

A New Approach to Study the Physical Stability of Monoclonal Antibody Formulations – Dilution from a Denaturant

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Abstract

The early-stage assessment of the physical stability of new monoclonal antibodies in different formulations is often based on high-throughput techniques that suffer from various drawbacks. Accordingly, new approaches that facilitate the protein formulation development can be of high value to the industry. In this study, a dynamic light scattering (DLS) plate reader is used to measure the aggregation (by means of the increase in the hydrodynamic radius (R_h)) of monoclonal antibody samples that were subject to incubation and subsequent dilution from different concentrations of a denaturing agent, i.e. guanidine hydrochloride. The increase in the R_h of the protein samples is dependent not only on the denaturant concentration used but also on the buffer in which the incubation/dilution was performed. We also compare the aggregation after dilution from a denaturant with other high-throughput stability-indicating methods and find good agreement between the techniques. The proposed approach to probe the physical stability of monoclonal antibodies in different formulation conditions offers a unique combination of features – it is isothermal, probes both the resistance to denaturant-induced unfolding and the colloidal protein stability, it is entirely label-free, does not rely on complex data evaluation, and requires very short instrument measurement time on standard equipment.

Keywords

Protein aggregation, Protein formulation, Protein folding/refolding, Light scattering (dynamic), Stability, Fluorescence spectroscopy

Abbreviations

- C_m concentration of denaturant required to unfold 50 % of the protein;
- dG the Gibbs free energy of unfolding (apparent values in this work);
- DLS dynamic light scattering;
- GuHCI guanidine hydrochloride;
- ICD isothermal chemical denaturation;
- mAb monoclonal antibody;
- T_m protein melting temperature;

Introduction

One of the goals in the (pre-)formulation development of new monoclonal antibodies (mAbs) is to find conditions that provide high protein physical stability. Such studies can be performed with high-throughput techniques which can be classified as isothermal or non-isothermal. Two of the most widely used non-isothermal techniques are differential scanning calorimetry (DSC) and differential scanning fluorimetry (DSF) ^{1–4}. In general, non-isothermal techniques suffer from drawbacks related to the heating of the sample. For example, aggregation during temperature ramps often hinders the thermodynamic evaluation of DSC data and affects the accuracy of the determined protein melting temperatures from both DSC and DSF ^{5,6}. Furthermore, non-isothermal techniques suffer from the fact that the properties of many excipients (e.g. the pH of amine buffers like histidine) change during heating, which can affect the obtained stability rankings based on protein melting temperatures ⁷.

Isothermal chemical denaturation (ICD) is a valuable technique that avoids the above-mentioned drawbacks of DSC and DSF ^{5,7}. Historically, ICD was not the method of choice for protein formulation studies mostly due to the tedious sample preparation ⁸. These limitations of ICD have recently been overcome by the use of equipment that can (semi-)automatically prepare and measure protein samples containing varying concentrations of a denaturating agent ^{5,7–10}. However, the accurate thermodynamic evaluation of ICD data assumes that the protein unfolding process is fully reversible, the system is in equilibrium and the denaturation graph fits a known model (e.g. two-state, three-state unfolding, etc.) ^{11,12}. A recent paper shows that neither reversibility nor equilibration times in an ICD experiment with mAbs are trivial ¹¹. In addition, multidomain proteins may also exhibit multiple transitions that can be close to each other or (partially) overlap which can introduce a large error to the parameters derived from the fit.

Last but not least, high-throughput DSF and ICD methods are usually based on intrinsic protein fluorescence measurements (i.e. observations due to a change in the tryptophan exposure after protein unfolding) ^{2,5,13}. This creates complications when no tryptophan is present or when the tryptophan in a certain protein domain is already solvent exposed in the native protein conformation.

Considering the above-mentioned issues, we saw a demand for novel approaches that can be used in a high-throughput manner to investigate the physical stability of new mAbs in different formulations. The hypothesis we present is that isothermal incubation and dilution of monoclonal antibodies from guanidine hydrochloride solutions with a certain concentration will lead to a substantial protein aggregation. Further, we propose that the amount and the size of the aggregates formed will depend on the formulation conditions and the physical stability of the mAbs (Fig 1) and that this aggregation will be in good agreement with other methods used to study the protein physical stability. Previously, dilution from a denaturant was used to probe the molten globule states and the stability during refolding of other proteins (e.g. human growth hormone^{14,15}, lysozyme¹⁶, recombinant human gamma interferon¹⁷, and others^{18–20}). However, previous work focuses mostly on proteins that are expressed as inclusion bodies in bacteria and the aim of such experiments was usually to achieve higher monomer yields after expression, solubilization and subsequent refolding. To best of our knowledge, we are the first to propose that the assessment of the aggregation after incubation and dilution from different concentrations of a denaturant can be used for high-throughput formulation studies of large proteins (i.e. mAbs, that are typically not expressed as inclusion bodies).

