

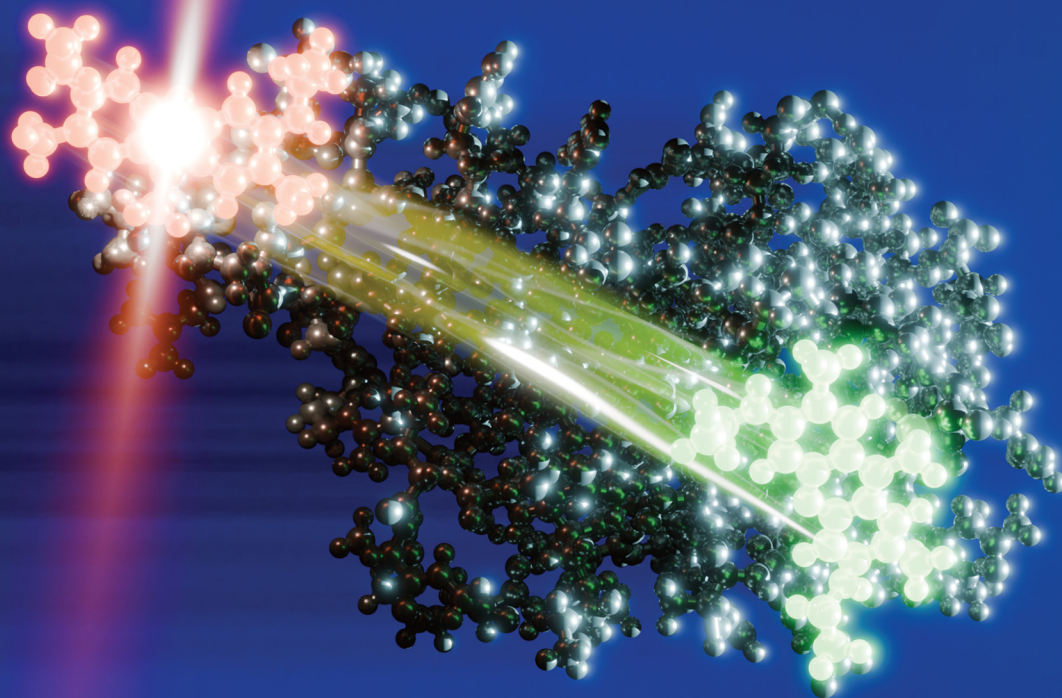
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Structural stability of chromophore-grafted Ubiquitin mutants in vacuum†

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Structural biology is witnessing a transformative era with gas-phase techniques such as native mass spectrometry (MS), ion mobility, and single-particle imaging (SPI) emerging as critical tools for studying biomolecular assemblies like protein capsids in their native states. SPI with X-ray free-electron lasers has the potential to allow for capturing atomic-resolution structures of proteins without crystallization. However, determining particle orientation during exposure remains a major challenge, compounded by the heterogeneity of the protein complexes. Gas-phase Förster resonance energy transfer (FRET) offers a promising solution to assess alignment-induced structural perturbations, providing insights into the stability of the tertiary structure under various activation methods. This study employs molecular dynamics (MD) simulations to explore chromophore integration's effect on ubiquitin's structure and alignment properties in vacuum. Ubiquitin serves as an ideal model due to its small size, well-characterized properties, and computational simplicity. By investigating chromophore placement, we identified optimal sites for monitoring gas-phase denaturation and unfolding processes, advancing SPI applications and a broader understanding of protein stability in the gas phase.

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1 Introduction

Structural biology has entered an exciting era where gas-phase techniques, such as native mass spectrometry (MS), ion mobility, and single-particle imaging (SPI), are emerging as invaluable

tools to study the structural dynamics of protein supramolecular assemblies, such as capsids, in their native and dynamic states. Recent advancements in SPI using X-ray free-electron lasers (XFELs) have underscored the importance of protein homogeneity in providing high-resolution structural insights. The principle of SPI lies in introducing individual protein molecules or complexes into an X-ray beam, where each is exposed to an ultrashort XFEL pulse. This pulse induces intense ionization and ultimately destroys the sample. However, the use of ultrafast pulses allows data collection from intact structures before disintegration, enabling the acquisition of single-particle X-ray scattering images.¹ To build a consistent 3D dataset, hundreds of thousands of measurements have to be collected to reduce signal-to-noise ratios and achieve reliable reconstructions.² The achievement of high-resolution SPI outcomes critically depends on the structural homogeneity of proteins introduced into the X-ray beam. The variability between individual protein molecules could pose a more significant challenge to SPI than radiation damage.³ Despite this, many feasibility studies have focused mainly on addressing radiation damage,^{1,4,5} with limited attention to the challenges posed by sample heterogeneity. To address this issue, the European consortium MS SPIDOC⁶ has proposed innovative methods to improve protein sample homogeneity for SPI experiments. These include native MS for mass selection, ion-mobility separation for conformational filtering, and the development of prototypes designed to couple with the EuXFEL.^{7,8} Furthermore,

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orienting proteins using external electric fields has been suggested to enhance SPI conditions.

