

Sampling the protonation states: the pH-dependent UV absorption spectrum of a polypeptide dyad

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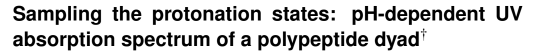
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When a chromophore interacts with several titratable molecular sites, the modeling of its photophysical properties requires to take into account all their possible protonation states. We have developed a multi-scale protocol, based on constant-pH molecular dynamics simulations coupled to QM/MM excitation energy calculations, aimed at sampling both the phase space and protonation state space of a short polypeptide featuring a tyrosine–tryptophan dyad interacting with two aspartic acid residues. We show that such a protocol is accurate enough to help in the interpretation of the experimental tyrosine UV absorption spectrum at both acidic and basic pH. Moreover, it is confirmed that radical tryptophan probably contributes to the peptide spectrum, thanks to a UV-induced electron transfer from tyrosine to tryptophan, ultimately shedding light on the complex pH-dependent behavior of the peptide spectrum.

1 Introduction

The classical atomistic modeling of a biological molecule like a polypeptide, a protein or a DNA double helix usually involves a converged sampling of its configuration space, i.e. atom positions and velocities. Molecular dynamics (MD) simulations, in which trajectories are generated by solving classical Newton equations, are clearly among the most popular available methods and take benefit of continuous improvements on both the software (eg replica-exchange, accelerated MD) and hardware sides (GPUs, Anton)¹⁻⁴. A typical MD simulation starts with some required input parameters: the force-field defining the atom-atom interaction energy and a set of atom coordinates and velocities used as initial conditions. The latter geometrical parameters are commonly obtained from available experimentally derived structures, often by means of X-ray diffraction or NMR spectroscopies⁵. However, no information is usually found regarding the distribution of the protonation states of titratable moieties like aspartic acid, lysine side-chains and similar in a protein. Hence the model needs to be complemented by an educated guess of these protonation states.

Most of the times, the protonation state of a titratable moiety is determined by comparing its pK_a with the pH of the system. Of course, experimental pK_a are macroscopic values which can barely be attributed to a given titratable site in a molecular system featuring several, and possibly interacting, sites. Hence empirical methods have been developed to give a quick and rough estimation of effective microscopic pK_a values for all the titratable sites in a system. For instance, the PropKa approach⁶ uses an available 3-dimensional structure to estimate amino-acidic pK_a values in a protein. More accurate methods based on continuum electrostatics can routinely be applied to various biomolecular systems. 7-10 On the other hand, the recently developed constant-pH molecular dynamics (CpHMD) method¹¹⁻¹⁴ has been especially designed to sample the protonation states of titratable amino-acids as a function of pH. Roughly speaking, this method introduces a Metropolis-based probability eventually allowing to change protonation states during the course of a normal MD simulation. This method has been shown to efficiently sample both the phase space and the protonation state space at the same time, given that sufficiently long trajectories are produced. Ultimately, the CpHMD simulations result in accurate pK_a predictions¹⁵.

Instead of using this information to decide on the most probable protonation states of the titratable sites, the same CpHMD trajectories can be exploited to give access to an ensemble of structures featuring a probability distribution of the protonation states at a given pH value, in agreement with the computed pK_a values. Furthermore, this ensemble can be used to calculate in a second step any molecular property whose value depends on the pH. This is precisely the target of the present study, in which the pH-dependent UV absorption spectrum of a small polypeptide is simulated for the first time. In such a case, the properties of interest (vertical excitation energies and oscillator strengths) have to be evaluated by a quantum mechanical method coupled to an approximate description of the interactions between the chromophore and its environment (QM/MM)¹⁶. To the best of



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[†] Electronic Supplementary Information (ESI) available: details of system setup and MD parameters. Simulation timings. Convergence of pK_a and statistical analysis of the trajectories. See DOI: 10.1039/cXCP00000x/

our knowledge, all the QM/MM models reported in the literature assume a single and constant (i.e. most probable) distribution of the protonation states (which will be called microstate in the following). In other words, the calculated molecular property is somehow biased towards this particular microstate.

The here-proposed CpHMD-then-QM/MM work-flow can be seen as the generalization of the routine MD-then-QM/MM approach^{17–19}, used when (classical) nuclear motion contributions to a given molecular property are needed. The successful application of such a work-flow relies on a statistically meaningful selection of snapshots along the MD trajectory. Moreover the number of such snapshots has to be large enough to ensure the convergence of the property averaged value, i.e. with a reasonable standard deviation. In the case of the CpHMD approach, reaching such a convergence is certainly more involved than in standard MD, since the phase space is complemented with the protonation state space¹³.

