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Structural Analysis of the Effect of a Dual-FLAG Tag on Transthyretin

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Abstract

Recombinant proteins have increased our knowledge regarding the physiological role of proteins; however, affinity purification tags are often not cleaved prior to analysis, and their effects on protein structure, stability and assembly are often overlooked. In this study, the stabilizing effects of an N-terminus dual-FLAG (FT₂) tag fusion to transthyretin (TTR), a construct used in previous studies, are investigated using native ion mobility-mass spectrometry (IM-MS). A combination of collision-induced unfolding and variable-temperature electrospray ionization is used to compare gas- and solution-phase stabilities of FT₂-TTR to wild-type and C-terminal tagged TTR. Despite an increased stability of both gas- and solution-phase FT₂-TTR, thermal degradation of FT₂-TTR was observed at elevated temperatures, viz., backbone cleavage occurring between Lys9 and Cys10. This cleavage reaction is consistent with previously reported metalloprotease activity of TTR [Liz et al. 2009] and is suppressed by either metal chelation or excess zinc. This study brings to the fore the effect of affinity tag stabilization of TTR and emphasizes unprecedented detail afforded by native IM-MS to assess structural discrepancies of recombinant proteins from their wild-type counterparts.

Graphical Abstract

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.biochem.0c00105.

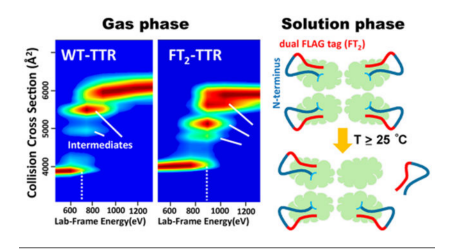
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Description of mass and charge dependency calculations, tuning parameters for mass spectrometry, schematic of the VT-ESI source, secondary structure analysis and melting experiments using CD spectroscopy, CIU plots of charge reduced WT-, CT-, and FT₂-TTR, mass spectrum of gas-phase activation of thermally heated FT₂-TTR, and mass spectrum of the isolated thermo-cleaved peptide without and with copper and zinc (PDF)



The biophysics of transthyretin (TTR), a homotetrameric protein complex with a β -sandwich fold,¹ involved in thyroxine (T₄) transport in cerebrospinal fluid (CSF) and retinol in blood, $^{2-5}$ has been the subject of many studies. $^{6-9}$ Although many studies have linked TTR to amyloid diseases, understanding the genetic and pathological maladies remains elusive. While it is generally agreed that TTR plays a key role in disease progression, how TTR structure and stability influence the mechanism(s) by which aggregates are formed remains poorly understood.¹⁰ Development of a biological mechanism for aggregation is difficult because it is still unclear as to which particular species, viz., transient intermediates (kinetic and/or thermodynamic), are involved; possibly of greater importance are the structure(s) and stabilities of these toxic species. X-ray crystallography and nuclear magnetic resonance (NMR), in concert with more traditional biophysical techniques, have provided atomistic level structure(s) for TTR. Unfortunately, traditional biophysical techniques are not well suited for studies of heterogeneous systems, especially those where low abundance transient intermediates are responsible for disease. For example, TTR functions as a metallopeptidase¹¹ when complexed with Zn(II), for which three distinct sites have been determined.¹² Excess Zn(II) binding decreases retinol transport function¹² and increases rates of fibril formation for the TTR L55P mutant.¹³ Moreover, changes in structure/stability occur upon binding Zn^{2+} , which might ultimately influence T_4 or retinol binding. The new evidence for metal-induced oxidation observed using native IM-MS underscores the importance of rigorous analytical measurements to understand protein behavior.¹⁴

An equally important aspect of understanding protein function and structure lies in the advances made in the expression and purification of recombinant proteins. Overexpressed proteins are typically expressed and isolated using a specific affinity tag, such as polyhistidine ($6 \times$ His), FLAG (DYKDDDDK), and Strep tag.^{15,16} Despite their applicability and advantages, the impact of tags on structure, stability, and function of proteins is often overlooked.¹⁷ For instance, an increase in backbone dynamics of tagged proteins has been reported¹⁷ based on an increase in B-factor of their crystal structures, which can promote in-body aggregation or decreased solubility. Booth et al. have reported slightly enhanced thermal stability of proteins without the N-terminal polyhistidine tag compared to tagged proteins.¹⁸ Yewdall and co-workers have also shown that tags can have aberrant effects on oligomerization of toroidal proteins and disfavor their self-assembly.¹⁹

Chemical modification of tags has been also reported for the well-studied polyhistidine tag in which *a-N*-6-phosphogluconoylation of the tag results in an increase in mass by 178 and 258 Da.²⁰ More recently, Liu et al. reported previously unidentified phosphorylation sites of a G-protein-gated inward rectifying potassium channel (GIRK2), and phosphorylation sites were found on both the surface exposed regions of the protein as well as the serine residues of the linker region between GIRK2 and TEV protease cleavage sites.²¹

Tag removal generally increases the purification time as well as sample purity, and several studies have reported beneficial or no effect of retaining the tag on proteins.^{17,22–25} The decision to cleave or retain the tag is difficult because molecular level details of the influence of the tags are lacking. Although structural effects of tags can be obtained using high-resolution biophysical methods such as X-ray crystallography and NMR, the requirements for crystallization and limitation on size makes these methods less appealing. Differential scanning calorimetry (DSC)²⁶ and circular dichroism (CD) spectroscopy,²⁷ which are experimentally less complicated, can be used to compare protein stabilities, but these methods do not offer molecular-level details. Fluorescence-based assays, i.e., differential scanning fluorimetry (DSF) or thermal shift assay (TSA),^{28,29} can indirectly measure protein stability through monitoring unfolding and subsequent binding of a fluorescent dye to the exposed hydrophobic residues, but the dyes used in this approach can also interact with the protein and thereby alter its stability.