Both static fields^{8,9} and laser fields^{10–12} have been explored for small molecules and have recently also been applied to proteins.¹³ Strong alignment fields inevitably affect protein structures,¹⁴ although simulations identify experimental conditions under which alignment occurs before denaturation.¹⁵

Beyond simulations, experimental tools are needed to evaluate alignment efficiency and measure structural perturbations induced by alignment fields. This is especially important in the gas-phase, where such tools are lacking. Förster resonance energy transfer (FRET) represents a promising technique to address these challenges.^{16,17} On the basis of dipole–dipole interactions between donor and acceptor chromophores, FRET is highly sensitive to changes in donor–acceptor distances, making it suitable for probing the effects of alignment fields on protein structure. Gas-phase FRET^{18–30} and variants of FRET, including action self-quenching,³¹ offer potential for sensitive and efficient measurements. Notably, these techniques, when applied to strategically engineered protein mutants, could elucidate stepwise unfolding processes under various activation methods, such as collisions, electric fields, or thermal activation. The grafting of chromophores at two intentional sites within the protein can provide insight into the differential stability of tertiary substructures in the gas-phase. Therefore, a deeper understanding of how chromophores influence protein stability and structural properties in the gas-phase is essential. Although similar questions have been extensively studied in solution-phase FRET, their relevance to gas-phase systems remains underexplored. Addressing this knowledge gap could provide valuable insights into the interplay between chromophore placement, protein stability, and alignment efficiency.

In this study, we employ classical molecular dynamics (MD) simulations to investigate the impact of chromophore integration on protein structure and alignment properties in vacuum. We focus on ubiquitin, a small globular protein, for several reasons: (i) its small size allows us to explore structural effects more prominently than in larger proteins; (ii) ubiquitin is well-characterized experimentally and has been studied using FRET related techniques,^{23,32} and (iii) its computational simplicity and extensive literature on vacuum stability make it an ideal model.^{3,9,33,34} We also focus on a model with the same chromophore on both mutation sites, to mimic action self-quenching experiments³¹ rather than tradition FRET experiments. This reduces complexity in both simulation and in the future real-world experiments that we plan to perform. The primary questions we aim to answer are: (1) how does the integration of chromophores affect ubiquitin's structure in the gas phase? (2) what is the impact of the placement of the chromophore on the properties of the field alignment of ubiquitin? (3) which chromophore sites are optimal for monitoring gas phase denaturation and unfolding processes? By addressing these questions, we hope to establish a robust framework for studying protein stability and alignment in the gas phase, advancing the application of SPI and related techniques to structural biology.

2 Methods

We conducted a series of classical MD simulations using the amber99sb force field,³⁵ incorporating ATTO520-cysteine adduct as the chromophore. This specific chromophore was chosen because it is known to work well in self-quenching experiments.³¹ The non-standard ATTO520-cysteine adduct was first geometry-optimized at the HF/6-31G* level³⁶ in Gaussian³⁷ with its total charge constrained to +1e. The resulting electrostatic potential was then used to fit RESP charges and used by Antechamber³⁸ to generate GAFF atom types and bonded parameters. All atom types and bonded terms were chosen to match amber99 conventions, ensuring seamless integration with the protein force field and avoiding spurious electrostatic polarization of neighboring residues. The force field parameters for the chromophore, including its functionalization onto cysteine residues, are provided in the “ATTO520 parametrization” section in the ESI† and all associated files are available in the public repository.³⁹

The initial wild-type (WT) ubiquitin structure was obtained from the Protein Data Bank (PDB ID: 1UBQ).⁴⁰ The mutants were generated by substituting, in the WT sequence, the amino acid at the position of interest with a cysteine residue, which was subsequently functionalized with the ATTO520 chromophore (Fig. 1). In each mutant, two residues were replaced by an ATTO520-tagged cysteine: residue 73 was replaced in all cases, and additionally one of the residues at position 6, 20, 35, or 48 was substituted. Accordingly, we labeled the mutants based on the non-73 replacement as ubi₆, ubi₂₀, ubi₃₅, and ubi₄₈.

To make the simulations similar to a potential experiment, an additional amino acid sequence of ELALVPR was added to the C-terminus of each variant. This is due to the anticipation of the necessary inclusion of a cleavable HIS-tag to the protein for production and purification purposes. This ELALVPR amino acid sequence would be left behind after purification and enzymatic cleavage reaction.

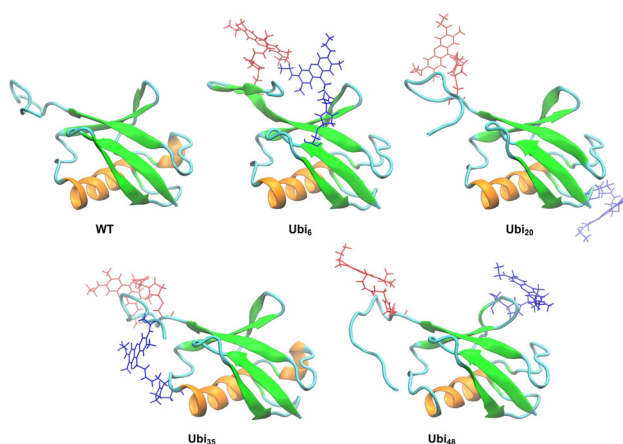


Fig. 1 Cartoon representation of ubiquitin variants structure prior to the vacuum exposure. Secondary structure elements are given, in orange, green and cyan for helix, extended and coil regions, respectively. The chromophores are shown in licorice representation. Their colors resemble the position in the linear sequence presented in Table 1.



The simulation strategy we used, identical for all the simulations, is the following. First, the ubiquitin structures are relaxed by performing a steepest descent minimization followed by a 60 ns long simulation in water at physiological pH and in the isothermal-isobaric ensemble. The temperature was kept constant at 300 K by using the v-rescale thermostat⁴¹ with a 0.1 ps coupling time. Pressure was kept constant at 1 bar by using the Berendsen barostat⁴² with a 1 ps coupling time and an isothermal compressibility of $4.5 \times 10^{-5} \text{ bar}^{-1}$. The tip4p water model was used for water molecules.⁴³ Periodic boundary conditions were imposed to the system and the Particle Mesh Ewald algorithm⁴⁴ was employed in dealing with the long-range Coulomb interactions. The MD integration time step was 2 fs. From each of these bulk simulations, five different structures, replicas, were extracted and used as starting points for the gas-phase simulations. For each replica, we then assigned the amino acids protonation states to match the net charge expected in mass spectrometry experimental data. We thus modeled any system to have a net charge of +6e, +7e, +8e. The choice of which amino acids are most likely to be protonated *in vacuo* was driven by their different gas-phase basicity.⁴⁵ A comprehensive list of all the amino acid charges for all the simulated systems is shown in Table 1. Relaxation in vacuum was performed using the steepest descent algorithm. We then performed 250 ps simulation with Berendsen thermostat,⁴² using 300 K, 400 K, 500 K, 700 K and 1000 K temperatures. Note that these “temperatures” are used purely as surrogates for increasing