The subject of our study is a β -hairpin 18-mer, named Peptide M, designed by Barry *et al.*²⁰ and containing two UV-absorbing chromophores: tyrosine (Y5) and tryptophan (W14). The ultraviolet absorption spectrum features a dependency upon the pH value: the trace obtained by subtracting the tryptophan spectrum (recorded in water) from that of Peptide M (i.e. the Y5 spectrum) is always dominated in the 250-to-350-nm region by the $\pi - \pi^*$ transitions of the phenol ring, but the λ_{max} undergoes a red-shift from ~ 283 nm at pH=5 to ~ 292 nm at pH=11. There is also an additional red-shift of a few nm at pH=5 with respect to tyrosine in water. Quoting²⁰, "the red shift of the tyrosine ultraviolet spectrum in Peptide M is attributable to the close proximity of the cross-strand Y5 and W14 to form a Y5-W14 dyad." It is rightful,

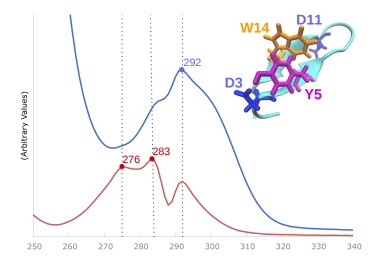


Fig. 1 Peptide M structure and experimental UV absorption spectra (in nm) 20 of Y5 in Peptide M at pH=5 (red) and pH=11 (blue).

being the experimental pK_a of the tyrosine side chain in water $\sim 10.9^{21}$, to attribute this behavior to the deprotonation occurring at basic pH; however, the presence of two other titratable residues, aspartic acids D3 and D11, contributes to the complexity of the protonation microstates landscape. The small size of the

peptide and the limited number of titratable amino-acids make this system the ideal case study for the development and testing of our method.

Besides the protonation states of the three titratable residues (D3, Y5 and D11), the proximity of the two chromophores (Y5 and W14) may result in complex interactions, e.g. charge transfer or excitonic coupling, which would also contribute to the peptide UV absorption spectrum. Hence, it is useful to summarize the experimental and theoretical results already available^{20,22}. In the former, several spectroscopic techniques have been used to characterize the peptide M. Besides the UV absorption spectra obtained at pH=5 and pH=11, NMR, ECD, EPR, UV Resonance Raman investigations have been also reported. The gathered information suggest: (i) the persistence of a β -hairpin structure, featuring a co-facial alignment of Y5 and W14 side-chains and a hydrogen bond between Y5 and R16; (ii) the existence of an excitonic coupling via a dipole-dipole interaction; (iii) the presence of radical tyrosine resulting from an electron transfer from Y5 to W14; and (iv) the indication of a charge transfer involving the radical dyad.

On the theoretical side²², MD simulations of different, but very close, peptides have been reported. Actually, in the so-called peptide A and peptide C, either histidine (peptide A) or cyclohexalanine (peptide C) replace tryptophan at position 14. Protonated (neutral), deprotonated (anionic) and radical states of tyrosine have been considered thanks to different sets of atomic point charges. While the β -hairpin-like motif is conserved (2 to 4 hydrogen bonds) when tyrosine is protonated or deprotonated, it is no longer the case with radical tyrosine which evolves to a random coil after 70 ns of simulation. Changes in microsolvation around tyrosine radical having been suggested for being responsible for such a behavior. It should be noted that the reported partial charges used to parametrize the forcefield actually indicate a phenol C-O bond dipole reversed in the radical state, with respect to both the neutral and anionic forms.

In the following, we briefly present some details regarding the CpHMD and QM/MM models we used in the present study. Then we describe the procedure to obtain the spectrum of peptide M at different pH values. We analyze the effect of the populated microstates before to conclude on the tyrosine protonation and electronic states as the main responsible for the experimentally observed λ_{max} shifts.

2 Models and computational details

2.1 QM Method

Quantum mechanical investigations on the tyrosine-tryptophan heterodimer (as well as the corresponding monomers) have been performed at the B3LYP/6-31G* level of theory, using the Gaussian16 package.²³ The geometry of the dimer corresponds to the average structure coming out of the NMR experiments by Pagba *et al.*²⁰ In the case of monomers, structures have been optimized to the minimum energy structure of the ground and the brightest excited state, using Time-Dependent DFT (TDDFT) and the Tamm-Dancoff approximation (TDA) in the region from 240 nm to 300 nm in the gas phase. All minima have been confirmed with normal mode analysis. In all cases, frequencies have been scaled by a factor of 0.96.²⁴ Vibrationally resolved spectra have been computed for monomers using the Herzberg-Teller approximation in the time-independent approach.^{25,26} The vibrational spectra have been broadened with a half-width at half-maximum of 500 cm⁻¹.

