The Gas-Phase Dipeptide Analogue Acetyl-phenylalanyl-amide: A Model for the Study of Side Chain/Backbone Interactions in Proteins

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The issue of the influence of the side chain/backbone interaction on the local conformational preferences of a phenylalanine residue in a peptide chain is addressed. A synergetic approach is used, which combines gas-phase UV spectroscopy as well as gas-phase IR/UV double-resonance experiments with DFT and post Hartree–Fock calculations. N-Acetyl-Phe-amide was chosen as a model system for which three different conformers were observed. The most stable conformer has been identified as an extended $\beta_1$ conformation of the peptide backbone. It is stabilized by a weak but significant NH–π interaction bridging the aromatic ring on the residue (i) with the NH group on residue (i+1), with the aromatic side chain being in an anti conformation. This stable conformation corresponds to the common NH(i+1)–aromatic(i) interaction encountered in proteins for the three aromatic residues (phenylalanine, tyrosine, and tryptophan), which illustrates the relevance of gas-phase investigations to structural biology issues. The two other less abundant conformers have been assigned to two $\gamma$-folded backbone conformations that differ by the orientation of the side chain. In all cases, the IR data provided spectroscopic fingerprints of these interactions. Finally, the strong conformational dependence of the fluorescence yield found for N-acetyl-Phe-amide illustrates the role of the environment on the excited-state dynamics of these species, which is often exploited by biochemists to monitor protein structural changes from tryptophan lifetime measurements.

1. Introduction

Amide–aromatic interactions have been early recognized in biology as significant interactions that are able to stabilize specific backbone secondary structures in proteins.1–3 Recent data mining investigations4 have shown that the Ar(i) $\cdots$ NH(i+1) interaction acting between an aromatic residue i and its neighbor residue i+1 (Figure 1) is the most abundant amide–aromatic interaction. Correlatively, those residues, that exhibit such an interaction, have a marked preference for an anti (trans) orientation of the aromatic side chain relative to the peptide backbone. Quantum chemists have investigated the role of such an amide–aromatic interaction by studying model molecular complexes. In particular, the binding and the most favorable geometries of an amide molecule relative to an aromatic ring, the so-called parallel and pointing conformations, have been studied.5,6 In model systems such as benzene–formamide, interaction energies on the order of 4 kcal/mol were found which are mainly due to dispersion and high-order electrostatic interactions such as quadrupole (benzene)–dipole (amide) interactions.

Figure 1. Upper panel: possible side chain/backbone local interactions for a phenylalanine residue within a peptide chain. Lower panel: schematic view of the several interactions controlling the conformational preferences of N-Ac-Phe-NH$_2$.

Gas-phase studies on aromatic amino acids with protected N- and C-termini provide an original approach to understand the role of local backbone/backbone and backbone/side chain interactions that goes beyond investigations of benzene–amide molecular complexes. In protected amino acids (Figure 1), among the nine possible conformations (according to Ramachandran terminology7 based on the $\varphi$ and $\psi$ dihedral angles), folded $\gamma$ conformations, which are similar to the central part of the $\gamma$-turns encountered in biology,8 are the only remarkable H-bonded structures expected. These $\gamma$-turns consist of three

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amino acid residues, where a H-bond bridges the backbone CO of residue (i−1) with the NH group of residue (i+1) via a seven-atom ring (referred to as C7). Recently, Lavrich et al. have provided spectroscopic evidence for the formation of such a C7 structure for the N-Ac-Ala-NH-Me dipeptide analogue in the gas phase. From the theoretical side, numerous ab initio calculations have shown that such C7 H-bonded conformations are the most stable forms in the gas phase for residues such as glycine (Gly) or alanine (Ala). In particular, the extended backbone structures (labeled βL) that correspond to the β-strands in proteins are more than 1.5 kcal/mol less stable, mainly because of the weakness of the CO(i)−NH(i) close contact attractive interaction (referred to as a C5 interaction; Figure 1).

The issue of the backbone/aromatic side chain interaction was also addressed theoretically, but so far experimental studies remain sparse. A few groups investigated modified amino acids with either tryptophan (Trp) or phenylalanine (Phe) as an aromatic side chain chromophore using UV and IR spectroscopy. These studies focused on the conformational landscape of dipeptide analogues and provided evidence for both folded (γ) and extended (βL) backbone conformations.

The goal of the present study is to investigate the role of the backbone/side chain interactions on the local conformational preferences of the peptide backbone. The small protected amino acid N-acetyl-phenylalanyl-amide (Figure 1), short: N-Ac-Phe-NH2 or NAPA, is used as a model system. The study is conducted in the gas phase to focus on intrinsic properties, free of any solvent effect. Two complementary approaches are used: quantum calculations and optical spectroscopic methods.

Thanks to progress in computer technology, quantum calculations can nowadays be performed on small model peptides, allowing us to explore their conformational landscape theoretically. When investigating weakly polar interactions such as backbone/side chain interaction, it is necessary to include electron correlation; consequently, only a few computational studies have been devoted to this issue. The goal of the accompanying paper is to give an accurate theoretical description of the here studied peptide.