levels of experimental internal energy deposition and should not be taken as literal gas-phase temperatures. The MD integration time step in vacuum was 0.5 fs. Long-range electrostatic forces were captured using no cutoffs for nonbonded interactions and no periodic boundary conditions were applied. A total of 375 independent simulations were performed (five molecules, three net charges, five replicas, and five temperatures). Bulk simulations were performed using Gromacs 2019.1⁴⁶ simulation package. Gas-phase simulations were performed using Gromacs 4.6⁴⁷ compiled in double precision.

3 Results

3.1 Validating the position of the chromophore (ATTO520) for ubiquitin

Experiments in solution have shown that adding chromophores to amino acids 6, 20, 35, 48, and 73 does not significantly affect the folding of ubiquitin, preserving a structure similar to that of the native form.^{23,48,49} To assess the dynamical behavior of these positions in gas-phase ubiquitin, we conducted simulations at various temperatures and compared the resulting structures. We used the same simulation protocol that was used in Mandl *et al.*³⁴ We then calculated the distances between all α -carbons and compared those distances between simulations at different temperatures. This way we could predict what parts of the structures of the protein are the most dynamic, *i.e.* which connections in the

Table 1 Linear sequences of WT ubiquitin and its mutants are shown. The positions of the mutations corresponding to the locations of the chromophores are marked with the letter C, their color reflect the cartoon representation shown in Fig. 1. The location of the amino acid partial charges along the linear sequence are indicated for the three different net charges of the simulated systems. The charges are given in units of elementary charge, *e*

	1234567890	1234567890	1234567890	1234567890	1234567890	1234567890	1234567890	1234567890	123
WT	MQIFVKTLTG	KTITLEVEPS	DTIENVKAKI	QDKEGIPPDQ	QLRIFAGKQL	EDGRTLSDYN	IQKESTLHLV	LRLRGGELAL	VPR
Ubi ₆	MQIFV C TLTG	KTITLEVEPS	DTIENVKAKI	QDKEGIPPDQ	QLRIFAGKQL	EDGRTLSDYN	IQKESTLHLV	LRC R GGELAL	VPR
Ubi ₂₀	MQIFVKTLTG	KTITLEVEP C	DTIENVKAKI	QDKEGIPPDQ	QLRIFAGKQL	EDGRTLSDYN	IQKESTLHLV	LRC R GGELAL	VPR
Ubi ₃₅	MQIFVKTLTG	KTITLEVEPS	DTIENVKAKI	QDKE C IPPDQ	QLRIFAGKQL	EDGRTLSDYN	IQKESTLHLV	LRC R GGELAL	VPR
Ubi ₄₈	MQIFVKTLTG	KTITLEVEPS	DTIENVKAKI	QDKEGIPPDQ	QLRIFAG C QL	EDGRTLSDYN	IQKESTLHLV	LRC R GGELAL	VPR
net charge = +6 e									
WT	00000+0000	+0000-0000	-00000+0+0	00+-0000-0	0+00000+00	000+000-00	00+-000000	0+0+000000	00+
Ubi ₆	00000+0000	+0000-0000	-00000+0+0	00+-0000-0	0+00000+00	000+000-00	00+-000000	0+++00-000	00+
Ubi ₂₀	00000+0000	+0000-0-0+	000-00+0+0	00+-0000-0	0+00000+00	-00+000-00	00+-000000	0+++000000	00+
Ubi ₃₅	00000+0000	+0000-0000	-00000+0+0	00+-+000-0	0+00000+00	-00+000-00	00+-000000	0+++00-000	00+
Ubi ₄₈	00000+0000	+0000-0000	000000+0+0	00+-0000-0	0+00000+00	-00+000-00	00+-000000	0+++00-000	00+
net charge = +7 e									
WT	00000+0000	+0000-0000	000000+0+0	00+-0000-0	0+00000+00	000+000-00	00+-000000	0+0+000000	00+
Ubi ₆	00000+0000	+0000-0000	-00000+0+0	00+-0000-0	0+00000+00	000+000-00	00+-000000	0+++000000	00+
Ubi ₂₀	00000+0000	+000000-0+	000-00+0+0	00+-0000-0	0+00000+00	-00+000-00	00+-000000	0+++000000	00+
Ubi ₃₅	00000+0000	+0000-0000	-00000+0+0	00+-+000-0	0+00000+00	-00+000-00	00+-000000	0+++000000	00+
Ubi ₄₈	00000+0000	+0000-0000	000000+0+0	00+-0000-0	0+00000+00	-00+000-00	00+-000000	0+++000000	00+
net charge = +8 e									
WT	00000+0000	+0000-0000	000000+0+0	00+-0000-0	0+00000+00	000+000000	00+-000000	0+0+000000	00+
Ubi ₆	00000+0000	+0000-0000	000000+0+0	00+-0000-0	0+00000+00	-00+000000	00+-000000	0+++000000	00+
Ubi ₂₀	00000+0000	+00000000+	000-00+0+0	00+-0000-0	0+00000+00	-00+000-00	00+-000000	0+++000000	00+
Ubi ₃₅	00000+0000	+0000-0000	-00000+0+0	00+-+000-0	0+00000+00	-00+000000	00+-000000	0+++000000	00+
Ubi ₄₈	00000+0000	+0000-0000	000000+0+0	00+-0000-0	0+00000+00	-00+000000	00+-000000	0+++000000	00+