2.2 CpHMD and MD Methods

We carried out the CpHMD method simulations in explicit solvent using a discrete protonation state model as presented by Roitberg *et al.*¹³ and implemented in the AMBER16 software suite²⁷. In this method, the standard molecular dynamics steps are performed in explicit solvent, and periodically interspersed with attempts to change the protonation state in GB implicit solvent, which avoids the problem of the solvent molecule orientation; these attempts are regulated by a Metropolis Monte Carlo approach. After a successful protonation state change, which is handled by changing the charges on each atom of the residue according to the designed force field (AMBER ff14SB²⁸), the solvent molecules and non-structural ions are restored and relaxed, keeping the solute frozen; then, the velocities of the solute atoms are recovered, allowing the standard dynamics to continue.

For this type of calculation, we made use of the replica exchange technique applied along the pH-dimension (pH-REMD), in order to enhance the sampling capabilities and get an acceptable convergence^{12,29} in the given time. Our simulations were carried out using periodic boundary conditions and with a total length of 40 ns, which we considered a good compromise between accuracy in the convergence and computational time; we used 8 pH-replicas, spanning from pH 3 to pH 6 and from pH 9 to pH 12 with one pH unit as interval. More details are provided in Supplementary Informations.

For the single microstates trajectories, we used the temperature replica exchange technique (T-REMD), aiming at overcoming small energy barriers and therefore exploring exhaustively the potential energy surface; we chose 6 temperature values from 260 and 360 K, with a 20 K interval.

Finally, in order to further sample the configuration space, we performed a regular MD simulation of peptide M in its most probable protonation state at pH=5 for half a microsecond and extracted 250000 snapshots for analysis purposes.

2.3 QM/MM Method

We extracted 10000 equally spaced snapshots from the trajectories at pH 5 and 11, and coupled each frame to the corresponding protonation microstate (either protonated or deprotonated). This data allowed us to get a spatial distribution of point charges.

We chose the tyrosine phenol as QM subsystem, inserting a hydrogen link-atom between $C\beta$ and $C\gamma$, and calculated the electrostatic potential acting on each QM atom using a direct sum approach over regularly spaced images of the primitive cell used in the MD simulations. 7^3 image boxes ensure the convergence of the electrostatic potential. In the case of an electrically charged system with total charge q_t , we neutralized each image by placing a $-q_t$ charge at its center. In other words, the electrostatic

potential experienced by the QM subsystem originates from the (charged) primitive cell and from neutralized images. Electrostatic embedding of the QM subsystem is realized thanks to the ESPF method³⁰, as implemented into our local version of Gaussian09³¹.

The λ_{max} and oscillator strengths for the first four excited states were calculated using the Gaussian09 package³¹ at the TDDFT/TDA B3LYP/6-31G* level of theory; this choice is justified by the aim of seeking qualitative and not necessarily quantitative accordance with the experimental data. More involved calculations using multi-configurational multi-reference wavefunction theory for QM or polarizable forcefield for MM would improve the accuracy of the pH-dependent simulated spectra, however at a computational cost which would be prohibitive for tens of thousands of vertical excitation energy calculations.

2.4 Spectrum Elaboration

The absorption spectra were generated at room temperature with normalized Lorentzian functions from the excitation energies for the first four excited states and the corresponding oscillator strengths using Newton-X $2.0^{32,33}$, which adopts the nuclear ensemble approach³⁴. Data of the experimental spectra published in²⁰ have been kindly provided by Prof. Barry³⁵.

3 Results

3.1 The tyrosine-tryptophan dyad

In their report, Barry et al²⁰ indicate that Y5 and W14 are interacting: "the red shift of the tyrosine ultraviolet spectrum in Peptide *M* is attributable to the close proximity of the cross-strand Y5 and W14 to form a Y5-W14 dyad." As a matter of fact, the UV absorption spectrum of Y5 is perturbed at pH=5, but not at pH=11, with respect to the reference spectrum in water. Together with other spectroscopic arguments, these perturbations are interpreted as the signature of the formation of a dyad. Actually, their close proximity may promote a photoinduced electron transfer between tyrosine and transient radical tryptophan,^{36,37} which pK_a is below 5.³⁸ Assuming that W14 can be oxidized by the UV laser used in the experiment²⁰ to form the radical form that we will denote W⁻⁺ in the following, it can then react with protonated Y5 (denoted Y–OH in chemical reaction 1) at pH=5 or deprotonated Y5 (denoted Y–O⁻ in chemical reaction 2) at pH=11.

$$W^{+} + Y - OH \Longrightarrow W + Y - O^{-} + H^{+}$$
(1)

$$W^{+} + Y - O^{-} \rightleftharpoons W + Y - O^{-} \tag{2}$$

Whatever the pH value, these chemical reactions result in the generation of radical tyrosine which spectroscopic signature may be significantly different not only from the neutral (protonated) Y5 one, but also from the anionic (deprotonated) Y5 one.

