Characterizing Thermal Transitions of IgG with Mass Spectrometry

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Abstract. Variable temperature electrospray ionization (ESI) is coupled with mass spectrometry techniques in order to investigate structural transitions of monoclonal antibody immunoglobulin G (IgG) in a 100-mM ammonium acetate (pH 7.0) solution from 26 to 70 °C. At 26 °C, the mass spectrum for intact IgG shows six charge states +22 to +26. Upon increasing the solution temperature, the fraction of low-charge states decreases and new, higher-charge state ions are observed. Upon analysis, it appears that heating the solution aids in desolvation of the intact IgG precursor.

Above ~ 50 °C, a cleavage event between the light and heavy chains is observed. An analysis of the kinetics for these processes at different temperatures yields transition state thermochemistry of $\Delta H^\ddagger = 95 \pm 10$ kJ mol$^{-1}$, $\Delta S^\ddagger = 8 \pm 1$ J mol$^{-1}$ K$^{-1}$, and $\Delta G^\ddagger = 92 \pm 11$ kJ mol$^{-1}$. The mechanism for light chain dissociation appears to involve disulfide bond scrambling that ultimately results in a non-native Cys$^{199}$–Cys$^{217}$ disulfide bond in the light chain product. Above ~ 70 °C, we are unable to produce a stable ESI signal. The loss of signal is ascribed to aggregation that is primarily associated with the remaining portion of the antibody after having lost the light chain.

Keywords: Mass spectrometry, variable temperature electrospray ionization, antibody degradation, dissociation kinetics

Introduction

The immunoglobulin G (IgG) antibody is a ~ 147-kDa protein in the immune system that is involved in antigen recognition and binding [1]. This molecule is often visualized by the “Y”-shaped diagram shown in Scheme 1. As shown, IgG is composed of a dimer of heterodimers (the heavy and light chains). The heterodimers are linked by two disulfide bonds. The light and heavy chains of each heterodimer are bound by a single disulfide bond. Together these regions create a highly specific antigen binding pocket called the FAB portion of the molecule that is critical for immune response [2]. In recent years, numerous monoclonal antibodies with therapeutic value have been introduced [3–6]. Because of this, an understanding of the structures and stabilities of these molecules is of fundamental importance.

Although calorimetric studies of antibodies are routinely carried out in the development and testing of new therapeutic antibodies, these methods provide information about only the stability of the ensemble average. That is, the structural change is observed as a two-state cooperative transition, and little is known about the nature of the configurations and mechanisms leading to denatured states [7, 8]. In the work presented below, we investigate the stability of IgG using a new, variable temperature electrospray ionization (vT-ESI) source coupled with mass spectrometry (MS) measurements [9, 10]. At elevated temperatures (above 50 °C), the MS measurements reveal that the light chain of IgG dissociates, through a mechanism that involves scrambling of the disulfide bonds, resulting in the formation of a light chain product that adopts non-native Cys$^{199}$–Cys$^{217}$ and...
Cys\textsuperscript{91}–Cys\textsuperscript{140} disulfide bonds. From an Arrhenius analysis of the kinetics of dissociation at varying temperatures, we derive transition state thermochemistry for dissociation process. This thermochemistry is discussed.

The present work builds on a number of new MS-based measurements that are being developed with the aim of understanding structures and structural transitions of biomolecules in solution as well as the gas phase. In the last decade, “native ESI” has enabled the study of large complexes [11–16]. Analyses of biomolecular conformations from solutions of varying composition and temperature now have an extensive history [17–20]. Differences in structures found under varying solution conditions can be investigated with a range of reaction chemistries and techniques, including isotopic hydrogen-deuterium (H-D) exchange [21–26]; fast photochemical oxidation of proteins [27–30]; chemical cross-linking [31–33] and other residue-specific modifications [34, 35]; and ion mobility measurements [9, 36–45]. Once ionized, an array of physical and chemical methods can be used to investigate biomolecular structure in vacuo. These include low-energy and high-energy collisions with buffer gasses [46–51] and surfaces [52–54]; photodissociation techniques [55–58]; measurements of collision cross-sections with many new ion mobility methods [59–66]; ion-molecule reactions, including proton transfer [67–70] and H-D exchange reactions [71–74]; ion-ion reactions [75–77]; and electronic and vibrational spectroscopies [78–81].