Native ion mobility-mass spectrometry (IM-MS),^{30,31} on the other hand, has emerged as a fast and powerful technique for structural biology and offers information regarding stoichiometry,^{32,33} protein–protein interactions,³⁴ and the thermodynamics and kinetics of ligand binding to proteins and other biomolecules.³⁵ The measurement of shape and conformation afforded by ion mobility bodes well for native IM-MS as a potential method to screen and assess protein stability. Specifically, collision-induced unfolding (CIU) offers invaluable information regarding solvent-free unfolding pathways and stability in the gas phase.^{36–38}

Recombinant TTR has been extensively used to investigate the underlying mechanism of fibril formation or subunit exchange studies; however, purification tags are often not cleaved. $^{39-41}$ Such modifications can alter structure and stability of TTR but such effects have not been extensively investigated. In an earlier study on the subunit exchange of wild-type TTR (WT),⁴² the effects of two different tags, dual-FLAG (FT₂) and C-terminal (CT), were examined. FT₂-TTR has a lower isoelectric point (pI) (see Table 1), which facilitates the separation of subunit exchange products using ion exchange chromatography. Rappley et al. have previously reported results from chromatography studies, where a peak that eluted before tetrameric FT₂-TTR corresponded to an impurity, which was believed to come from the loss of one FT₂ tag.⁴³ Interestingly, we have also noted a backbone cleavage of FT₂-TTR, viz., Lys9-Cys10 (24 °C), which complicates subunit exchange experiments and hinders kinetics modeling.⁴²

Here, we report native IM-MS studies aimed at characterizing the impact of these fusion tags on TTR and the mechanism of the underlying cleavage for FT_2 -TTR. This study utilizes CIU measurements to analyze the gas-phase stability of TTR, and the solution-phase

stabilities are studied using a variable-temperature electrospray ionization (VT-ESI) source and collision-induced dissociation (CID). Overall, our results reveal detailed information on the impact of an affinity tag, FT_2 , on TTR structure and the mechanism of thermo-cleavage.

EXPERIMENTAL SECTION

Materials.

Alcohol dehydrogenase (ADH), concavalin A (ConA), pyruvate kinase (PK), sodium iodide, zinc acetate, copper acetate, tartaric acid, *N*-ethylmaleimide (NEM), tris (2-carboxyethyl)phosphine hydrochloride (TCEP), triethylammonium acetate (TEAA), diethylenetriaminepentaacetic acid (DTPA), and formic acid (FA) were purchased from Sigma-Aldrich (St. Louis, MO). Ammonium acetate (AA) was purchased from EMD (Millipore). Thyroxine (T₄), dimethyl sulfoxide (DMSO), acetonitrile (ACN), methanol (MeOH), and platinum wire (99.9%) were purchased from Alfa Aesar (Ward Hill, MA).

Protein Purification.

Methodologies used for the expression of the TTR variants have been described previously. ⁴² Briefly, plasmids for TTR variants were transformed into *E. coli* cells. Colonies were grown in LB media (IBI Scientific) at 37 °C until an OD₆₀₀ of 0.6–1.0. Isopropyl β -*d*-1thiogalactopyranoside (IPTG) (Enzo Life Sciences Inc.) was used at a final concentration of 0.5 mM to induce cells. Incubation was performed for 3–4 h at 37 °C. Cells were then harvested and centrifuged to remove cellular debris and loaded on a HisTrap HP column. Bound proteins were eluted with high imidazole buffer (500 mM) and immediately loaded to a desalting column (HiPrep 26/10). The His tag and fused protein (MBP for WT- and FT₂-TTR and GFP for CT-TTR) were cleaved using TEV protease overnight at 4 °C. Sizeexclusion chromatography (SEC) was used to separate aggregates and cleaved tags from TTR variants, and purified proteins were flash frozen and stored at –80 °C.

Native Ion Mobility-Mass Spectrometry.

A Synapt G2 traveling wave ion mobility separation (TWIMS) instrument (Waters, U.K.) was used for the native IM-MS experiments. A static nano-ESI source and in-house pulled borosilicate capillary tips were used to introduce TTR samples. Important tuning parameters are shown in Table S1. Mass calibration was performed using NaI, 2 mg/mL in ACN/H₂O/FA (49:49:1) for 250–3000 *m/z*. This calibration is sufficient to accurately assign and perform top down analysis for thermo-cleaved peptides (see Supporting Information). Cesium iodide, 2 mg/mL in ACN/MeOH/H₂O (30:30:40), was used as the calibrant for intact protein analysis (250–4500 *m/z*). The MS spectra for intact proteins were minimally smoothed without background subtraction.