In order to disentangle pH effects from Y5–W14 interaction ones, we first report results from the absorption spectra of the chromophoric moieties of tryptophan and tyrosine in different protonation states in the 240 to 340 nm energy window as reported experimentally²⁰ (see Figure 2.) The experimental absorption spectra of tryptophan and tyrosine are compared to the theoretical spectra of indole and phenol, either protonated (tyrosine most abundant form at pH=5), deprotonated (tyrosine most abundant form at pH=11) or radical.

Theoretical spectra qualitatively reproduce the order and shape of the main transitions, although they are displaced by 6-33 nm with respect to experiments. This is because the theoretical simulations do not include solvent effects or the side chain of aminoacids. For the sake of comparison, the band maxima for the theoretical spectra has been shifted to match the experimental maxima of aminoacids. The spectrum of indole (253 nm) has been shifted by 23 nm (tryptophan, exp. 276 nm), the spectrum of phenol (242 nm) has been shifted 33 nm (tyrosine pH=5, exp. 275 nm) and the spectrum of phenoxide anion (287 nm) has been shifted 6 nm (tyrosine pH=11, exp. 293 nm). The spectrum of the phenoxyl radical (266 nm) has been shifted 29 nm to the experimental value of phenoxyl at cryogenic temperatures (exp. 295 nm),³⁹ as information of tyrosyl radical in solution is not available. The intensities of all theoretical spectra have been decreased (keeping the ratio between the different peaks) to match the experimental intensity of tyrosine spectrum in solution at pH=11.

The vibrationally resolved theoretical spectrum of indole matches well the three peak structure of experimental tryptophan spectra, both at acidic and basic pH. Indeed, the indole ring remains protonated in both cases. The three peak structure involve in-plane vibrations involving the N1-C7a stretching (729, 1158 and 1407 cm⁻¹). The tryptophan spectra in the peptide is essentially equivalent to its spectrum in water. Accordingly, the coupling with the tyrosine residue is weak enough to remain hidden in the vibrational broadening.

At variance with tryptophan, the tyrosine absorption spectra features strong dependence on the pH. At pH=11, the tyrosine is deprotonated in water. The corresponding absorption spectrum matches well with the theoretical spectrum of the phenoxide anion, with a single broadened peak with little vibrational structure. The spectrum of tyrosine in the peptide M shows essentially the same band maximum, with a larger vibrational resolution. We cannot exclude the involvement of the radical phenoxyl form. However its contribution would be probably hidden in the phenoxide spectrum, due to the very similar absorption maxima. At acidic pH, the phenolic form can also coexist with the phenoxyl radical. The theoretical spectrum of phenol matches the spectrum of tyrosine at pH=5, showing a double peak vibrational shape, which origins from in-plane stretchings of C-O(H) and the benzene ring (810 and 981 cm^{-1}). However, the tyrosine spectrum in the peptide features a dissimilar structure, showing a peak at 292 nm. In Ref.²⁰, this peak was attributed to excitonic coupling. From our simulations, we cannot exclude that this peak origins from the presence of a non-negligible concentration of phenoxyl radical at acidic pH.

In order to determine the excitonic coupling strength between tryptophan and tyrosine, we compare the theoretical absorption spectrum of the dimer with respect to the monomers (see Tab. 1). We consider three dimers consisting of either phenol, phenoxide anion or phenoxyle radical with tryptophan. In this comparison, we only take the vertical electronic transition, as the vibrationally

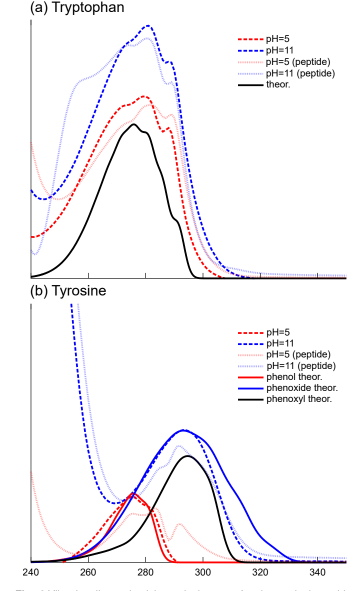


Fig. 2 Vibrationally resolved theoretical spectra for phenoxyl, phenoxide, phenol and indole, together with the experimental spectra at different pH values²⁰. Theoretical spectra (full lines) has been shifted to match the experimental band maxima. Dashed lines: spectra in water. Dotted lines: spectra in peptide M

resolved approach cannot simply be applied to the dimer due to a lack of excited state structural reference close enough to the ground state. From these results, we can infer that the excitonic couplings remain small, introducing shifts of 5 nm on average. Therefore, we suggest that the extra peak appearing in the tyrosine spectrum in the peptide M at pH=5 is due to the presence of tyrosyl radical. At pH=11, the same peak would strongly overlap with the anionic one. Of course, these results essentially come out of gas phase calculations. This is the reason why we now proceed with results obtained i) tyrosine in water and ii) tyrosine in peptide M in water.