In this work, we use a dynamic light scattering plate reader to assess the aggregation (i.e. the increase in the hydrodynamic radius) of two mAbs after dilution from different concentrations of guanidine hydrochloride (GuHCI) in different buffers. We confirm our hypothesis that conditions that provide higher physical stability of the protein will require higher GuHCI concentrations to induce protein aggregation after dilution of the denaturant. We also show that conditions that provide higher protein colloidal stability will result in a smaller increase of the R_h after dilution from GuHCI. We compare the proposed approach with other established high-throughput methods (e.g. DSF, ICD) to find agreement between the predictions. The method we investigate provides a unique combination of features – it is an isothermal technique that simultaneously probes the resistance to GuHCI-induced unfolding and the colloidal https://doi.org/10.1016/j.xphs.2018.08.004

protein stability (i.e. the level of aggregation after dilution from GuHCI). In addition, no complex data evaluation or a fitting to a certain protein unfolding model is required (in contrast to ICD). Additionally, the approach is label-free and does not rely on intrinsic protein fluorescence measurements (contrary to DSF and ICD). Finally, the proposed approach offers a high potential for scale down and full automation with the existing infrastructure in many protein formulation laboratories.

Materials and Methods

Monoclonal Antibodies

Both monoclonal antibodies (LMU-1 and PPI03) used in this work belong to the IgG1 subclass. SDS-PAGE of the protein bulk shows only bands for the mAb monomer and mAb fragments. Relative monomer area of both bulks is > 99.5 %, measured by size exclusion chromatography.

The mAb buffer was exchanged by dialysis at 20 - 25 °C using Spectra/Por® 8000 MWCO dialysis tubing (Spectrum Laboratories Inc., Rancho Dominguez, USA). The sample to buffer ratio was 1:200 and two buffer exchanges were performed 3 and 8 hours after the beginning. After the last change, dialysis was continued for another 16 hours. The final buffers of both proteins contained 10 mM citrate or 10 mM histidine with pH 5 or 5.75. Also, all LMU-1 samples contained 0.05 % w/v polysorbate 20, while all PPI03 samples were free of surfactants.

For the experiment where we compare the effect of additives, 2x stock solution of the respective additive (i.e. sucrose, trehalose, arginine hydrochloride or proline) in the respective buffer were mixed with the protein to obtain a final concentration of 200 mM sucrose, 200 mM trehalose, 200 mM arginine hydrochloride or 200 mM proline. For the final protein concentrations in the different experiments see the Results and Discussion section.

The protein concentration was measured with a Nanodrop 2000 (Thermo Fisher Scientific, Wilmington, USA). Reagent chemicals from the highest grade available were purchased from Sigma Aldrich (Steinheim, Germany) or VWR International (Darmstadt, Germany). Highly purified water (HPW) was used for the preparation of all buffers.

Sample Preparation, Incubation and Dilution from Different Concentration of Guanidine Hydrochloride

Samples containing protein (LMU-1 or PPI03) and 0, 1, 2, 3, or 4 M guanidine hydrochloride in buffer were prepared by mixing 25 µL of protein solution in the respective buffer (or in the respective buffer with an additive i.e. 200 mM sucrose, 200 mM trehalose, 200 mM arginine hydrochloride or 200 mM proline) with 75 µL of guanidine hydrochloride solution in the same buffer (or in the same buffer containing an additive – see above). After 24 hours of incubation with the denaturant, the samples were diluted 10 times by rapid addition of the respective buffer (or the respective buffer containing an additive – see above) and incubated for another 24 hours. For protein concentration in the final samples after the last dilution refer to the Results and Discussion section. The pH of the samples was controlled at each step using an appropriately calibrated InLab[™] Nano pH Electrode with a SevenEasy pH meter (Mettler Toledo, Germany). The pH reported is the pH of the samples after the final dilution from the denaturant (±0.1 pH units). Finally, the samples were centrifuged at 8200 x g for 10 minutes and measured with a DLS plate reader (see next sections). All experiments were performed in triplicates.