Theoretical masses and isoelectric points (pI) of the proteins were calculated using the ExPASy server. Proteins (final concentration of 5 μ M) were buffer exchanged into 200 mM AA (pH = 6.8) using Biospin columns, 6 kDa cutoff (BioRad). TEAA (100 mM) was used to generate charge reduced mass spectra of TTR. Stock solutions of T₄ (12 mM) were prepared in DMSO and further diluted with 200 mM AA (final DMSO concentration <5%). For CIU experiments, mass selected ions were activated via collision with buffer gas (argon)

to increase the internal energy of ions, which subsequently results in gas-phase unfolding. CIUSuite 2^{44} was used to generate CIU heat maps of the collision cross section (CCS) versus lab-frame energy (E_{lab} = collision voltage × charge). CCS calculations were performed as described elsewhere.⁴⁵ ADH, ConA, and PK were used as calibrants, and corresponding CCS values were obtained from literature.⁴⁶

Circular Dichroism Spectroscopy.

CD spectra were collected using a Chirascan CD spectrometer from Applied Photophysics, Ltd. with a 10 mm path length quartz cell for a protein concentration of 3.6 μ M in 10 mM Na₂HPO₄, pH 7.4. Data are averaged from two replicates collected with a 0.5 nm bandwidth ranging from 200 to 250 nm at room temperature. CD melting experiments were performed using a Peltier sample holder to heat the sample from 30 to 90 °C at 3 °C/min at 220 nm.

Variable-Temperature Electrospray Ionization Apparatus.

VT-ESI experiments were performed using a home-built ESI heater, adapted from a similar design previously employed via Sterling et al.⁴⁷ (Figure S1). The heater consists of a capillary holder that is designed to also hold a thermally conductive, electrically insulating ceramic housing that surrounds the ESI needle. A kanthal wire is coiled around the ceramic cylinder, and the resistively heated kanthal wire transfers heat to the ceramic housing and the solution contained in the ESI needle. This device can heat from just above ambient to more than 100 °C. Solution temperatures were calibrated by inserting a microthermocouple into the ESI needle that contained only AA. Fresh TTR samples were loaded for each temperature measurement (n = 3) to avoid metal-induced oxidation during ESI.¹⁴

Quantification of Thermo-Cleaved Peptide of FT₂-TTR.

To assess the effect of solution conditions on FT₂-TTR cleavage upon heating, protein aliquots (9 uL, 13 μ M) in 200 mM ammonium acetate were mixed with 1 μ L of zinc acetate (100 μ M, 1000 μ M, and 28 mM), copper acetate (100 μ M), DTPA (10 mM), and TCEP (10 mM). Mixtures were incubated for 3 days at 50 °C and subsequently separated using Ziptip (C18 column, Thermo Fisher) and eluted with a 20 μ L solution containing 30% ACN and 0.1% FA. The measured signal for samples (n = 3) collected with three different tips was converted to a relative amount of the thermo-cleaved peptide.

Separation of the Thermo-Cleaved Peptide from FT₂-TTR.

Both size-exclusion chromatography (16/600 75 pg column eluted with 20 mM Tris, 50 mM NaCl, pH = 7.4) and hydrophobic interaction chromatography (HiTrap Phenyl HP, 5 mL column (GE Healthcare) eluted with 20 mM Tris, 500 mM NaCl, pH = 8.0) were used to separate the thermo-cleaved peptide from thermally heated FT₂-TTR. Due to the strong interaction between the thermo-cleaved peptide and truncated TTR, both experiments failed to separate them. Denaturing conditions using ZipTip with elution solution consisting of 10% acetonitrile and 0.1% FA were used to elute only the peptide (the usage of more hydrophobic solvent (ACN > 10%) led to elution of peptide and unfolded monomers). The solvent was evaporated using a Savant Speed Vac (Thermo Savant, Holbrook, NY), and 200 mM ammonium acetate was used to dissolve the peptide.

Zinc and Copper Binding Affinity to the Thermo-Cleaved Peptide.

Purified thermo-cleaved peptide samples were incubated with 2 μ M and 20 μ M zinc acetate and copper acetate. The mass spectrum of the sample incubated with 20 μ M copper showed complete binding to the peptide, whereas, for samples incubated with 2 μ M and 20 μ M zinc acetate, a similar relative abundance of the holo peak was observed. To avoid nonspecific binding during nanoelectrospray ionization, similar experiments were performed in the presence of 1 mM tartaric acid as a competitive binding specie to metals,⁴⁸ and no significant differences were observed with or without it.

RESULTS

FT₂ Tag Increases TTR Resistant to Unfold in the Gas Phase.

CIU is a process by which ions are activated by energetic collisions with a neutral target gas, such as Ar, and changes in the CCS of the ions are monitored by using IM-MS.49 This approach is increasingly used to investigate the effects of ligands,^{37,50} small molecule,^{51,52} and/or metal binding⁵³ on the structure and stabilities of the gas-phase ions. Here, CIU is used to compare the stabilities and structures of WT-, CT-, and FT₂-TTR. Figure 1 contains CIU heat maps for the 15⁺ ions of WT-, CT-, and FT₂-TTR sprayed from ammonium acetate. Unfolding transitions of the native conformer (N) produce several intermediates, labeled I1, I2 (and I3 in case of FT2-TTR), and the unfolded conformer (U). Interestingly, the CIU heat maps for CT- and WT-TTR are quite similar; however, FT₂-TTR appears to stabilize TTR against unfolding, as evidenced by the shift in the collision energy required to form the first intermediate (I₁), i.e., $E_{lab} \sim 870 \text{ eV}$ for FT₂-TTR vs ~700 eV and ~730 eV for WT and CT-TTR, respectively. Also, the CIU heat maps CT-WT and FT₂-WT clearly show that the unfolding pathway of WT-TTR is similar to CT-TTR but very different from FT₂-TTR (Figure 1D,E). The enhanced stability of FT₂-TTR is even more pronounced for charge-reduced TTRs 11⁺ ions as FT₂-TTR retains its native conformer even at the highest activation energy (1900 eV), whereas WT-TTR unfolds to I₁ at ~800 eV (see Figure S2). The increased stability of FT₂-TTR, compared to two other proteins (Figure S3), independent of their charge and mass, was confirmed (see Supporting Information). Note that we did not include data for ions that have even charges as these signals overlap with signals for dimer ions (Figure S4).