Experimental

Variable Temperature Electrospray Ionization

The solution temperature of the ESI emitter is controlled using a home-built variable temperature electrospray ionization emitter [9]. This interface holds a borosilicate glass ESI emitter which has been pulled to a narrow inner diameter of ~1 to 5 µm using a Flaming/Brown P-97 pipette tip puller (Sutter Instruments, Novato, CA, USA). The emitter interface is made of a thermally conductive ceramic block that is resistively heated using a cartridge heater. An ESI voltage of 0.7 to 1.0 kV is applied to a platinum wire that is inserted into the back of the emitter, making electrical connection with the solution. The temperature is measured (to a precision of ±0.5 °C) using a K-type thermocouple that is inserted into the ceramic block.

Instrumentation

Initial experiments were performed on Waters Synapt G2 and remaining kinetics experiments were performed on Waters Synapt G2S instrument. Both instruments were used with the source interlocks overridden to accommodate the vT-ESI source [9, 82]. Source pressures and voltages that minimize ion activation were used for initial studies, summarized below [83, 84]. Backing pressures were increased to ~8 mbar using Speedivave (Edwards, Burgess Hill, UK). Gas control was optimized to minimize ion activation and increase ion transmission, including flow rates in the source (20.9 mL/min), trap (10 mL/min), helium cell (180 mL/min), and IMS cell (90 mL/min). We additionally optimized and used instrument settings, including sampling cone voltage (62 V), extraction cone (1.6 V), source temperature (50 °C), cone gas (10 L/h), flow gas (0.6 bar), purge gas (100 L/h), and trap collision energy (12.6 collision energy). Finally, TriWave DC voltages were also optimized: entrance voltage (3.2 V), bias (45.3 V), trap DC (1.7 V), and exit voltage (1.3 V). We note that there are different source configurations between the Synapt G2 and the Synapt G2S that can contribute to charge state shifts, presumably due to collisional activation. This may be the origin of the charge state shift between the two sets of mass spectra presented. The analysis presented below uses only the time-of-flight mass analyzer. That is, the quadrupole is fixed to transmit all ions. The data were originally acquired as nested ion mobility mass spectra; however, only the mass dimension was used for this analysis as there were no resolvable changes in drift time distributions for these species.

Analysis of the Data

Each dataset was collected by systematically increasing the temperature from 26 to 70 °C. Samples were allowed to incubate at each temperature for at least 3 min. After this time, the mass spectra recorded at low solution temperatures (26 to 45 °C) do not appear to change for an extended period, and it appears that we have reached an equilibrium (on this timescale). At higher temperatures, the antibody dissociates. Kinetics experiments of this process were carried out by increasing the temperature to a set point and then collecting a series of 3-min acquisitions until spray was lost. Unidec [85] (Oxford, UK) was used for deconvolution of native charge state distributions. Data from both experiments were exported using TWIM extract (University of Michigan, Ann Arbor, MI) and
processed using Origin2018 (OriginLab Corporation, North-ampton, MA, USA). Kinetic data were fit using a first-order reaction rate of formation. Measured rate constants, $k$, were plotted as a function of the temperature in an Arrhenius plot and fit linearly to obtain transition state chemistry.

**Sample Preparation**

Immunoglobulin G (IgG1, universal antibody standard, human, ≥ 90% purity) was purchased from Sigma Aldrich (St. Louis, MO, USA). IgG (1 mg) was resolubilized in 100 mM ammonium acetate solution (pH 7.0, 500 μL). Sample was buffer exchanged using a spin concentrator (molecular weight cutoff = 30,000 Da, 100 to 500 μL × 3, Millipore Sigma, Burlington, MA, USA). IgG was brought to a final concentration of 80 μM in 100 mM ammonium acetate (Sigma Aldrich, St. Louis, MO, USA).