	Trp-Phenol	Trp-Phenoxide	Trp-Phenoxyl
Trp	271 (265)	270 (265)	275 (265)
Tyr	242 (244)	348 (353)	264 (269)

Table 1 Excitation energies for the three heterodimers (in nm) compared to the same monomer excitations (in parenthesis).

3.2 Aqueous tyrosine absorption spectrum

As a first test of our CpHMD-then-QM/MM protocol, we have calculated the absorption spectra of tyrosine amino-acid in water at pH=5, 9, 10 and 11. The experimental pK_a of tyrosine side-chain is about 10. Accordingly, the calculated spectra change from a single band around 245 nm, corresponding to neutral tyrosine, to two bands with maxima at 245 nm and 318 nm, the latter being the signature of the tyrosine anionic form, which intensity increases with the pH. Conversely, the intensity of the neutral spectrum decreases with the pH. Of course, the calculated spectra

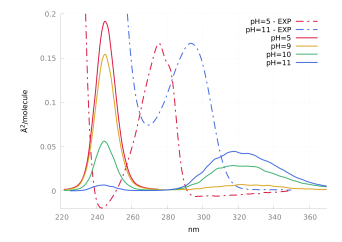


Fig. 3 Experimental $^{\rm 20}$ and calculated spectra of tyrosine in water at different pH values

are shifted with respect to the experimental ones, similarly to the gas phase vibrationally-resolved spectra reported in Section 3.1. Again, these shifts are mainly due to the low level of theory used to describe the tyrosine vertical excitations. There is now a 30 nm blue-shift for neutral tyrosine and a 27 nm red-shift for its anionic form. Even if exaggerated, the neutral-to-ionic red-shift is reproduced by our calculations. It demonstrates that the protonation state populations produced by CpHMD simulations are converged enough to catch the principal trend.

3.3 The peptide M structure

According to Pagba's RMN analysis²⁰, the peptide M adopts a stable β -hairpin structure at pH=5. We first tested the ability of the chosen forcefield to reproduce this preferred conformation by analyzing a 500 ns long MD trajectory of peptide M (D3 and D11 deprotonated, Y5 protonated) in explicit water solvent. According to Figure 4, the peptide M structure is characterized by a turn involving residues N9, G10 and D11, which is connected to an anti-parallel β -sheet that expands to Y5 and I15 most of the time. Even if the β -sheet sometimes expands until R16, this event re-

mains quite scarce. Indeed, the average distance between R16 (N ε H) and Y5 (O) is 15.2 Å with a standard deviation lower than 3 Å. This distance is much larger than the value reported experimentally (8.2 Å). On the other hand, the experimental distance (6.4 Å) between Y5 (O) and W14 (NH) is correctly reproduced in our simulation (7.1 \pm 2.0 Å on average, 6.3 Å in the case of the most abundant cluster centroid, see Figure 5).

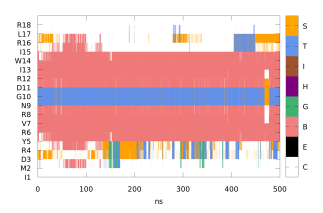


Fig. 4 Secondary structure analysis of the 500 ns trajectory (S: bend; T: turn; I: π (3-14) helix; H: α -helix; G: 3-10 helix; B: anti-parallel β -sheet; E: parallel β -sheet; C: random coil).

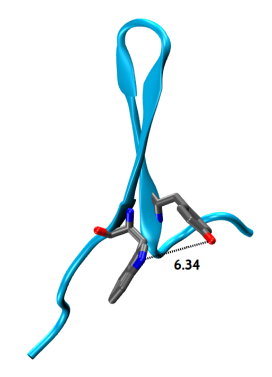


Fig. 5 Centroid structure of the principal cluster, highlighting the Y5–W14 distance.

The nature of peptide M principal secondary structure can be assessed by the number of hydrogen bonds connecting its composing residues. As shown in Figure 6, 3 to 4 hydrogen bonds exist on average, taking place mainly in R4–I15, R6–I13, R8–D11

pairs of residues. The resulting structural information clearly in-

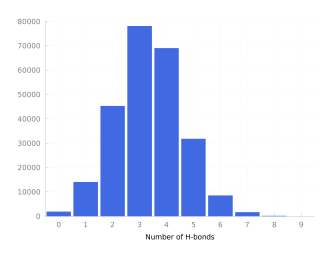


Fig. 6 Histogram analysis of the number of hydrogen bonds structuring the peptide M.

dicates that the simulated peptide M keeps a β -hairpin geometry, even if the peptide tails are somehow randomly oriented, at variance with the available NMR data²⁰.