Isothermal Chemical Denaturation (ICD)

Protein in formulation buffer and various amounts of buffer and 6 M guanidine hydrochloride solution in the respective buffer were combined in a 384- non-binding well plate (Corning, USA) using the Viaflo Assist pipetting station equipped with a 16-channel 12.5 µL and a 125 µL pipette (Integra Biosciences, Konstanz, Germany) as already described ⁷(Dilution schemes are available in the supplementary data as Tab S1 and S2). The pH of the samples was controlled after preparation as described in the previous section. Next, the well plate was sealed. After 24 hours incubation at room temperature, the protein fluorescence intensity at 330 nm and 350 nm was measured after excitation at 280 nm with a FLUOstar Omega microplate reader (BMG Labtech, Ortenberg, Germany). The intrinsic fluorescence intensity ratio F350/330 (the fluorescence intensity at 350 nm divided by the fluorescence intensity at 330 nm) was plotted against the denaturant concentration to obtain isothermal chemical denaturation graphs. All samples were prepared in triplicates and the best fit to a three-state model of each replicate in the CDpal software was used to determine the apparent Gibbs free energy of unfolding (dG) and the denaturant concentration needed to unfold 50 % of the protein (C_m). Other unfolding models available in the CDpal software (e.g. two-state, three-state with dimerization of the intermediates, etc.) were also investigated but showed worse fit quality than the three-state model after an f-test comparison. The three-state model was also used by other groups to evaluate isothermal chemical denaturation data with mAbs ^{11,21}. Since all samples were evaluated with the three-state model, two C_m values were determined – C_{m1} for the transition taking place at lower denaturant concentration and C_{m2} for the other transition.

Dynamic Light Scattering (DLS)

 $25 \ \mu$ L of each sample were pipetted in triplicates into a 384- clear bottom well plate (Corning, USA) and the plate was centrifuged at 2000 rpm for 2 minutes using a Heraeus Megafuge 40 centrifuge equipped with an M-20 well plate rotor (Thermo Fisher Scientific, Wilmington, USA). Next, each well was capped with approximately 5 µl of silicon oil and centrifuged again at 2000 rpm for 2 minutes. 10 acquisitions of 10 seconds at 25 °C were taken with the Dyna Pro II DLS plate reader (Wyatt Technology Europe, Dernbach, Germany). The Dynamics V7.8 software was used for all the calculations. The autocorrelation function (ACF) of each sample was calculated from the fluctuation of the light scattering intensity. Cumulant analysis was used to derive the apparent coefficient of self-diffusion (D) and the polydispersity index (PDI). The viscosity of each sample was measured with a falling ball viscosimeter, AMVn (Anton Paar GmbH, Ostfildern-Scharnhausen, Germany). The Stokes-Einstein equation was used to calculate the apparent hydrodynamic radius (R_h) at 25 °C from the D and the sample viscosity. As an additional part of the data analysis, the sample size distribution was calculated from the regularization method. Unless otherwise stated, the R_h values reported in this work are derived from the cumulant analysis.

To determine the interaction parameter (k_D) , different concentrations of LMU-1 (from 1 to 15 g/L) and PPI03 (from approximately 1 to 10 g/L) in each buffer were prepared and measured as described above. The interaction parameter was calculated using the expansion of the apparent diffusion coefficient:

$D=D_0(1+k_Dc)$

where D_0 denotes the diffusion coefficient of an isolated scattering solute molecule in a solvent and c is the protein concentration. The linear fits of the concentration dependence of D_0 from which k_D was determined can be found in the supplementary data (Fig S3 and Fig S4)

Differential Scanning Fluorimetry (DSF)

All samples were diluted to a protein concentration of 1 g/L with the respective buffer and filled into 9 μ L microcuvette arrays (Unchained Labs, USA). The Optim® 1000 system (Unchained Labs, USA) was used to apply a temperature ramp of 1 °C/min starting from 25 °C and ending at 100 °C. Protein fluorescence spectra were collected during the ramp after excitation at 266 nm. The system software was used to calculate the intrinsic fluorescence intensity ratio from the fluorescence intensity at 350 nm and 330 nm (F350/330) against temperature. The protein melting temperatures (T_m) were determined

from the maximum of the first derivatives of these graphs using the Optim® 1000 software (Avacta Analytical, United Kingdom). T_{m1} was assigned to the transition at lower temperature and T_{m2} was assigned to the transition at a higher temperature.

Statistical Data Analysis and Comparison

The mean values, standard deviations, 95 % lower confidence interval (LowCI) and 95 % upper confidence interval (UpCI) of the means were calculated with Origin 2018. A significant difference between two values was considered when the 95 % confidence intervals of two means were not overlapping. In the case of slightly overlapping confidence intervals, we could not prove a significant difference, although the lack of such cannot be confirmed due to the small samples size and the approach we used²².