The observation of a third intermediate (I₃) in the CIU for FT₂-TTR suggests that the FT₂ tag alters the protein unfolding pathway. Ruotolo et al. previously reported the correlation between the number of domains and the observed CIU intermediates.^{54,55} The difference heat maps (Figure 1D,E) suggest that intermediates from WT and CT most likely are produced from the unfolding of similar domains of TTR. The calculated relative ratios of CCS values for unfolding products, CT- and FT₂-TTR, compared to WT-TTR (see Table S2 and Figure S5 for more detail), show that I₁ in FT₂ appears at lower CCS, compared to WT-and CT-TTR, suggesting that it might originate from the unfolding of a new domain, resulting from the FT₂ tag infusion in the protein. CD spectra of three proteins (Figure S6A) showed no significant changes in the secondary structure of TTR variants, which cannot provide any insight into observed discrepancies in CIU experiments.

Temperature Effect on the Quaternary Structure of FT₂-TTR.

VT-ESI, in combination with native IM-MS, can provide similar information as calorimetric methods,^{56–58} specifically on thermodynamics and melting point temperatures of protein and protein complexes.³⁵ This apparatus (Figure S1) is used to measure the effects of temperature on the quaternary structure of TTR and to compare the thermal stability of TTR with various tags. An increase in the average charge state (z_{avg}) of monomeric proteins has been correlated to protein unfolding;^{56,59} however, only subtle shifts in the z_{avg} of TTR upon heating are observed in this study. This observation reflects the high thermostability of TTR $(T_{\rm m} > 98 \,^{\circ}{\rm C})$;⁶⁰ however, previous studies have shown that the tetramer-dimer and dimermonomer equilibrium is highly temperature-dependent.⁴² For example, higher dissociation rates are observed for dimer to monomer reactions as the solution temperature is increased, ⁴² and higher temperatures decrease the rate of tetramer to dimer reactions. Collisioninduced dissociation (CID) of TTR results in monomer ejection in the gas phase, which requires interrupting interactions in both the dimer-dimer and monomer-monomer interfaces. Upon transition to the gas phase, ions can retain their solution-phase structure; ^{61,62} thus, CID of TTR variants originating from solutions that have different temperatures can be used to track changes in the quaternary structure upon heating. This approach provides yet another means to compare stabilities of protein complexes, viz., untagged vs tagged TTRs.

Here, in VT-ESI experiments, the relative abundances of tetrameric ions versus collision voltage are used to measure the point, denoted as I_{50} , at which 50% of the tetramer remains intact (Figure 2A). Higher I50 values indicate more energy is required for gas-phase dissociation of the protein. Similar to unfolding energies, the I_{50} values are normalized for differences in mass due to tag infusion (Figure 2B). For WT- and CT-TTR, similar normalized I₅₀ values were obtained over a wide range of temperatures (25-78 °C), indicative of their similar and high thermal stabilities (Figure 2B). An increased stability for FT₂-TTR, relative to WT-TTR (I_{50} of 0.95 \pm 0.05 vs 0.81 \pm 0.007 V/kDa), is consistent with CIU results, and similar results were obtained for charge reduced ions (Figure S7A). These results indicate higher relative gas- and solution-phase stabilities for FT₂-TTR, compared to WT- and CT-TTR. While no significant change in I_{50} was observed for WT- and CT-TTR over a range of temperatures (25-78 °C), the stability of FT2-TTR gradually decreases with increasing temperatures. Measurements shown in Figure 2 were performed in three replicates (n = 3) for each temperature (25, 42, 62, and 78 °C), and data reproducibility is diminished for FT₂-TTR at elevated temperatures (Table S3). The thermal stability of TTR samples was also investigated by using CD spectroscopy to monitor secondary structure changes (Figure S6B). In these experiments, no melting was observed for three variants in the temperature range of 30–90 °C; however, there was a similar reproducibility issue and slightly lower stability of FT₂-TTR (Figure S8).

VT-ESI was also used to examine the effects of the TTR-T₄ complex; T₄ is known to stabilize the TTR quaternary structure. At 25 °C (Figure 2B), the I_{50} values for apo/holo proteins are 0.82:1.3, 0.82:1.4, and 0.88:2.1 V/kDa for WT-, CT-, and FT₂-TTR, respectively. Note the ~2-fold increase for FT₂-TTR, compared to WT-TTR and CT-TTR, upon T₄ binding. Enhanced stabilities were also obtained for the TTR-T₄ complex,

compared to apo proteins at higher temperatures (Figure 2B). Such differences can be due to a higher affinity of FT_2 -TTR to T_4 ; however, our results showed a relatively similar binding affinity for WT and FT_2 -TTR and a slightly higher affinity for CT-TTR. Thus, these data suggest that the increased temperature destabilizes the FT_2 -TTR quaternary structure, whereas it has no effect on WT-TTR or CT-TTR.