**Results and Discussion**

*Changes in ESI Charge State Distribution of IgG Precursor with Temperature*

Figure 1 shows representative mass spectra for intact IgG recorded upon electrospraying a 100 mM ammonium acetate solution at 28, 32, 42, and 52 °C. At low temperatures (28 to 42 °C), a narrow distribution of IgG charge states from +22 to +26 and centered at +24 is observed. As the solution temperature is increased, each charge state becomes noticeably shaper and the distribution of charge states changes. From ~28 to 42 °C, the abundances of the lower-charge state +22 and +23 species decrease with increasing temperature; in this same temperature region, the populations of +24 through +27 also increase. Above 42 °C, the +24 species decreases in abundance and the +25 species is favored. This ion reaches a maximum abundance at ~54 °C and decreases above this temperature. The +26 and +27 continue to increase until ~70 °C, where we no longer maintain a stable ion signal. At this temperature, the clear solution becomes turbid due to the formation of insoluble aggregates [86, 87], likely originating from IgG unfolding. The slight shift in charge state observed here is similar to that seen when tetrameric concanavalin A is heated, which was coupled to a structural change [88]. While we do not observe a change in the collision cross-sections for IgG, there have been reports that IgG undergoes structural changes between 25 and 70 °C under acidic pH [89, 90]. In these cases, highly charged MS peaks emerged at elevated temperatures, indicating that the protein had unfolded. We also find that the peaks decrease in width at elevated temperatures, suggesting that IgG emerges from hot electrospray droplets with fewer species adducted.

**High-Temperature Dissociation of the Light Chain**

At low temperatures, IgG remains stable for long times. We monitored the mass spectra of a solution at 45 °C for up to ~15 h, and it shows no measurable change. At elevated temperatures, IgG is known to dissociate by loss of the light chain [91, 92]. Figure 2 shows mass spectra acquired after incubating solutions at 65 °C for 3, 18, or 42 min. At relatively short incubation times (3 min), the mass spectrum is dominated by peaks associated with the IgG precursor. At longer times (18 min as shown in Figure 2), the relative abundances of the
IgG peaks decrease and a new, well-defined set of peaks \( m/z < 3000 \) are observed. These peaks increase in magnitude with increasing incubation time. The theoretical molecular weight of the light chain species is 22,942 Da. Using this value, we determine that the major peaks correspond to +9 through +13 charge states of the light chain. Once the charge states are assigned, our experimental measurement yields \( m = 22,943 \pm 1 \) Da, in close agreement with the theoretical value. It is interesting that we do not observe the complementary remaining IgG fragment, which should have \( m \sim 124 \) kDa, although aggregation of this species is known to occur rapidly [87, 91].

**Mechanism of Light Chain Dissociation**

Before describing the kinetics experiments, we first present a possible mechanism for the covalent bond cleavage leading to the release of the light chain species. Dissociation of the light chain from the heavy chain must involve cleavage of the Cys\(^{217}\)–Cys\(^{224}\) disulfide bond, between the two chains (Scheme 1). It is established that during the dissociation of the light chain, disulfide bonds can scramble [91, 93, 94]. The precursor IgG antibody used in our study has a single free Cys\(^{91}\) residue on each of the light chains. As we think about thermodynamic considerations, we see that disulfide scrambling upon dissociation of the light chain could stabilize the products. That is, if only the native Cys\(^{224}\)–Cys\(^{217}\) bond between the heavy and light chains is cleaved, three free, unbound Cys residues would be available (Cys\(^{224}\) on the heavy chain; Cys\(^{217}\) and Cys\(^{91}\) on the light chain). Consider one scenario. After the Cys\(^{224}\)–Cys\(^{217}\) bond is cleaved, the freed light chain might refold and in doing so scramble its disulfide bonds in order to stabilize this product. For example, if the Cys\(^{199}\)–Cys\(^{140}\) disulfide bond was also to cleave, we might form a non-native Cys\(^{199}\)–Cys\(^{217}\) linkage. In this case, the newly freed Cys\(^{140}\) residue could form a second disulfide bond with the Cys\(^{91}\) residue and the resulting covalent Cys\(^{140}\)–Cys\(^{91}\) bond would further stabilize the light chain product. With this change, the only free Cys residue is located on the heavy chain (Cys\(^{224}\)) and the light chain is no longer covalently linked to the heavy chain. Overall, this process is a disulfide mixing [95, 96] step with nearby Cys residues both making and breaking covalent bonds, a process that has been described previously for light chain dissociation from IgG under mildly denaturing conditions.

![Figure 2. Mass spectrum showing formation of light chain charge states +9 through +13 after incubation at 65 °C for 3, 18, and 42 min](image)

![Figure 3. Light chain signal monitored over time at 60, 62, and 65 °C, as open squares, triangles, and circles, respectively. Change in light chain abundance with time is shown modeled with first-order kinetics (black lines)](image)
conditions [91, 97–100]. It is likely that at elevated temperatures, a similar disulfide bond scrambling will occur.