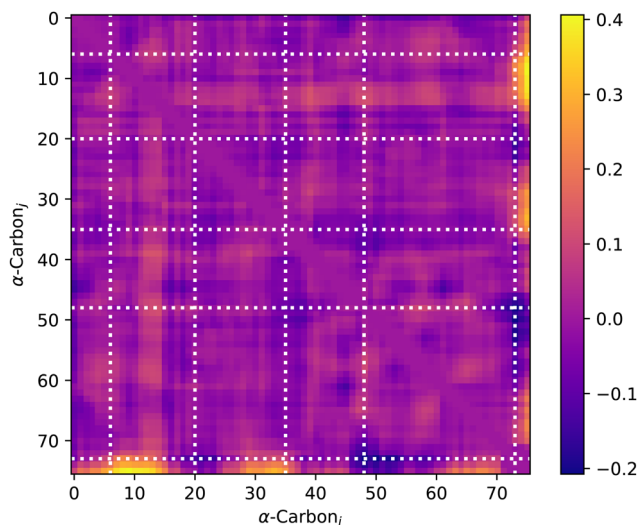


Fig. 2 Average distance difference between all alpha carbons in WT ubiquitin, comparing simulations at 325 K and 275 K. White markers indicate the positions where the chromophores are placed.

protein would be the first ones to break. Fig. 2 shows the difference in distances between the protein simulated at 325 K and 275 K. The results clearly indicate that some parts of the protein become more dynamic with temperature than others. The regions that we found to be most dynamic are the distances between alpha carbons 5–15, 28–36, and 62–71. These findings seem to be in general agreement with earlier observations, including experimental results²³ and simulations of unfolding induced by an external electric field.^{14,15}

The experimentally feasible chromophore positions (at amino acid 6, 20, 35, 48 and 73) overlap reasonably well with the protein regions our simulations identify as the most dynamics parts of the proteins, see Fig. 2. Therefore, for the rest of the study, we focused on four ubiquitin mutants (ubi₆, ubi₂₀, ubi₃₅ and ubi₄₈), along with WT ubiquitin. Henceforth, we will refer to both WT ubiquitin and the mutants as variants. Fig. 1 and Table 1 illustrate the variants and the positions in the sequence where the chromophores were introduced.

3.2 Structural stability of the variants in vacuum

After determining the optimal locations for the chromophores, we conducted simulations involving five replicas of each variant across three distinct charge states, each simulation lasting 250 ps. To replicate the energy absorption similar to a FRET experiment and potentially provide guidance for future experiments, we conducted simulations at varying temperatures. We subsequently analyzed the final trajectories using five distinct methods, aiming to understand the stability and similarities in behavior among the different variants when in a vacuum environment.

Root mean square deviation (RMSD). For every charge state (+6e, +7e, +8e) and temperature (300–1000 K), the second halves of the five replica trajectories were concatenated. Cartesian coordinates of backbone atoms were reshaped into 3*N*-dimensional vectors and subjected to *k*-means clustering with *k* = 20 (scikit-learn v1.5).⁵⁰ Clusters were ranked by population; the

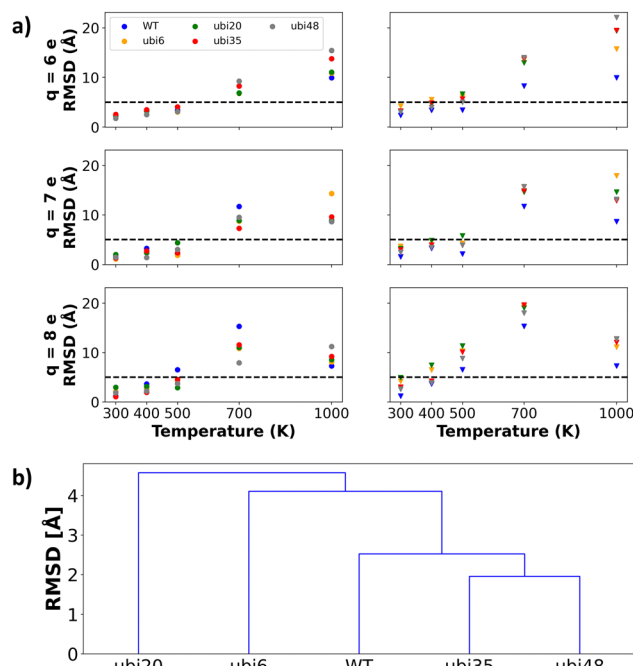


Fig. 3 (a) In the left column, intra-variant RMSD values (extracted from the diagonal of the pairwise RMSD matrix) are plotted for each ubiquitin variant, reflecting the mean structural variability across replica simulations. In the right column, WT-referenced RMSD values, calculated relative to the wild-type structure, illustrate the divergence of each variant from the reference. In both columns, each row corresponds to a system at a distinct net charge ($q = +6e, +7e, +8e$), and RMSD values are plotted as a function of simulation temperature. RMSD values are expressed in Å, with lower values (<5 Å) indicating high structural similarity and higher values (>5 Å) reflecting significant divergence. The complete pairwise RMSD matrix is presented in Fig. S1 of the ESI† while additional detailed comparisons among the replicas can be found in (Fig. S2–S16b, ESI†) global hierarchical clustering of the five ubiquitin variants using the UPGMA algorithm on the distance matrix D defined in eqn (1). Branch heights (in Å) correspond to the average C_{α} RMSD between variant pairs: shorter branches denote greater structural similarity.

five largest ones were retained, and the frame closest to each cluster centroid was extracted as the representative structure, thereby filtering out the sparsely populated clusters that correspond to conformational outliers. We used the representative structures to calculate RMSD of C_{α} atoms, and compare the values between the replicas and among variants. The results are presented in Fig. 3(a). Further, Fig. S1–S16 in ESI† show the pairwise RMSD between all four mutants and the wild-type ubiquitin, averaged over the five replica of the simulations.