3.4 Microstate populations

Titration curves for all titratable residues in the system are the first useful information coming out from CpHMD simulations. Their shapes not only give qualitative informations regarding the convergence of the simulations, it also may indicate non-Henderson-Hasselbalch^{40,41} behaviors possibly arising when titratable sites are strongly interacting. In the case of Peptide M (Figure 7), we first produced 12 ns long trajectories for pH ranges 3–6 and 9–12. The smooth sigmoidal shape of the 3 titration curves corresponding to D3, Y5 and D11 is a good indication of a converged exploration of both the phase and protonation state spaces. Because the sigmoidal shape does not always imply that

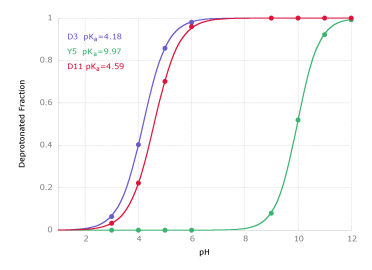


Fig. 7 Titration curves (deprotonated fraction as a function of pH) from CpHMD trajectories for the three titratable residues D3, Y5 and D11 in Peptide M.

the corresponding titratable sites are non-interacting with other ones⁴², we have performed Hill fitting^{40,43} of the titration curves, resulting in Hill factor n_h equal to 0.96, 1.07 and 0.93 for D3, Y5 and D11 respectively. In other words, these titratable residues are interacting negligibly in the protonation state space. The subsequent analysis of the fitted curves allows to determine the microscopic pK_a value of each titratable residue. As expected, the pK_a value of Y5, close to 10 and not far from the reference pK_a of tyrosine (in water), is higher than the aspartic acids ones (about 4). This result implies that at pH=5, the pH value at which the Peptide M absorption spectrum has been experimentally determined, the deprotonated form of both D3 and D11 dominates. Of course, at this pH value, Y5 is always protonated. On the other hand, at pH=11, corresponding to the second experimental absorption spectrum value, both D3 and D11 are always deprotonated while Y5 is predominantly in the deprotonated form.

After having established qualitatively the relative populations of the various microstates, we have expanded to 40 ns the trajectories corresponding to the same pH ranges 3–6 and 9–12. The D3 pK_a value converges to 4.11, while D11 pK_a is evaluated to 4.27. The detailed analysis of the microstates is reported in Figure 8. First, it should be noted that, in principle, the position

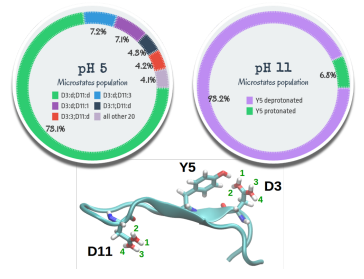


Fig. 8 D3, Y5 and D11 microstate populations at pH=5 and pH=11. At pH=5, Y5 is always protonated. At pH=11, D3 and D11 are always deprotonated. "d" stands for deprotonated while a number 1–4 stands for protonated in the corresponding position in Peptide M as indicated in the lower part).

of the acidic proton on one or the other oxygen atom of D3 (or D11) are equivalent. However, the small difference in the corresponding populations (Figure 8), about 0.1%, indicates that the trajectories are sufficiently converged to obtain reliable population estimates. Y5 is always protonated at pH=5 while D3 and D11 are predominantly (73%) deprotonated. On the other hand, there exists a noticeable population (14%) in which D11 is protonated on one of the two oxygen atoms. The same occurs with D3, but to a lower extent (8%). All other 20 possible micro-states population amounts to 4%.

At pH=11, both D3 and D11 are fully deprotonated, while Y5

is 93% deprotonated. Accordingly, only 2 micro-states are populated. Compared to pH=5 populations, this situation looks easier to handle.

With respect to the pK_a value of isolated aspartic acid in water (3.9), D3 closer pK_a value implies a slightly larger stabilization of D11 protonated form which may be attributed to enhanced interactions between D11 and other components of Peptide M. Figure 9 (complemented with Table 2) reports a selected set of average distances at pH=5 and pH=11. First, it should be noted that the

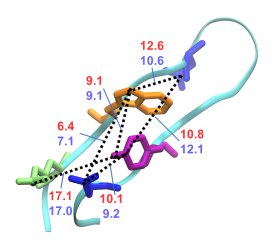


Fig. 9 Selected average distances (in Å) at pH=5 (in red) and pH=11 (in blue) between D3 (blue, bottom left), Y5 (purple), D11 (blue, top right), W14 (orange) and R16 (green).

pH does not seem to modify the average distance (9.1 Å) between Y5 and W14. However, the corresponding fluctuations are larger at acidic pH than at basic one. Regarding the distances between the two aspartic acids (D3 and D11) and the members of the dyad (Y5 and W14), they show different behaviors with respect to the pH. While Y5 is always closer to D3 than D11, the distance between Y5 and D3 decreases with increasing pH while the distance between Y5 and D11 increases at the same time. On the other hand, W14 is always much closer to D3 than D11. When going to acidic to basic pH, the distance between W14 and D3 slightly increases while the the distance between W14 and D11 decreases by 2 Å. Finally, at variance with results indicated by Pagba et al²⁰, our simulations do not show evidence of strong (hydrogen-bond) interactions between Y5 and R16, the corresponding distance being always larger than 17 Å.