Backbone Cleavage Observed only for FT₂-TTR upon Heating.

The temperature-dependent degradation of FT₂-TTR was reported in a previous paper, describing subunit exchange experiments.⁴² To better understand the underlying mechanism(s) and to identify the cleavage site, mass spectra of both WT- and FT₂-TTR were acquired for samples incubated at 50 °C for 18 h. Relatively broad peaks were observed for heated WT-TTR solution (3300–4500 *m/z*) compared to freshly prepared samples (Figure 3A). The observed peak broadening is due to adduction of unidentified salts to the intact molecule,⁵⁸ which are removed upon mild gas-phase collisional activation (data not shown). The native MS spectrum of heated FT₂-TTR solution contains abundant signals for multiply charged ions (2+ (1483.632 *m/z*), 3+ (989.420 *m/z*), and 4+ (594.051 *m/z*)), corresponding to an ion with MW = 2965.264 (theoretical: 2965.229) (Figure 3B). In-source activation resulted in the removal of adducts from ions in high *m/z* (3000–5500) and resolved ions, corresponding to the tetrameric FT₂-TTR with zero, one, and two truncated subunits, as well as the released intact and truncated monomers (Figure S9).

Top down tandem mass spectrometry was used to identify the 2965.264 ion as GSDYKDDDDKDYKDDDDKGPTGTGESK⁹ that is formed by the cleavage of Lys9-Cys10 in FT₂-TTR, hereafter denoted as the "thermo-cleaved" peptide (Table S4). Further evidence in support of this assignment is the number of Na⁺ and K⁺ adduct ions bound to the thermo-cleaved peptide (up to 11 sodium). The formation of salt adducts is consistent with previous studies,^{63,64} in which the number of salt adduction is correlated to the number of acidic residues (eight aspartic acids in FT₂ tag). Two other degradation products were also observed with MW = 2750.12 (theoretical: 2750.10) and 2837.15 Da (theoretical: 2837.13) formed by the cleavage of Glu7 and Ser8 with a relative abundance of ~12% and 4%, respectively. Due to their low abundance, the following discussion is limited to the major thermo-cleaved peptide (Lys9-Cys10 cleavage).

Why Does FT₂ Increase TTR Stability in Gas and Solution Phases?

Results from CIU and CID experiments demonstrate increased stability of FT_2 -TTR compared to WT- and CT-TTR at room temperature. This enhanced stability is most likely due to the FT_2 interaction with the TTR backbone, but it is unclear as to which amino acid residues are involved in the interactions. To answer this question, a solution containing WT-TTR and the thermo-cleaved peptide was incubated at 24 °C for 18 h (200 mM AA). Interestingly, binding of the thermo-cleaved peptide was not detected, which points to a potential intramolecular interaction of the FT_2 tag (DYKDDDDKDYKDDDDK) with the N-terminal residues of TTR (GPTGTGESK) in the peptide. Nevertheless, we cannot rule out the possibility of structural discrepancies between the intact and isolated tag due to the rigorous isolation process (see Experimental Section). The organic solvent and low pH

required to separate the thermo-cleaved peptide may have induced an irreversible change on the structure of the tag, which can prevent its interaction with similar residues on WT-TTR.

Metal-induced oxidation of WT-TTR¹⁴ was also observed in the presence of the thermocleaved peptide. Cys-10 oxidation and following Cys10-Pro11 cleavage results in tetrameric WT-TTR with truncated subunits shown in Figure 4 and was also evident in monomers obtained from CID of WT-TTR (data not shown). We further characterized this enhanced oxidation in the presence of the peptide (see Supporting Information), and our results revealed the higher affinity of the peptide to copper (Figure S10), which can potentially (Figure S11) increase the metal-induced oxidation of TTR as previously reported.¹⁴

Mechanism of Thermo-Cleavage Observed for FT₂-TTR.

Thermal degradation of the peptide bond in C-terminus aspartic⁶⁵ acid and N-terminus cysteine⁶⁶ has been previously reported for proteins incubated at elevated (220 °C) temperatures. Basile et al. showed that cysteine oxidation is a prerequisite for cysteine degradation and, under a nitrogen atmosphere, is completely diminished.⁶⁶ Observed thermo-cleavage (Lys9-Cys10) in our experiment can be the result of thermal decomposition of N-terminus cysteine reported by Basile; however, alkylation of Cys10 with NEM has no significant effect on the amount of thermo-cleaved product. Also, upon addition of TCEP, which inhibits Cys-10 oxidation,¹⁴ the abundance of the thermo-cleaved peptide was increased (Figure 5). These results suggest that cysteine degradation is not responsible for observed thermo-cleavage of FT₂-TTR. Moreover, WT-TTR was not susceptible to thermal decomposition, which raises the question regarding the role of the FT₂ tag on TTR propensity for thermo-decomposition in solution.

Relation between Metalloprotease Activity of TTR and Thermo-Cleavage.

