \( [\eta_i] \) is sensitive to shape [24]. For mAbs, which have an axial ratio \( \sim 5 \), both the measured and predicted value of \( [\eta_i] \) are \( \sim 6 \text{ mL/g} \) [24, 25], resulting in \( k_H = -6 \text{ mL/g} \), which is in excellent agreement with the Yadav et al. prediction of \(-5.34 \text{ mL/g} \) [5].

The data in Figure 3A were fit to a general logarithmic function simply to guide the eye and show the trend of the diffusion data, with no theoretical justification. At this time, the fact that the lines in Figure 3A all cross at the same value as \( D_0 \) (\( D_m \) extrapolated to zero protein concentration) must be considered an intriguing observation. Furthermore, for the first time we report the \( Z_{DHH} \) range over which the interactions contributing to the \( k_D \) go from net attractive to net repulsive, giving an indication the minimum mAb surface charge is needed for good solubility and low viscosity. It must be kept in mind, however, that greater \( Z_{DHH} \) may be needed to overcome the intrinsic attractive interactions between IgGs [19].

CONCLUSION

This brief report highlights a novel methodology that may be used to improve mAb screening assessments, by being able to differentiate within candidates that have physical stability attributes dominated by electrostatic conditions and appear not to strongly self-associate at the solvent conditions used. Although some surface charge on a molecule might be good to prevent aggregation [26], non-uniform distribution of charge across the molecule can cause dipoles that have adverse effects on high concentration properties such as solution viscosity [16] and solubility [27]. This result may provide a starting point to identify the minimum amount of charge desired on the surface of mAb candidates to guide earlier protein design and engineering decisions.

Figure 3. Protein concentration and diffusion coefficient data plotted versus surface charge. (A) The diffusion coefficient for the seven mAbs in 15 mM histidine buffer, pH 6 plotted as a function of \( Z_{DHH} \) at various protein concentrations. The dotted lines are general logarithmic model fits to the data (\( D_m = a^*\ln(b^*Z_{DHH}) \), where a and b are the constants) are to guide the eye and all have \( R^2 \) values above 0.97. Each vertical grouping of data points at constant \( Z_{DHH} \) corresponds to a different mAb identified by the numeral above the data grouping, consistent with the mAb numbering in Figures 1 and 2. Note that mAb3 and mAb5 both had a \( Z_{DHH} \) of \( \sim 11 \). (B) The \( K_D \) is a more conventional representation of the protein concentration and diffusion coefficient data shown in panel A as a way to understand protein interactions in solution. The \( K_D \) was measured at 15 mM histidine, pH 6, and plotted against \( Z_{DHH} \). The horizontal blue dashed line represents the point at which thermodynamic and hydrodynamic interactions contribute equivalently to the \( K_D \), and the horizontal red dashed line represents the \( K_D \) of \( \sim -5.34 \text{ mL/g} \) that represents no attractive or repulsive interactions between IgG proteins. 14 The pink shaded area between the vertical pink dashed lines shows the \( Z_{DHH} \) regime in which the \( K_D \) inflection point happens as it goes from attractive to repulsive.

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The authors declare no conflict of interest and may own AbbVie stock. Dana Filoti is an employee of AbbVie. Thomas M. Laue is a Carpenter Emeritus Professor at the University of New Hampshire. Joshua Laber is currently an employee of Nektar Therapeutics.

ETHICS AND CONSENT
This work has been conducted by respecting the ethical scientific standards.

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ANIMAL RESEARCH
No applicable. No animals have been used for conducting this research.

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