3.5 Analyzing Y5 spectrum at pH=5

The computed UV absorption spectrum of Y5 in Peptide M at pH=5 is reported in Figure 10. It includes 2 bands between 200 and 300 nm, with λ_{max} values equal to 244.6 and 201.1 nm. Given the TDDFT 33 nm blue-shift already documented in Section 3.1, the shifted λ_{max} value corresponding to the first absorption reproduces almost quantitatively the 276 to 283 nm experimental value. Similarly, the second maximum is in agreement with the experimental λ_{max} at 227 nm.

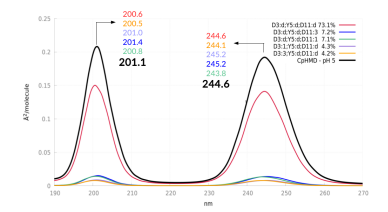


Fig. 10 Computed UV absorption spectrum for Y5 in Peptide M and individual contributions from each important microstate at pH=5 weighted by their respective population, obtained by means of CpHMD simulations.

In order to disentangle the contribution of each microstate, we have performed single microstates MD simulations with T-REMD, i.e. with a fixed distribution of protonation states, and calculated the corresponding UV spectra (Figure 10). Obviously, the main contributions originate from the most abundant microstate in which D3 and D11 are deprotonated, while Y5 is protonated. Accordingly, the pH=5 spectrum of Y5 could be satisfactorily modeled using this single microstate. However, it is interesting to have a look to the other contributions, i.e. the microstates in which either D3 or D11 are protonated. When considering the first absorption band, D3 protonation is responsible for either a 0.8 nm blue-shift or a 0.6 nm red-shift, depending on the location of the proton. Similarly, D11 protonation comes together with a ± 0.5 shift, depending on which D11 oxygen atom the proton is located on. In the case of the second absorption band, three protonation possibilities induce a 0.2 to 0.8 red-shift with respect to the most probable microstate. When considered all together, these small contributions just cancel out (first absorption band) or add a 0.5 red-shift (second band).

Variations of λ_{max} between different protonation microstates remain small. However they demonstrate that the present CpHMD-then-QM/MM protocol is able to capture subtle λ_{max} changes caused by modifications of the protonation states. The distance between Y5 and D3 increases when D3 becomes protonated (0.6 to 1 Å, depending on the oxygen atom where the proton is bonded to). The same result gets out when D11 is protonated, with a 0.8 Å lengthening (see Table 2). Of course, other interactions between Y5 and the rest of the peptide and/or the solvent are also modified when a titratable residue changes its protonation state at a given pH. Therefore it is very difficult to find qualitative relations between structural modifications and variations of λ_{max} .

3.6 Comparing Y5 spectra at pH=5 and pH=11

The computed UV absorption spectra of Y5 in Peptide M at pH=5 and pH=11 are reported in Figure 3.6 in the range 230-320 nm, together with the experimental spectra reproduced from ²⁰. For the sake of fair comparison between the various forms of Y5, it

Table 2 Selected average distances and standard deviations (in Å) at pH=5 (also decomposed according to the most important microstates, see Figure 8 for notation) and pH=11 between D3 (C_{γ}), Y5 (O) , D11 (C_{γ}), W14 (N_{ε}) and R16 (N_{ε}).

	Y5 ⋯ W14	Y5 ⋯ D3	Y5 ··· D11	W14 · · · D3	W14 · · · D11	Y5 ··· R16
pH=5	9.1 ± 2.2	10.1 ± 2.1	10.8 ± 1.3	6.4 ± 2.9	12.6 ± 1.6	17.1 ± 2.3
D3:d,D11:d	8.8 ± 0.9	9.7 ± 1.9	10.7 ± 0.8	6.8 ± 1.9	10.5 ± 0.7	19.2 ± 2.1
D3:d,D11:3	11.1 ± 1.0	10.4 ± 1.0	11.5 ± 0.9	4.5 ± 1.7	13.6 ± 1.5	18.3 ± 0.9
D3:d,D11:1	8.5 ± 0.7	10.3 ± 0.8	11.5 ± 0.8	5.9 ± 1.2	11.1 ± 0.8	15.0 ± 1.5
D3:1,D11:d	8.5 ± 0.5	10.7 ± 0.7	10.7 ± 0.7	6.6 ± 1.3	10.6 ± 0.6	20.4 ± 1.6
D3:3,D11:d	8.0 ± 0.7	10.3 ± 1.4	11.7 ± 0.8	5.5 ± 1.4	10.4 ± 0.7	17.5 ± 1.9
pH=11	9.1 ± 1.6	9.2 ± 1.7	12.1 ± 1.0	7.1 ± 3.1	10.6 ± 1.4	17.0 ± 2.5