Results and Discussion

Protein Aggregation and R_h after Dilution from Different Concentrations of Guanidine Hydrochloride

The R_h of LMU-1 and PPI03 after incubation with different concentration of guanidine hydrochloride and subsequent dilution is dependent not only on the denaturant concentration but also on the formulation used (Fig 2A and 2B). The increase in the R_h is due to the formation of more (and/or larger) soluble aggregates, which was confirmed with size exclusion chromatography coupled to multi-angle light scattering detector (see Fig S6 in the supplementary data). The differences in the Rh are most pronounced after dilution from 3 M guanidine hydrochloride for both proteins. A comparison between the different LMU-1 samples diluted from 3 M guanidine hydrochloride shows that the lowest Rh is measured in 10 mM histidine pH 5.75 with a mean value of 19.43 nm and a lower and upper 95 % confidence interval of 18.81 nm and 20.06 nm respectively , followed by 10 mM citrate pH 5.75 (Mean = 24 nm with LowCI = 23.1 nm and UpCI = 24.89 nm), 10 mM histidine pH 5 (Mean = 29.6 nm with LowCI = 27.86 nm and UpCI = 31.34 nm) and 10 mM citrate pH 5 (Mean = 43.1 nm LowCI = 40.08 nm and UpCI = 46.11 nm) (Fig 2A). Similar observations are made for PPI03 (Fig 2B). In the case of PPI03 the following means and 95 % confidence intervals of the hydrodynamic radius after dilution from 3 M GuHCl were measured – in 10 mM histidine pH 5.75 (Mean = 5.37 nm; LoCl = 5.22 nm; UpCl = 5.51 nm), in 10 mM citrate pH 5.75 (Mean = 5.97 nm; LoCI = 5.82 nm; UpCI = 6.11 nm), in 10 mM histidine pH 5 (Mean = 6.87 nm; LoCl = 6.72 nm; UpCl = 7.01 nm) and in in 10 mM citrate pH 5 (Mean = 7.43 nm; LoCI = 7.29 nm; UpCI = 7.58 nm). The PDI of all samples is between 0.05 and 0.2 when the proteins are diluted from up to 1 M GuHCl, while the PDI of all samples diluted from 3 M or more GuHCl is between 0.2 and 0.4. An example of the ACFs with the corresponding cumulant fits and the size distribution from the regularization analysis can be found in the supplementary data (Fig S8). Important to mention, none of the samples formed a pellet of insoluble matter after dilution and centrifugation. In addition, the absorption of the protein in the supernatant was measured at 280 nm and no loss of soluble protein was observed (data not shown). Also, the R_h in the samples after 24 hours of incubation in GuHCl prior to dilution is measured and can be found in the supplementary data (Fig S7). Further, we studied whether the Rh changes during longer incubation time and found no significant differences in the R_h of the samples for up to 1 week after dilution from GuHCI (data not shown).

Isothermal Chemical Denaturation

The isothermal chemical denaturation graphs obtained with the intrinsic protein fluorescence intensity ratio (F350/330) show that both mAbs exhibit complex unfolding behaviour which fits well to a three-state transition model (see Fig S1 and S2 in the supplementary data). A visual observation of the ICD graphs shows that higher concentration of guanidine hydrochloride is needed to unfold both proteins at pH 5.75 compared to pH 5 (Fig 2A and 2B), which indicates higher resistance to the GuHCI-induced unfolding of the mAbs at the higher pH. To further support this statement, in the case of LMU-1, the mean C_{m2} values are significantly higher at pH 5.75 compared to pH 5 with non-overlapping confidence intervals (Tab 1A). In the case of PPI03, the difference between the C_{m2} in 10 mM histidine pH 5 and 10

mM histidine pH 5.75 is also significant, although the same statement is difficult to make for the results in 10 mM citrate due to the slightly overlapping confidence intervals (Tab 1B). Comparing the apparent dG values at different pH values reveals that in the case of LMU-1 there is a significant difference (nonoverlapping confidence intervals) between the dG measured in 10 mM citrate pH 5 and in 10 mM citrate pH 5.75, which indicates higher conformational stability of LMU-1 at the higher pH. In the case of PPI03, although the mean values for the dG differ at pH 5 and pH 5.75, a significant difference cannot be proved due to the overlapping confidence intervals (Tab 1B). Further, the ICD graphs of the proteins obtained in histidine or citrate with the same pH almost completely overlap (Fig 2C and 2D). Important to note, when comparing the stability of the two mAbs in 10 mM citrate or 10 mM histidine with the same pH, there is not a single case in which a significant difference between ICD data obtained at the same pH can be confirmed due to the overlapping confidence intervals (Tab 1A) and 1B).

Colloidal Stability of the mAbs in Different Buffers

The colloidal stability of both mAbs in citrate and histidine buffer was assessed by means of the interaction parameter k_D . It has been shown that k_D describes the interparticle interaction in protein solution ²³. Highly positive k_D is attributed to repulsive interactions, whereas highly negative is related to attractive interactions ²⁴. The k_D of both LMU-1 and PPI03 is positive in histidine buffer and negative in citrate buffer (Tab 1A and 1B). This indicates that both proteins have higher colloidal stability in 10 mM histidine buffer compared to 10 mM citrate in the studied pH range. However, the k_D is not a parameter that can assess the conformational stability of the protein and it is demonstrated that both the conformational and colloidal protein stability is important for the aggregate formation in solution ²⁵.