TTR is a known metalloprotease for which the preferential site for substrate cleavage has been determined as Lys > Ala > Arg > Leu Met > Phe.⁶⁷ A similarity between this trend and lysine cleavage observed in our studies prompted us to investigate the correlation between metallopeptidase activity of FT2-TTR. The protease activity of untagged TTR is inhibited when metal is removed from solution using EDTA or *ortho*-phenantroline.¹¹ To test whether thermo-cleavage is related to peptidase activity of TTR, FT₂-TTR was incubated at 50 °C in the presence of DTPA (1 mM), a more potent metal chelator, and the degradation was decreased but was not completely inhibited compared to the control sample (TTR in 200 mM AA) (see Figure 5). The addition of divalent metals, i.e., Zn(II), Co(II), and Fe(II), can restore protease activity.¹¹ For example, the addition of an equimolar concentration of zinc produced more thermo-cleaved peptides (~1.7-fold), but excess zinc (1:280 molar ratio) decreased the cleavage product in agreement with previous results that excess zinc exhibits inhibitory effects.¹¹ This inhibition can be attributed to structural changes of TTR upon zinc binding to other sites, i.e., EF loop and increased backbone flexibility.¹² Unlike zinc, a low concentration of Cu(II) has shown an inhibitory effect on TTR proteolytic activity;¹¹ however, incubation of FT₂-TTR with an equimolar concentration of Cu(II) increased the amount of cleavage product (Figure 5).

DISCUSSION

Schneider et al.⁴¹ reported similar thermodynamic stability of WT and FT₂-TTR using GdmHCl unfolding experiments. More recently, FT₂-TTR was shown to be more prone to aggregation (5–10%) and slightly less thermodynamically stable than WT-TTR.⁶⁸ The CIU and VT-ESI studies reveal increased gas- and solution-phase stabilities for FT₂-TTR at room temperature (Figure 1 and Figure 2); conversely, CD spectroscopy and VT-ESI suggest a decreased thermostability that might explain previously reported aggregation propensities of FT₂-TTR.⁶⁸ A recent study by El-Baba et al.⁵⁶ illustrates increased sensitivity of native IM-MS compared to other methods; they used VT-ESI combined with native MS and IM-MS and discovered several new intermediates in the unfolding pathway of ubiquitin. Here, we used a similar approach in an effort to better assess the effects of tags on the TTR structure.

Collectively, the experimental results suggest that the mechanism of increased stability of FT_2 -TTR, compared to WT- and CT-TTR at room temperature (Figure 2), can be attributed to interactions between the FT_2 tag and N-terminal domain of TTR, which is favorable due to their close proximity. Charged residues of the N-terminus, e.g., E and K, can readily form salt bridges with the FT_2 tag by hydrogen bond interactions involving the lysine and aspartic acid residues. The increased gas-phase stability of FT_2 -TTR compared to WT- and CT-TTR in CIU experiments (Figure 1) is consistent with previous studies where electrostatic interactions are enhanced and are dominant for desolvated ions in the gas phase.⁶⁹

Similar to other proteins, the N-terminal domain of TTR is very dynamic, and only a single crystal structure is atomically resolved containing this domain (PDB 1ttc).⁷⁰ Despite such flexibility, this domain has a huge impact on TTR structure, and several hydrogen bonds have been assigned between residues Thr59 and Thr60 and N-terminal residues, i.e., Gly4, Gly6, and Ser8 (Figure S12). Previous studies also point to involvement of the N-terminus in metal binding,¹² ligand binding,^{4,71} and amyloidosis. Specifically, Cys10 is involved in the first zinc binding site,¹ and its modifications, i.e., oxidation, sulfonation, and cysteinylation, have been shown to alter amyloidogenesis of wild-type TTR and mutants.^{14,72–74} FT₂ tag interaction with the N-terminal domain of TTR can decrease the flexibility of this domain and thus increase the TTR stability.

Despite increased stability at room temperature, the observed thermo-cleavage of only the N-terminus of FT_2 -TTR indicates that amino acid residues in the tag destabilize the secondary structure of TTR at elevated temperatures. This result was further confirmed with CD spectroscopy as some batches showed slightly lower thermostability (Figure S8). A comprehensive study on proteome scale analysis of thermostability of different cells has shown higher relative abundance of aspartic acid residues in thermally unstable proteins.⁷⁵ Our data corroborate the destabilization effect of multiple aspartic acid residues on the FT₂ tag, which decreases FT₂-TTR thermal stability in solution. Our data suggest that the N-terminal domain of TTR is not a suitable region for structural modification. Also, tag-induced structural change and the infusion domain, N-terminus vs C-terminus, are needed to be studied case by case for any protein of interest.

Previously, Gouvea et al.⁷⁶ have studied the metalloprotease activity of TTR and demonstrated that a single basic residue, probably lysine, and one acidic residue are involved in the proteolysis site of TTR. Liz et al.¹¹ also identified the active site of TTR using the site-directed mutagenesis approach, where Glu89 plays a key role in stabilizing His88 in Zn coordination. Thus, several lysine and aspartic acid residues in the sequence of the FT₂ tag can influence and/or create a new binding site for the metalloprotease activity of TTR.