From the figures, we can clearly see that for all charge states, the simulations at temperatures up to 500 K, the average RMSD between the structures is mostly below 10 Å. Within the same mutants, the RMSD is mostly below 5 Å, showing low sample heterogeneity. This holds for all three charges.

Another general trend is that the RMSD seems to increase slightly with the charge. This is in line with experimental evidence of many gaseous proteins, including ubiquitin.^{51–54} This statement is not true for all proteins at all temperatures and charges, but it is true if we consider the average RMSD for all proteins at a given charge state and temperature.



To assess the overall structural relationships among our five ubiquitin variants, we first constructed a “global” distance matrix D . Each entry

$$D_{i,j} = \frac{1}{N_{\text{cond}}} \sum_{q,T \leq 500 \text{ K}} \overline{\text{RMSD}}_{(i,j)}(q,T) \quad (1)$$

is the mean of the block-averaged C_α RMSDs between variant i and variant j , taken over all charge states q and temperatures $T \leq 500$ K. Here $\overline{\text{RMSD}}_{(i,j)}(q,T)$ denotes the average of the submatrix comparing the five frames of variant i to those of variant j under condition (q,T) , and N_{cond} is the total number of (q,T) pairs. We chose the 500 K cutoff because our initial RMSD analyses showed that, below this temperature, all variants retained the native fold across charge states—thus focusing the clustering on biologically relevant conformational variability rather than high-temperature denaturation. This global distance matrix D was then subjected to agglomerative hierarchical clustering using the Unweighted Pair Group Method with Arithmetic Mean (UPGMA⁵⁵) as implemented in the SciPy library.⁵⁶ The resulting dendrogram (Fig. 3(b)) visualizes the mutual structural similarity of the five variants, with branch heights corresponding to mean-RMSD distances in Å. Across all charges and temperatures, the dendrogram shows:

- ubi₃₅ and ubi₄₈ form the tightest pair (smallest branch height ≈ 1.9 Å), indicating these two site-specific variants are most structurally similar.

- WT joins that cluster next (merge height ≈ 2.5 Å), showing that the native protein more closely resembles the ubi_{35/48} mutants than the ubi₆ or ubi₂₀ variants.

- ubi₆ merges at a higher RMSD (≈ 4.1 Å), and ubi₂₀ is the most divergent, joining only at the top of the tree (≈ 4.7 Å).

Thus, on average across all simulated conditions, ubi₃₅ and ubi₄₈ preserve the native fold most faithfully, while ubi₆—and especially ubi₂₀—induces the largest conformational shifts. Table 1 lists the modification sites; no obvious positional trend explains the observed deviations from WT, implying a complex interplay of local and long-range effects.

Root mean square fluctuations (RMSF). The RMSF of each residue was computed for all protein variants as a function of net charge and temperature. For each variant, the five replica trajectories were aligned to their respective initial structures to eliminate global translational and rotational motion. RMSF values were then calculated for each residue over the final 125 ps of simulation time and averaged across the five replicas. This procedure yielded a converged and representative RMSF profile for each variant, allowing for direct comparison across different charge states and temperatures. The resulting averaged RMSF profiles, resolved per residue, are shown in Fig. 4. From the RMSF plots, one can notice a common trend: as the net charge increases, so does the structural deviation. Moreover, as expected, the RMSF increases with the simulation temperature. The general trend is that the dynamics are similar for the different variants and they align with the behavior of the