Thermal Stability of the mAbs in Different Buffers

Both LMU-1 and PPI03 have higher melting temperatures (T_{m1}) at pH 5.75 compared to pH 5 (Tab 1A and 1B). This indicates higher physical stability of both mAbs at pH 5.75 and corresponds well to the higher concentration guanidine hydrochloride needed to unfold both proteins at higher pH. The mean values of the melting temperatures of both proteins in histidine almost always appear lower compared to citrate counterparts most probably due to the buffer pH shift of histidine during heating⁷. However, a significant difference between the melting temperatures of the mAbs in either 10 mM citrate or 10 mM histidine with the same pH cannot be proved due to the small sample size and the overlapping confidence intervals (Tab 1A and 1B). The thermal denaturation graphs of each mAb sample can be found in the supplementary data (Fig S5).

The Relation Between the $R_{\rm h}$ after Dilution from GuHCl and Other Parameters

The increase of R_h after dilution from a certain concentration of guanidine hydrochloride (in our case 3 M) will depend on the degree of protein unfolding at these conditions. In the case of LMU-1 and PPI03 samples with pH 5, the isothermal chemical denaturation curves are shifted to lower denaturant concentrations compared to samples with pH 5.75, which indicates a lower resistance to the GuHCl-induced unfolding of these proteins at the lower pH (Fig 2C and 2D). Respectively, a higher R_h (more and larger aggregates formed) is measured after dilution from 3 M guanidine hydrochloride in buffers with pH 5 compared to dilution in buffers with pH 5.75 (Fig 2A and 2B).

When the degree of protein unfolding (or structural perturbation) at a given denaturant concentration is the same (e.g. buffers with the same pH or higher denaturant concentrations where the protein is fully unfolded regardless of the pH i.e. at 4 M GuHCl), the protein aggregation and the measured R_h after dilution will depend mostly on the colloidal stability and the aggregation propensity of the (partially) unfolded protein after dilution in these conditions. For example, the ICD curves of LMU-1 in citrate or histidine buffer with the same pH almost completely overlap (Fig 2C and 2D). However, the R_h of LMU-1 after dilution from 3 M GuHCl is always lower in histidine buffer compared to citrate counterparts with the same pH (Fig 2A). Similar observations are made also with the other mAb - PPI03 (Fig 2B). Also, at 4 M GuHCl, there is the same degree of unfolding (according to the intrinsic protein fluorescence ratio – Fig 2C and 2D) for both proteins regardless of the formulation pH and buffer. Our assumption is that

in this case the aggregation after dilution will be driven mostly by differences in the protein colloidal stability and we see that the R_h after dilution of the proteins from 4 M GuHCL is always lower in histidine compared to citrate, regardless of the pH. This behaviour is in good agreement with the high k_D values of both proteins in histidine which indicates a higher protein colloidal stability in this buffer in comparison to citrate.

Effect of Different Additives on the Change in the R_h after Dilution from GuHCI

We further tested if the herein proposed approach could be used to study the effect of different additives (i.e. 200 mM sucrose, 200 mM trehalose, 200 mM arginine hydrochloride or 200 mM proline) on the increase in R_h (i.e. aggregation) of the mAbs after dilution from GuHCI. To study this, we focused on dilution from 3 M guanidine hydrochloride since this was the concentration of denaturant where we observed the largest differences in the earlier experiments. Next, we performed the dilution in a buffer i.e. 10 mM histidine pH 5, in which the physical stability of both mAbs is neither the worst nor the best according to the different techniques we used in this work. We reported the effect of the additives as the difference in the R_h after dilution with the additive compared to dilution without an additive (Figure 3). We observed that for both LMU-1 and PPI03 all additives tested were significantly better compared to the histidine buffer alone (Figure 3). However, in the case of LMU-1 200 mM sucrose and 200 mM arginine chloride caused the largest decrease in the R_h i.e. the highest stabilizing effect (Figure 3 – Left), while in the case of PPI03 200 mM sucrose and 200 mM trehalose were the most stabilizing excipients according to the proposed approach (Figure 3 – Right).

Conclusion and Outlook

In this work, we demonstrated with two mAbs that the DLS assessment of the aggregation (by means of the R_h) after dilution from different concentrations of guanidine hydrochloride is a promising approach to probe the protein physical stability in different formulations.

A formulator would then aim to i) find buffers and pH that shift the increase in R_h after dilution to higher guanidine hydrochloride concentrations (i.e. find conditions that provide higher resistance to guanidine hydrochloride unfolding); and ii) find conditions that minimize R_h dilution (i.e. find conditions that provide higher colloidal stability and less aggregation during dilution from a denaturant);

The proposed method could be used alone or as a complementary technique. If used alone, we suggest that at least several guanidine hydrochloride concentrations (in a step of 0.5 or 1 M) are tested to find conditions where the increase in the R_h is largest after dilution. Next, this denaturant concentration could be used to test the influence of different buffers and additives on the increase in the R_h . In the case of the antibodies that we tested, such denaturant concentration is usually around 3 M guanidine hydrochloride.