The fact that FT_2 -TTR is detected intact after incubation for several days at high temperatures suggests that only a fraction of protein has peptidase activity. Previous studies have shown that only ~7% of TTR is enzymatically active,¹¹ which can be due to several metal binding sites, i.e., three for zinc binding,¹² as well as conformational heterogeneity of TTR revealed via NMR studies.⁷⁷ While mechanistic details of the thermocleavage of FT₂-TTR are not resolved, inhibition of thermocleavage by DTPA and excess zinc points to correlation with the proteolysis activity of TTR. However, complete inhibition was not achieved in any solution tested in this study, which implies that other pathways are involved in the observed thermo-cleavage. Controversial results, in terms of observed increases in protease activity of FT₂-TTR in the presence of copper and reported inhibitory role of copper in TTR metalloprotease activity,¹¹ are likely due to the higher affinity of the FT₂ tag toward copper (Figure S10). Copper sequestration by the tag can diminish the inhibitory effect of copper, which we attribute to the substitution of zinc in the active site. Reduction of endogenous copper(II) to (I) by TCEP can justify more degradation of FT₂-TTR with TCEP addition, which diminishes the inhibitory effect of copper.⁷⁸

CONCLUSION

Purification tags have undoubtedly facilitated the purification of recombinant proteins; however, they can alter the dynamics and function of proteins. The results presented here clearly illustrate the stabilizing effects of the dual-FLAG tag on the quaternary structure of TTR at room temperature, whereas, at elevated temperatures, metal ions (Zn and Cu) present in the solution are directly linked to a backbone cleavage reaction between Lys9 and Cys10. While metal removal or excess zinc inhibited cleavage similar to the previously reported metalloprotease activity of TTR, equimolar zinc or copper increased the thermal cleavage. The mechanism of metal-induced cleavage was revealed by combinations of variabletemperature native MS and IM-MS. This study shows that the N-terminal domain of TTR is not suitable for affinity tag insertion. This study further illustrates the utility of native IM-MS for detailed structural analyses of proteins.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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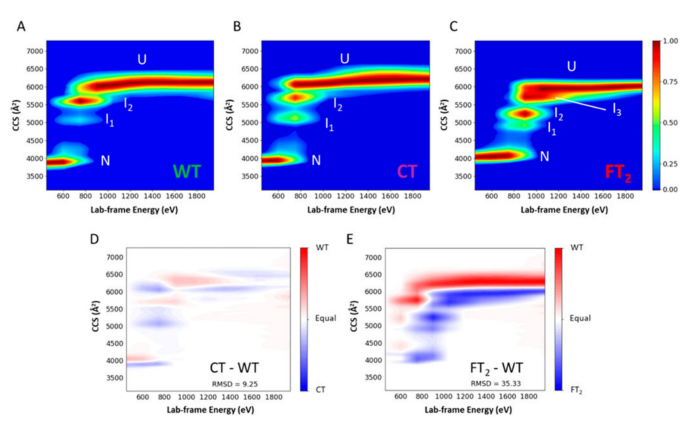


Figure 1.

Different gas-phase stabilities and intermediates were observed for TTRs (untagged and tagged) in the gas phase. Lab-frame energy is plotted versus the collision cross section (CCS) of (A) WT-TTR (B) CT-TTR, and (C) FT₂-TTR (15+) in 200 mM ammonium acetate at 25 °C. Difference CIU plot obtained from te subtraction of (D) CT-TTR and WT-TTR and (E) FT₂-TTR and WT-TTR after adjustment of the CCS value of the native conformer of WT-TTR to match the CCS value of the native conformer of CT-TTR or FT₂-TTR. CIU plots of WT- and CT-TTR are similar in terms of the number of intermediates and the relative energy required for unfolding at each step (see text for more detail), whereas FT₂-TTR shows a third intermediate and different unfolding energies.

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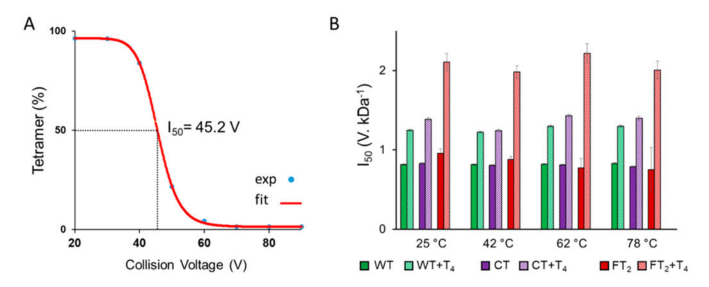


Figure 2.

Solution-phase stability analysis of TTR variants using a VT-ESI source. (A) Relative abundance of tetrameric WT-TTR to total ions as a function of collision voltage (25 °C). Data are fitted with an exponential fit to get I_{50} , which represents the voltage that 50% of tetramers remains intact. (B) I_{50} values for WT-, CT-, and FT₂-TTR at 25, 42, 62, and 78 °C without (solid) and with (dashed) T₄ in a 1:2 molar ratio (TTR/T₄). Measurements are obtained for three different replicates (n = 3) with error bars shown as ± standard deviation.