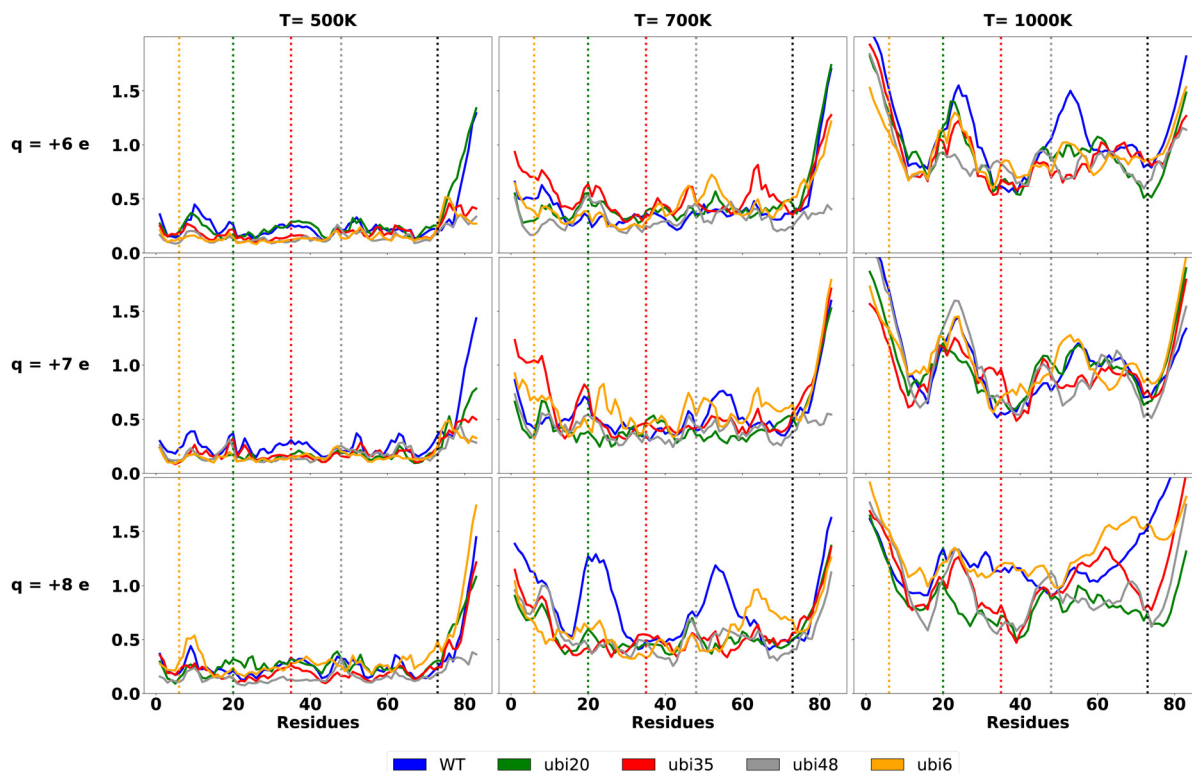


Fig. 4 RMSF of ubiquitin mutants residues as a function of temperature and net charge of the simulated systems. WT, ubi₆, ubi₂₀, ubi₃₅, and ubi₄₈, are respectively presented in blue, orange, green, red and grey. Vertical lines denote the positions in the amino acid sequence of the chromophores.



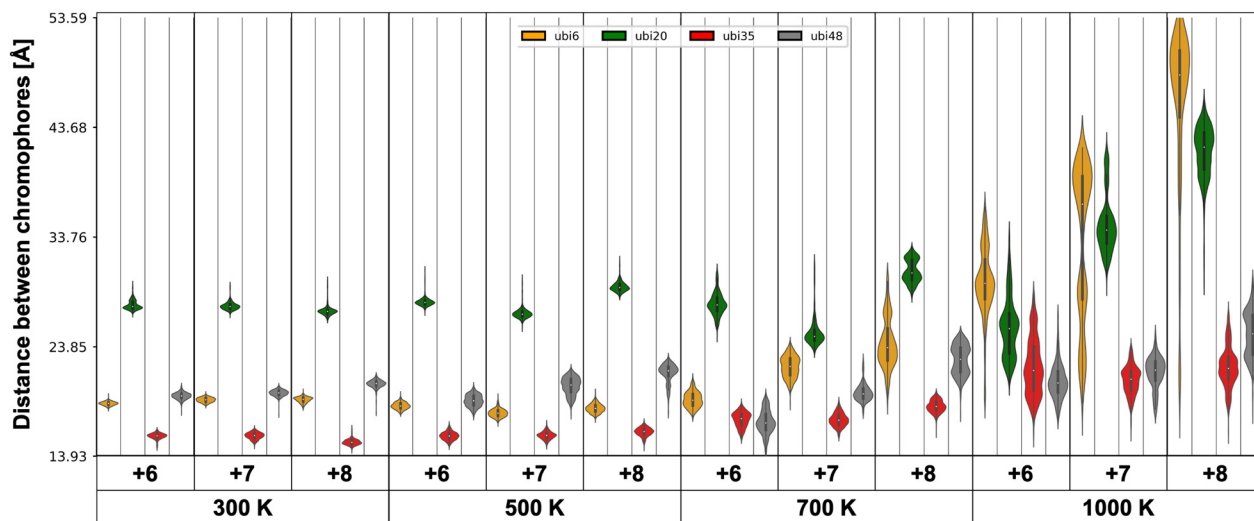


Fig. 5 Violin plots depicting the distribution of distances between chromophores are presented for simulations conducted at indicated temperatures (300 K, 500 K, 700 K, and 1000 K) and for the three different net charges of simulated systems. ubi₆, ubi₂₀, ubi₃₅, and ubi₄₈, are respectively presented in orange, green, red and grey.

WT. However, if we look closer at which residues show the highest RMSF values, we find that the WT sticks out at +8 and 700 K, and at +6 and 1000 K. In both cases, residue numbers 20–25 and 53–57 for WT show a higher RMSFs than ubiquitins attached to chromophores. At higher temperatures, it seems like the RMSF for all systems becomes similar again.

Interchromophore distance. Fig. 5 shows the distribution of the distance between the chromophores over the full simulation. The width of the violins indicates the probability of finding a system in that state during the simulation. The distance between the chromophores is defined as the distance between the two centers C_x in the chromophores.

The results indicate that at temperatures up to 700 K, the distance between the chromophores is rather well defined and stable. The effect of increasing the charge state on the interchromophore distance is barely noticeable at 500 K (and only for mutant ubi₄₈) and becomes more significant starting from

700 K. In other words, it is only at higher temperatures that charge-state-induced unfolding becomes significant. Overall, this suggests that the experimental selection of an arbitrary charge state (usually guided by pragmatic signal-to-noise considerations) may not be of paramount importance for experiments targeting the structural characterization of a given protein sample. This is promising for experiments measuring the chromophore distances, such as FRET performed at room temperature. At 1000 K, on the other hand, the spread in distances is wide, and it will be hard to draw any reasonable conclusion about the state of the molecule based solely on the chromophore distances. At this temperature the proteins are denatured, and from the figure it is clear that higher charges result in a more unfolded protein.