Alternatively, the method we show could be used as a complementary technique to isothermal chemical denaturation (e.g. to distinguish if there is a difference in the protein physical stability in conditions that exhibit overlapping curves in an isothermal chemical denaturation experiment). In such case, we suggest that the concentration of guanidine hydrochloride that can be used is around the denaturant concentration where the protein is partially unfolded according to the ICD experiment.

The approach we propose offers a unique combination of features: i) the method is isothermal (problems arising from sample heating are avoided); ii) it can distinguish between overlapping curves in an ICD experiment due to the different colloidal stability and different level of aggregation after dilution from such samples; iii) the method is label-free and independent of the intrinsic protein fluorescence; iv) very short instrument measurement time for one sample is required; v) there is a high potential for scale down and automatization with the help of already available equipment in many protein formulation laboratories;

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Conflict of Interests

L.G. and D.R. are employees of Wyatt Technology Europe GmbH.

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Figure legend

Figure 1. Schematic explanation of the working hypothesis - Impact of the physical stability of a mAb on the protein aggregation after incubation and subsequent dilution from a denaturant.

Dilution with buffer providing Ulution with purier providing Incubation in guanidine More and/or larger hydrochloride Dilution with buffer providing high protein physical stability aggregates Native mAb monomers mAb monomers with perturbed structure detected by intrinsic protein fluorescence changes Less and/or smaller protein aggregates

Figure 2. Hydrodynamic radius Rh from dynamic light scattering of LMU-1 (A) and PPI03 (B) after incubation and dilution from different concentrations of guanidine hydrochloride in 10 mM histidine pH 5 (red squares), 10 mM histidine pH 5.75 (blue circles), 10 mM citrate pH 5 (yellow diamonds), and 10 mM citrate pH 5.75 (green triangles). The protein concentration in the measured samples in (A) and (B) is 1 g/L and 0.2 g/L for LMU-1 and PPI03 respectively; Isothermal unfolding of LMU-1 (C) and PPI03 (D) in presence of different concentrations of guanidine hydrochloride measured by the change in the intrinsic protein fluorescence intensity ratio (F350/330). The protein concentration in the measured samples in (C) and (D) is 1 g/L and 0.5 g/L for LMU-1 and PPI03 respectively;



Figure 3. Effect of various additives on the change in the hydrodynamic radius R_h from DLS of LMU-1 (left) and PPI03 (right) after dilution from 3 M guanidine hydrochloride in 10 mM histidine pH 5. All measurements were corrected for viscosity as described in Materials and methods. For clarity reasons, the values with additives are represented as a difference from the value without an additive. Negative values indicate that the R_h is lower after dilution with the additive compared to dilution without an additive. The circles represent the mean value of the measurements, the bars represent the upper and lower 95 % confidence intervals. The final protein concentration after dilution is 1 g/L for both LMU-1 and PPI03.





Table 1. Physical stability parameters of LMU-1 (A) and PPI03 (B) in different buffers and pH. The apparent Gibbs free energy of unfolding (dG) and the melting denaturant concentrations for the first and second unfolding (C_{m1} and C_{m2}) are obtained with isothermal chemical denaturation as described in Materials and methods. The melting temperatures of the protein (T_{m1} and T_{m2}) are obtained from the maximum of the first derivative of the protein unfolding measured with differential scanning fluorimetry as described in Materials and methods. All values (except the k_D) are means of triplicates and the 95 % lower and 95 % upper confidence intervals are reported in the brackets (Lower CI 95 % - Upper CI 95 %) next to each mean.