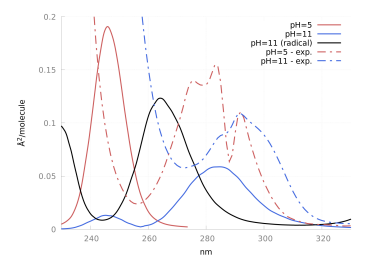


Fig. 11 Y5 spectra in Peptide M at pH=5 and pH=11 (anionic and radical Y5), both calculated and experimental 20 .

must first be noted that we needed to apply a rigid shift to the raw spectra. Indeed, the electron density localized in the phenol moiety is interacting strongly with the closest water molecules. Their MM point charges may, in turn, induce an overpolarization of the Y5 electron density. As a matter of fact, we have compared QM/MM and QM-only vertical excitation energies obtained from 10 selected snapshots, treating the closest water molecules (within a 3 Å distance from Y5 oxygen atom) either with point charges or quantum-mechanically. Inspection of Table 3 shows that i) the QM/MM approach works well when Y5 is protonated or radical, ii) an average 16 nm blue-shift has to be applied when Y5 is deprotonated. Accordingly, the spectra reported in Figure

Table 3 Influence of the level of theory (fully QM or QM polarized by water point charges (QM+q)) describing the Y5 – water interactions on the first vertical transition (averaged over 10 snapshots, in nm).

	QM	QM+q	Δ
Protonated Y5	237.7	236.3	-1.4
Deprotonated Y5	278.4	294.0	15.6
Radical Y5	251.8	250.3	-1.5

have been obtained by rigidly shifting the calculated bands using the values in Table 3. The radical spectrum has been obtained using the hypothesis that the Y5 anion form is transformed into the radical one, thanks to an electron transfer from Y5 to W14, as discussed above. Going from pH=5 to pH=11, the experimentally reported red-shift is reproduced by our calculations, whatever the form of Y5 at pH=11 (anionic or radical). With respect to the experimental spectra, all the theoretical spectra are blue-shifted, as expected from the gas phase results already discussed in 3.1. However, at variance with these gas phase spectra, now the anion and radical spectra λ_{max} are separated by 20 nm. Moreover, while the gas phase spectra take into account the homogeneous broadening due to the Y5 vibrational motion, the ones obtained for the peptide M take into account the inhomogeneous broadening due to the interaction between the chromophore and its environment. For this very reason, we can't apply the shifts discussed in 3.1. Nevertheless, comparison with the experimental spectra suggests that the Y5 radical form may contribute to both pH=5 and pH=11 spectra, in amounts which obviously depend on the experimental conditions (laser intensities, number of scans, ...) that our calculations cannot account for.

4 Conclusions

In this article, we have reported a new multi-scale protocol developed for simulating the pH-dependent photophysical properties of a peptide featuring a tyrosine-tryptophan dyad in interaction with two titratable aspartic acid residues. The modeling work-flow features two main steps: (i) the sampling of both the phase space and the protonation state space of the peptide by CpHMD simulations, (ii) the calculation of the tyrosine UV absorption spectrum by means of QM/MM calculations.

Using the replica-exchange approach, CpHMD-based pK_a values of the three titratable residues are converged in tenths of ns, with uncorrelated snapshots separated by 1 ps. Using the ESPF method, the QM/MM calculations can be achieved on thousands of protonated or deprotonated tyrosine side-chains polarized electrostatically by their environment (peptide and water molecules).

At pH=5, if tyrosine in Peptide M is protonated (neutral), its interaction with aspartic acid or aspartate residues in various minor microstates induces small deviations from the principal microstate. At pH=11, tyrosine in Peptide M is mostly deprotonated, while interacting with deprotonated aspartate residues. However, peptide M experimental UV absorption spectrum cannot be explained without assuming that (i) tryptophan can be ionized, eg by the UV-light source and (ii) radical tryptophan is reduced by an electron transferred from tyrosine which UV spectrum signature reflects its radical nature, ultimately confirming the existence of the tryptophan–tyrosine dyad.

Experimentally, it may be possible to estimate the radical concentration by modifying it (changing the laser intensity or generating the tyrosyl radical by other chemical reactions, e.g. using an organometallic species⁴⁴) and recording the corresponding absorption spectra. It may be also possible to use the spin-trapping technique⁴⁵: the quantity of trapping agent, e.g. a nitrone which would transform into a nitroxide, would be directly proportional to the tyrosyl concentration.

In principle, the reported modeling protocol can be applied to the calculation of any pH-dependent molecular property, especially when it depends on a larger protonation state space, as it is the case in proteins which may feature a very large number of titratable residues.

Conflicts of interest

There are no conflicts to declare.

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