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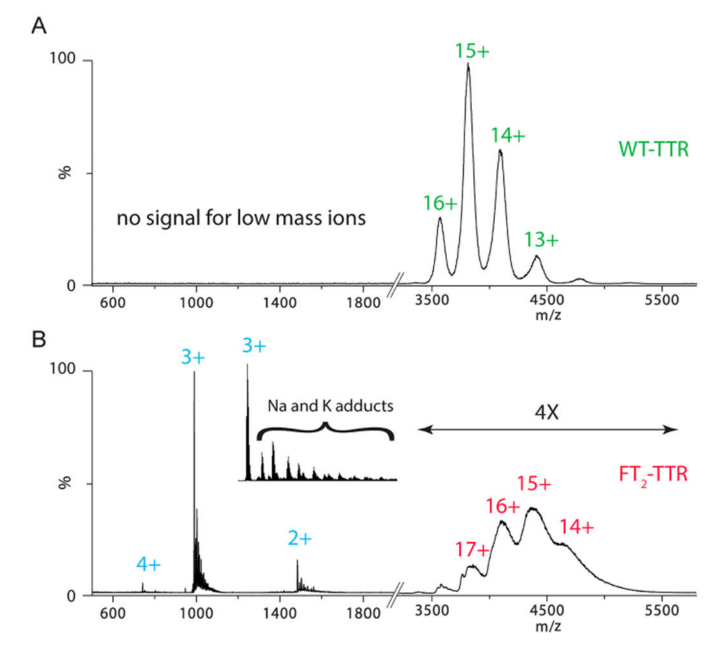


Figure 3.

Thermo-cleavage of FT₂-TTR at above ambient temperature. Mass spectra of (A) WT-TTR and (B) FT₂-TTR incubated at 50 °C for 18 h. Only FT₂-TTR upon heating produces a thermo-cleaved peptide, which is extensively bound to Na and K ions (charge states are shown in blue).

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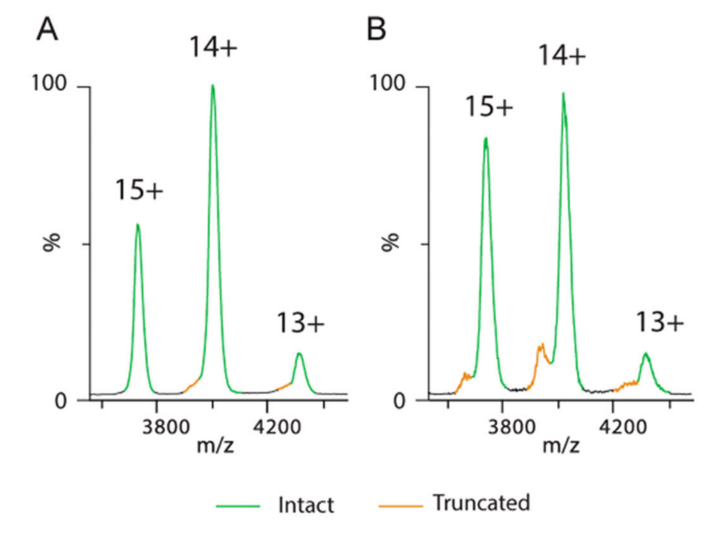


Figure 4.

Addition of the thermo-cleaved peptide results in metal-induced oxidation of WT-TTR. Mass spectra of WT-TTR incubated at 24 °C for 72 h. (A) Without and (B) with the thermocleaved peptide (GSDYKDDDDKDYKDDDDKGPTGTGESK⁹).

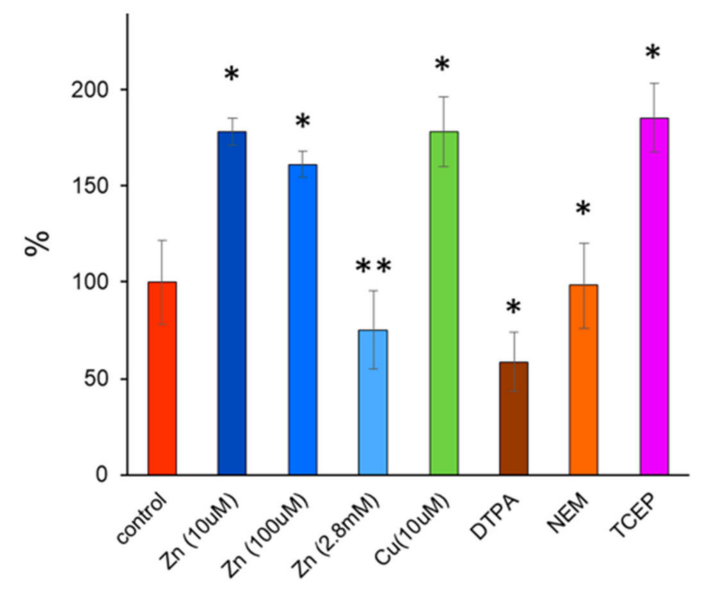


Figure 5.

Effect of metal and additives on the amount of FT₂-TTR thermo-cleavage. The relative abundance of the thermo-cleaved peptide in each solution was converted to a percentage of activity compared to the control solution (200 mM ammonium acetate). * P < 0.005 and ** P < 0.05. Error bars are shown as ± standard deviation (n = 3).

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Summary of Fusion Tags for WT-TTR, CT-TTR, and FT_2 -TTR^{*a*}

	sednence	MW (monomer, Da) pI	pI
WT-TTR	GSGPTGTGESKCPLMPKE	13905	5.31
CT-TTR	CT-TTR MGPTGTGESKCPLMPKEASGENLFYQ	14903	5.15
FT_{2} -TTR	GSDYKDDDDKDYKDDDDKGPTGPKE	15895	4.69

^aThe amino acid sequence shown in bold is the remainder after TEV protease cleavage. For CT-TTR, bold amino acids were used to increase the TTR molecular weight for the subunit exchange experiments.⁴² The italic amino acid sequence corresponds to the FT2 tag in the N-terminus of FT2-TTR