Table 1A LMU-1	dG, kJ/mol	C _{m1} , M	C _{m2} , M	k₀, mL/g	T _{m1} , ℃	T _{m2} , °C
histidine pH 5	94.91 (90.06 - 99.77)	1.76 (1.61 - 1.92)	2.44 (2.40 - 2.49)	37.9	62.20 (61.56 - 62.83)	77.5 (76.90 - 78.20)
citrate pH 5	76.84 (60.35 - 93.34)	1.67 (1.21 - 2.13)	2.38 (2.33 - 2.43)	-1.18	63.52 (62.17 - 64.88)	78.5 (77.41 - 79.58)
histidine pH 5.75	106.10 (98.70 - 113.49)	1.92 (1.74 - 2.10)	2.66 (2.61 - 2.72)	40.2	66.82 (65.09 - 68.55)	79.7 (79.23 - 80.26)
citrate pH 5.75	111.84 (100.98 - 122.70)	1.94 (1.80 - 2.08)	2.60 (2.56 - 2.63)	-8.46	68.29 (66.41 - 70.18)	79.3 (77.45 - 81.15)
Table 1B						
Table 1B PPI03	dG, kJ/mol	C _{m1} , M	C _{m2} , M	k _D , mL/g	T _{m1} , ℃	T _{m2} , ℃
Table 1B PPI03 histidine pH 5	dG, kJ/mol 75.06 (63.83 - 86.29)	C _{m1} , M 1.86 (1.62 - 2.09)	C _{m2} , M 3.00 (2.99 - 3.01)	k _D , mL/g 42	T _{m1} , °C 62.87 (61.48 - 64.25)	T _{m2} , °C 76.4 (74.67 - 78.13)
Table 1B PPI03 histidine pH 5 citrate pH 5	dG, kJ/mol 75.06 (63.83 - 86.29) 74.84 (50.41 - 99.27)	C _{m1} , M 1.86 (1.62 - 2.09) 1.99 (1.79 - 2.21)	C _{m2} , M 3.00 (2.99 - 3.01) 3.01 (2.81 - 3.20)	k _D , mL/g 42 -0.58	T _{m1} , °C 62.87 (61.48 - 64.25) 63.88 (63.31 - 64.44)	T _{m2} , °C 76.4 (74.67 - 78.13) 77.99 (75.84 - 78.29)
Table 1B PPI03 histidine pH 5 citrate pH 5 histidine pH 5.75	dG, kJ/mol 75.06 (63.83 - 86.29) 74.84 (50.41 - 99.27) 87.56 (74.70 - 100.42)	C _{m1} , M 1.86 (1.62 - 2.09) 1.99 (1.79 - 2.21) 2.28 (1.78 - 2.77)	C _{m2} , M 3.00 (2.99 - 3.01) 3.01 (2.81 - 3.20) 3.09 (3.04 - 3.14)	k _D , mL/g 42 -0.58 46	T _{m1} , °C 62.87 (61.48 - 64.25) 63.88 (63.31 - 64.44) 68.26 (68.13 - 68.39)	T _{m2} , °C 76.4 (74.67 - 78.13) 77.99 (75.84 - 78.29) 77.01 (77.68 - 78.30)

Supplementary data

"A New Approach to Study the Physical Stability of Monoclonal Antibody Formulations – Dilution from a Denaturant"

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SUPPLEMENTARY FIGURES:

Figure S1. Unfolding of LMU-1 by guanidine hydrochloride in different buffers and pH values. The black points represent the experimentally measured values. The red line is the theoretical fit to a three-state model. The inset shows the corresponding residuals from the fit.



Figure S2. Unfolding of PPI03 by guanidine hydrochloride in different buffers and pH values. The black points represent the experimentally measured values. The red line is the theoretical fit to a three-state model. The inset shows the corresponding residuals from the fit.





Figure S3. Concentration dependence of the diffusion coefficient of LMU-1 in different buffers and linear fits from which the interaction parameter k_D was derived





Figure S5. Thermal unfolding curves obtained from the change in the intrinsic protein fluorescence (F350/330) ratio of LMU-1 (left) and PPI03 (Right) during heating in different buffers – 10 mM histidine pH 5 (black squares), 10 mM histidine pH 5.75 (red circles), 10 mM citrate pH 5 (blue triangles up), 10 mM citrate pH 5.75 (green triangles down).



Figure S6. Size exclusion chromatography and multi-angle light scattering with quasi-elastic light scattering detection (SEC–MALS-QELS).

A. Chromatograms of LMU-1 in 10 mM histidine pH 5 after dilution from 0 M (black), 1 M (red), 2 M (blue) and 3 M (green) guanidine hydrochloride – UV detection at 280 nm (left) and QELS signal (right). The aggregate peak and the light scattering from the aggregate peak increases when LMU-1 is diluted from higher guanidine hydrochloride concentration, which corresponds well to the increase in the R_h measured by DLS.



B. Chromatograms of LMU-1 in 10 mM histidine pH 5.75 (black) or 10 mM citrate pH 5.75 (red) after dilution from 3 M guanidine hydrochloride - UV detection at 280 nm (left) and QELS signal (right). The relative area of the aggregate peak after LMU-1 diluted from either citrate or histidine is similar, but the light scattering from the aggregate peak is higher when LMU-1 is diluted in citrate in comparison to histidine buffer, which indicates that the aggregates formed after dilution in citrate are larger. This corresponds well with the lower R_h measured by DLS after protein dilution in histidine compared to the R_h after dilution in citrate.