Secondary structure. In order to analyze the presence of secondary structure, we have plotted which residue is part of what type of secondary structure. The secondary structures

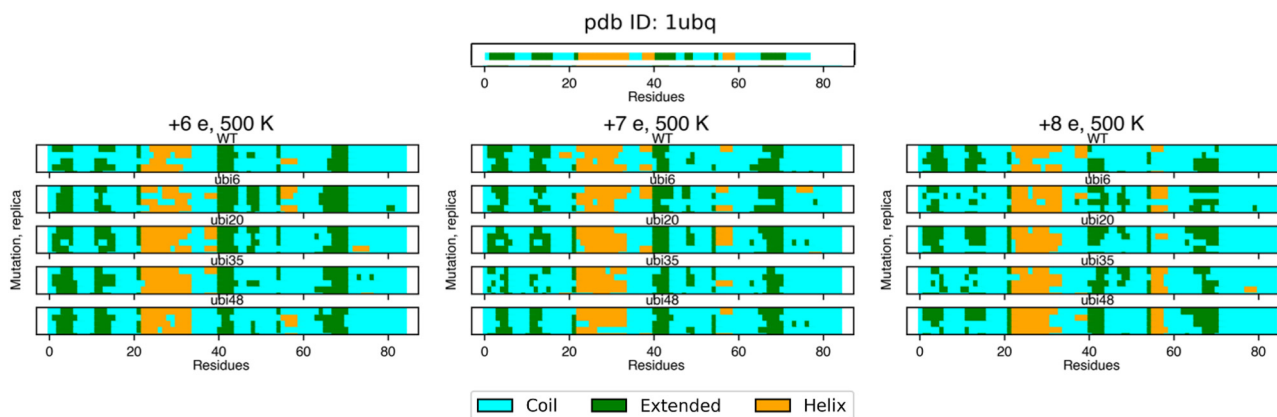


Fig. 6 Secondary structure analysis of ubiquitin is shown for the crystal structure, PDB ID: 1UBQ (top) and simulations performed at 500 K (bottom). Each line represents a replica for a specific mutant and net charge of the systems, where coil, extended, and helix structures are depicted in cyan, green, and orange, respectively.



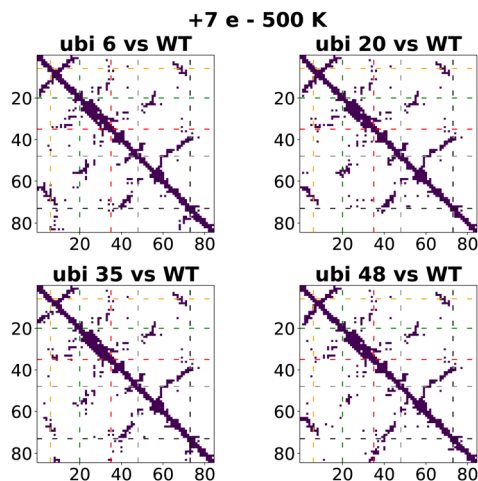


Fig. 7 Contacts matrix for simulations performed at 500 K for systems of net charge $+7e$.

shown were calculated using the DSSP algorithm,⁵⁷ and almost identical results (not shown) were also obtained using the STRIDE algorithm.⁵⁸ Fig. 6 shows the secondary structure during the simulations, for the five replicas of all the variants and charges, at 500 K. The panel on the top shows the secondary structure in the ubiquitin crystal structure (PDB ID: IUBQ⁴⁰), for reference. The figure shows that the helix is less stable, for all charge states and variants, than the extended regions (containing β -ladder and β -bridges), but in general the secondary structure seems quite stable. We did not see any evidence of the so called A-state, that has been reported in the literature.^{23,59} Instead, our simulations indicate that the secondary structure in vacuum is stable, independent of charge state.

Tertiary structure. As a last analysis we looked at the contact maps, which is a way to visualize the stability of the tertiary structure. Fig. 7 shows a comparison between the WT and the four mutants, at $+7$ and 500 K (all other comparisons are included in the ESI[†]). This analysis confirms the earlier observations (RMSD and inter-chromophore distances); the structures seem to agree well up to temperatures of 500 K.

Dipole orientation. The possibility of orienting proteins by aligning their intrinsic dipole with an external electric field has been proposed as a promising and powerful application in gas-phase imaging techniques,^{8,9,15,60–62} providing valuable insights into protein structure. For this purpose it is important to note that some chromophores, such as ATTO520 used in this study, carry a positive charge. Thus, when functionalized onto a protein residue, charged chromophores can potentially enhance the protein's intrinsic dipole, allowing for orientation with a smaller magnitude of electric field.

In Fig. 8, we present the intrinsic dipole of ubiquitin as a function of time, temperature, and net charge in the simulated systems. The dipole moment $\vec{\mu}$ was computed as

$$\vec{\mu} = \sum_{i=1}^N q_i (\vec{r}_i - \vec{r}_{\text{COM}}),$$

where q_i is the partial charge of atom i and \vec{r}_i is its position, \vec{r}_{COM} is the center of mass of the protein, and N is the total number of atoms. This definition ensures

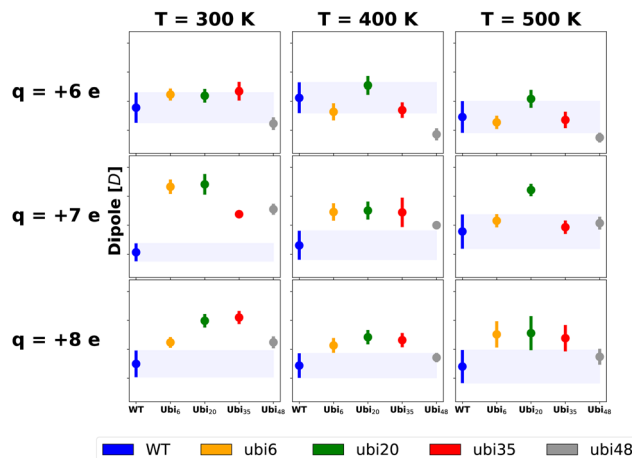


Fig. 8 Protein total dipole moment. Each panel represents a specific net charge of the systems. Within each panel, each subplots corresponds to temperatures of 300 K, 400 K, and 500 K (from top to bottom). The subplots display ubiquitin variants dipole mean values with error bars representing the standard deviation. To enhance visibility, the range of dipole values exhibited by the WT is indicated by a blue filled area.

translational invariance and is commonly used in simulations of charged systems.