With this idea in mind, we carried out studies to identify the location of the non-native disulfide bond. After incubation at 65 °C for 90 min, we alkylated the free Cys residues with iodoacetamide. The products of antibody dissociation were then proteolytically digested and the tryptic peptides that were formed were analyzed using a combination of chromatographic separations with MS detection (see supporting information). An analysis of the cross-linked peptides provides evidence for the non-native Cys\(^{199}\)–Cys\(^{217}\) disulfide bond. Although we anticipate that the Cys\(^{91}\)–Cys\(^{140}\) disulfide bond should also stabilize this fragment, we did not detect this cross-linked peptide in our analysis. We note that it would be difficult for us to fragment this species with our experiment due to its size (60 amino acids); so, the dearth of experimental information does not rule it out entirely. Overall, this result supports the idea that the mechanism for light chain dissociation involves disulfide bond scrambling.

**Kinetics Measurements at Varying Temperatures**

We next carried out a series of kinetics experiments at specified temperatures 57, 60, 62, and 65 °C using the vT-ESI source. Because we do not observe the IgG fragment that complements the light chain, we report kinetics based on only the increase in the light chain signal and the decrease of the intact IgG precursor (see supporting information for details). As mentioned above, it is interesting that we do not observe the complementary heavy chain fragment. We suspect that this species aggregates soon after the dissociation process. At our longest times at each temperature, where the dissociation of the intact IgG precursor approaches completion, the ESI signal is lost, consistent with this idea, and the experiment is terminated.

Examples of kinetics data recorded at several temperatures (60, 62, and 65 °C) are shown in Figure 3 for the increase in the light chain abundance with increasing time (the 57 °C data are not included because the figure appears crowded; but, an example for these data can be found in the supporting information). Examination of the kinetics shows that this process follows a simple, first-order reaction rate and can be modeled with Eq. (1),

\[
B_t = I_0 \times e^{-kt},
\]

where \(B_t\) is the intensity of the light chain signal at time \(t\), \(I_0\) is the final light chain signal upon reaction completion, and \(k\) is the rate constant.

**Transition State Thermochemistry Associated with Formation of the Light Chain**

The rate constants obtained from the kinetics experiments shown above were used to generate the Arrhenius plot in Figure 4. This plot yields a pre-exponential factor \((A = 4.33 \pm 0.8 \times 10^{13} \text{ s}^{-1})\) and activation energy \((E_a = 96 \pm 28 \text{ kJ mol}^{-1})\). This value of \(A\) indicates that accessing the transition state is very efficient—occurring near the vibrational frequency that is expected for a simple bond cleavage of a small molecule. We suggest that this may indicate that the transition state involves a very localized motion associated with cleavage of the native Cys\(^{224}\)–Cys\(^{217}\) bond in the intact precursor IgG. These values can be converted into transition state thermochemistry, yielding \(\Delta G^\ddagger = 92 \pm 11 \text{ kJ mol}^{-1}\), \(\Delta H^\ddagger = 95 \pm 10 \text{ kJ mol}^{-1}\), and \(\Delta S^\ddagger = 8 \pm 1 \text{ J mol}^{-1} \text{ K}^{-1}\). The large enthalpic barrier is consistent with the cleavage of a covalent disulfide bond. Furthermore, the relatively small entropy change at the transition state suggests little change in structure. Overall, this thermochemistry is consistent with a sequential process. First, the native Cys\(^{224}\)–Cys\(^{217}\) bond in the IgG precursor is cleaved. Upon cleavage, the light chain fragment refolds such that at least one and possibly two new non-native disulfide bridges (the scrambled Cys\(^{217}\)–Cys\(^{199}\) bond that was detected and the Cys\(^{140}\)–Cys\(^{91}\) bond that we anticipate could be formed, but was not directly detected) are formed, stabilizing the light chain product. The complementary heavy chain product of dissociation, having the single reduced free Cys\(^{224}\), rapidly aggregates and is not detected in our experiments.

**Conclusions**

Variable temperature ESI and mass spectrometry have been used to investigate thermal transitions in the IgG antibody. It is found that IgG dissociates through loss of the light chain, a process that involves disulfide bond scrambling. Kinetics studies at multiple temperatures were used to determine transition state thermochemistry of \(\Delta H^\ddagger = 95 \pm 10 \text{ kJ mol}^{-1}\), \(\Delta S^\ddagger = 8 \pm 1 \text{ J mol}^{-1} \text{ K}^{-1}\), and \(\Delta G^\ddagger = 92 \pm 11 \text{ kJ mol}^{-1}\).

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