Description of the SEC-MALS-QELS method:

The system consisted of Agilent 1260 infinity II pump (Agilent G7111B, Santa Clara, CA, USA) with an online degasser (Agilent G7111B), and a temperature controlled autosampler (Agilent G7129A) at 4°C. The separation was performed with a SUPERDEX 200 INCREASE 10/30 GL column. A TREOS II MALS detector (Wyatt Technology, Santa Barbara, USA) was connected to the system. The MALS detector measured the intensity of the scattered light at 4 scattering angles, 3 angles were used for the static light scattering – QELS). In addition, a variable wavelength detector (Agilent G7114A) operated at 280 nm and a differential refractive index detector (Optilab T-rEX,Wyatt Technology Corp.) were connected to the system. Data collection and processing were performed using the ASTRA software, Version 7.1 (Wyatt Technology, Santa Barbara, USA). The aqueous mobile phase consisted of PBS buffer in HPLC-grade water with 200 ppm of sodium azide (Sigma–Aldrich, St. Louis, USA). The reagent was purum grade and the mobile phase was filtered through Durapore VVPP 0.1 m membrane filters (Millipore Corporation, Billerica, MA, USA).

Figure S7. Hydrodynamic radii of LMU-1 (left) and PPI03 (Right) measured with DLS after 24 hours of incubation in different concentrations of GuHCL and different buffers – 10 mM histidine pH 5 (red), 10 mM histidine pH 5.75 (blue), 10 mM citrate pH 5 (yellow) and 10 mM citrate pH 5.75 (green). The protein concentration is 10 g/L in all LMU-1 samples and 2 g/L in all PPI03 samples. The viscosity of the respective GuHCL concentration was measured as described in Materials and methods and used in the calculations of the R_h. The large differences between the R_h of the proteins in the absence of GuHCL are due to effects from the k_D (see Figure S3 and S4).



Figure S8. Example of autocorrelation functions and the corresponding cumulant fit (black line) from DLS measurements of LMU-1 (A) and PPI03 (B) samples after dilution from different GuHCL concentrations. Size distributions from the regularization analysis of the LMU-1 (C) and PPI03 (D) samples after dilution from different GuHCL concentrations. All experiments for this figure were performed in 10 mM histidine pH 5. The concentration of LMU-1 and PPI03 are 1 g/L and 0.2 g/L respectively.



SUPPLEMENTARY TABLES:

Sample	Buffer	6 M Guanidine	Final guanidine	Protein	Total
No	volume (µL)	hydrochloride in	hydrochloride	solution	sample
		buffer volume	concentration (M)	volume	volume
		(µL)		(µL)	(µL)
1	72	0	0	8	80
2	67	5	0.38	8	80
3	64	8	0.60	8	80
4	61	11	0.83	8	80
5	58	14	1.05	8	80
6	56	16	1.20	8	80
7	54.4	17.6	1.32	8	80
8	52.8	19.2	1.44	8	80
9	51.2	20.8	1.56	8	80
10	49.6	22.4	1.68	8	80
11	48	24	1.80	8	80
12	46.4	25.6	1.92	8	80
13	44.8	27.2	2.04	8	80
14	43.2	28.8	2.16	8	80
15	41.6	30.4	2.28	8	80
16	40	32	2.40	8	80
17	38.4	33.6	2.52	8	80
18	36.8	35.2	2.64	8	80
19	35.2	36.8	2.76	8	80
20	33.6	38.4	2.88	8	80
21	28.6	43.4	3.26	8	80
22	23.6	48.4	3.63	8	80
23	18.6	53.4	4.01	8	80
24	13.6	58.4	4.38	8	80

Sample	Buffer volume	6 M Guanidine	Final guanidine	Protein	Total
No	(μL)	hydrochloride in	hydrochloride	solution	sample
		buffer volume (µL)	concentration	volume	volume
			(M)	(µL)	(µL)
1	72	0	0	8	80
2	67	5	0.38	8	80
3	63.5	8.5	0.64	8	80
4	60	12	0.90	8	80
5	56.5	15.5	1.16	8	80
6	54.5	17.5	1.31	8	80
7	52.5	19.5	1.46	8	80
8	50.5	21.5	1.61	8	80
9	48.5	23.5	1.76	8	80
10	46.5	25.5	1.91	8	80
11	44.5	27.5	2.06	8	80
12	42.5	29.5	2.21	8	80
13	40.5	31.5	2.36	8	80
14	38.5	33.5	2.51	8	80
15	36.5	35.5	2.66	8	80
16	34.5	37.5	2.81	8	80
17	32.5	39.5	2.96	8	80
18	30.5	41.5	3.11	8	80
19	28.5	43.5	3.26	8	80
20	26.5	45.5	3.41	8	80
21	24.5	47.5	3.56	8	80
22	22	50	3.75	8	80
23	18	54	4.05	8	80

 Table S2. Dilution tables for an isothermal chemical denaturation experiment with PPI03