We observe that for the cases of net charges $+7e$ and $+8e$ at temperatures of 300 K and 400 K, the protein's intrinsic dipole in the variants consistently surpasses that of the WT, thus confirming our speculation. In the case of a net charge of $+8e$, we also observed that the dipole of the mutants is larger compared to the WT at a temperature of 500 K. However, this trend is not observed for the net charge of $+7e$. Among the mutants, ubi_{48} exhibits the least influence on its dipole due to the presence of the chromophore. Thus, the dipole enhancement effect is more pronounced at high charge state and low temperature, which is interesting for experiments performed at room temperature on sample with similar dimensions as ubiquitin. This effect might nevertheless be less pronounced for larger proteins and protein complexes classically investigated by native mass spectrometry and which present typical mass and charge states both increased by 10- to 100-fold factor with regards to ubiquitin.

4 Discussion and conclusion

We have used molecular dynamics to perform gas-phase simulations of ubiquitin with the ATTO520 chromophore attached at four pairs of positions to investigate how the chromophore affects the dynamics and structure of the protein, comparing these results to the dynamics and structure of wild-type ubiquitin. We selected ubiquitin because it is a well-characterized protein and relatively small, which enabled us to simulate the aforementioned systems, each represented with three different net charges and at several temperatures. While the simulated time scales are shorter than those typically explored experimentally, they were sufficient to capture key dynamic and



structural features, offering insights that are highly relevant to experimental studies.

In classical molecular dynamics the charges are fixed during the simulations, and any effects caused by charge migration (due to a proton or an electron changing its position) are neglected. This includes processes such as proton hopping or charge scrambling, which are known to occur in the gas phase and can influence the detailed structure and dynamics of highly charged or partially unfolded proteins. While recent methods have begun to address such effects,⁶³ there is currently no widely accepted or computationally efficient approach to accurately model charge mobility within standard MD frameworks. Nonetheless, fixed-charge MD simulations have been shown to offer qualitatively reliable insights into gas-phase protein behavior, particularly in capturing relative stability trends.^{64–67}

Our simulations are also limited in the sense that they only simulate one charge distribution per total charge state, whereas in experiments over ensembles of molecules (such as mass spectrometry experiments where all ions with a given mass-to-charge ratio are simultaneously selected), multiple populations of charge distributions might be present at the same time, therefore also potentially affecting the secondary and tertiary structures of the protein, producing conformation ensembles within the same charge state. Our simulations only consider one charge distribution per molecule and charge state *i.e.*, our model does not allow for any charge transfer, which is a limitation. However, by including an initial equilibration in solution at physiological pH, we selected charge distributions that we believe provide a realistic and representative model of experimental conditions.

Furthermore, it is worth pointing out that the applied simulation “temperatures” serve as tunable proxies for varying levels of internal vibrational energy deposition (*e.g.*, in action-FRET experiments), rather than literal gas-phase temperatures. Because each protein’s energy uptake and redistribution depend on chromophore placement, net charge, and the specifics of the excitation mechanism, there is no one-to-one mapping between an MD temperature and a precise experimental energy input, reinforcing their role as qualitative surrogates for relative activation.

Our simulations show that introducing the chromophore affects the dynamics of the protein. For all chromophore positions and charges, a general conclusion is that the structures are well kept and similar to the WT up to 500 K (moderate energy deposition).

Moreover we notice that the mutant where the distance between the two chromophores is the most affected by temperature is ubi₆. At low temperatures (up to 500 K) the distance is below 20 Å, but at 1000 K (high energy deposition) the distance is around 30 Å, Fig. 5. This indicates that ubi₆ might be the best mutant to use for measuring the structural stability in an FRET experiment. This conclusion holds for all charge states, but is more obvious at +7e and +8e compared to +6e. In ubi₂₀ the chromophores are also separated by more than 30 Å at 1000 K, but since they start further apart than the chromophores in ubi₆ the difference is less. Interestingly, ubi₆ and ubi₂₀ are also the mutants that show the greatest structural

deviation from WT in our clustering analysis, suggesting a correlation between chromophore displacement and overall structural divergence. Judging from our simulations it is not straightforward to use chromophores at different positions in ubiquitin to monitor the temperature-induced unfolding using FRET. Our simulations do not reveal a pattern where the two chromophores in the protein are separated in a way that corresponds to their position in the amino acid sequence, at least at the temperatures when the protein structure is close the original, folded, state. At 1000 K, however the distance between the chromophores mirrors their position in amino acid chain. Adding the chromophores in general increases the dipole of the system, which could be beneficial for dipole orientation, however our simulations only show an increase for the two higher charge states (see Fig. 8). Since the structure of the proteins is not drastically affected by the presence of the chromophores, it might be a good idea to consider adding chromophores to proteins for dipole orientation.

In summary our main findings are: (i) all mutants are structurally stable at temperatures up to 500 K; (ii) below this temperature threshold, dendrogram analysis reveals that ubi₃₅ and ubi₄₈ retain conformations most similar to WT, while ubi₂₀ diverges significantly; (iii) functionalization with chromophores increases the overall dipole moment without substantially altering the protein structure, making them promising tools for monitoring and potentially assisting dipole alignment in gas-phase proteins.

Author contributions

Conceptualization: CC, EDS, JCKK, LMA, EGM; methodology: CC, EDS, TM, EGM; formal analysis and investigation: CC, EDS, TM; visualization: EDS; writing – original draft: CC, EDS; writing – review & editing: All the authors contributed to the editing of the manuscript. All authors read and approved the final manuscript.

Data availability

The data supporting this article have been included as part of the ESI,† or can be requested from the authors. The main software, GROMACS is available from <https://www.gromacs.org>.

Conflicts of interest

There are no conflicts to declare.

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