On the experimental side, gas-phase spectroscopy on biological molecules made significant progress over the past decade. The combined use of laser-desorption with supersonic expansion methods leads to an efficient cooling that nevertheless enables conformational relaxation toward the lowest energy structures of the system. Resonant two-photon ionization (R2PI) spectroscopy allows then for a sensitive detection and in some cases it even allows to distinguish the UV spectral signatures of several conformers populated in the jet. The use of double-resonance techniques, such as gas-phase resonant ion-dip infrared (RID-IR) spectroscopy, can then lead to conformer specific IR spectra and in some cases to a complete characterization of the H-bonding. Following such a strategy, Gerhards and co-workers published very recently a study of N-Ac-Phe-NHMe (short: NAPMA), a dipeptide model system that is very similar to that studied in the present experiment. Zwier and co-workers investigated another dipeptide model system, N-Ac-Trp-NHMe (short: NATMA) as well as its nonmethylated derivative (N-Ac-Trp-NH2, short: NATA). In both cases, detailed assignments of the observed conformers in terms of intramolecular H-bonding are proposed; the NH−π interaction, however, was not discussed thoroughly.

The present paper presents experimental data on NAPA together with a comparison to theoretical results. The experimental strategy combines both UV and IR spectroscopy. UV spectroscopy of the Phe chromophore is sensitive enough to the phenyl environment to spectrally distinguish between gas-phase conformers, and the specific UV signature of each conformer allows us to estimate their abundance in the supersonic expansion. Conformer selective UV excitation enables us then to record IR spectra of each conformer using the RID-IR technique. Two spectral regions have been investigated: the amide A (NH stretch: 2.8−3.2 μm) and the amide I and II regions (CO stretch/NH bend: 5−8 μm). The first region is very sensitive to the presence of H-bonds and leads to a straightforward, qualitative diagnosis about the H-bonding of each conformer observed, as it directly gives the number of intramolecular H-bonds in the molecule. The combination of the two spectral regions then enables us to distinguish among several possible backbone arrangements. To derive information in particular concerning the location of the H-bonds in the molecule, the amino acids used in the study are amidated on the C terminus, even though methyl-amidated peptides have a closer resemblance to a peptidic linkage. We shall see that for such amidated peptides, IR spectroscopy brings additional information about the involvement of the amino group of the C-end amide in the H-bonding.

The experimental gas-phase conformational abundances and the measured IR spectra will be compared to the relevant theoretical results that are presented in detail in the accompanying paper. In addition, the NAPA data will be compared to theoretical results on simpler related systems that do not have an aromatic side chain (N-Ac-Gly-NH2 and N-Ac-Ala-NH2) and the results will be discussed in terms of aromatic/backbone interaction effects.

2. Experimental Section

The R2PI and RID-IR spectroscopy experiments were carried out on two molecular beam/laser desorption setups located in Saclay and in Nieuwegein. NAPA was purchased from Sigma and used without further purification. In Saclay, the peptide powder was mixed and pressed with spectroscopic-grade graphite powder to produce a pellet, which is fixed below the pulsed gas nozzle at a distance of ca. 2 mm. The second harmonic of a Nd:YAG laser was sent on the pellet surface from the end of a 550-μm diameter transport optical fiber located above it at ca. 3 mm. In Rijnhuizen, the sample powder is deposited on a graphite surface and desorbed by the mildly focused output of the fundamental output of a Nd:YAG laser. Argon was used as carrier gas (typically 3−5 bar backing pressure).

In Saclay, the infrared radiation (2.8−3.2 μm) was generated as the idler of a Nd:YAG-pumped LiNbO3 OPO (Euroscan/BM Industries). A line width of 1 cm<sup>−1</sup> was achieved by an intracavity IR Pérot-Fabry plate. The spectral scan of the OPO, controlled by a LabView operating computer, was carried out by tilting the plate, simultaneously with the LiNbO3 crystal. A typical energy of 1−3 mJ per pulse could be obtained, except in the 3470−3500 cm<sup>−1</sup> energy range where the OPO emission vanishes because of an intense IR absorption in the crystal. The IR light was mildly focused (f = 500 mm) onto the supersonic jet between the extraction plates of a time-of-flight mass spectrometer. Peptide molecules were photoionized via an R2PI scheme using the output of an excimer-pumped (Lambda-Physik EMG 200) dye laser (Lambda-Physik FL 2002E) after a time delay of 100 ns.

In Nieuwegein, mid-IR radiation was generated by a free-electron laser FELIX. Its output consists of microsecond-long bursts (macropulses) of light that come at a repetition rate of 5 Hz. Each macropulse consists of a train of picosecond-long
micropulses, with a spacing of one nanosecond between micropulses. In the present experiment, FELIX was optimized for an intense output in the 5–8-μm region (typically 50 mJ per pulse); the line width employed was typically 1% fwhm, corresponding to ~15 cm⁻¹ in the region scanned. The unfocused IR beam crossed the molecular jet and is aligned counterpropagating to the focused UV photoionization beam of a Nd:YAG-pumped dye laser (Spectra Physics GCR-100/PDL3), operating at 10 Hz.

Both R2PI spectra and RID-IR spectra were measured. In both cases, the mass-selected ion signal was acquired on a LeCroy digital oscilloscope, averaged over 100 laser shots, transferred to a PC, and measured as a function of laser frequency. In Nieuwegein, a homemade synchronization interface between FELIX and the UV laser enables to average simultaneously “IR on” and “IR off” signals on the oscilloscope. In Saclay, this is done sequentially, using a PC controlled shutter.

The setup in Saclay is also fitted with a fluorescence detector, described elsewhere, which includes as a filter a broad bandwidth monochromator followed by photomultiplier (Hamamatsu). The fluorescence excitation spectrum of NAPA was recorded under the same jet conditions as its R2PI spectrum.

3. Results

3.1. UV Spectroscopy. The frequency positions of the bands of the chromophore (the phenyl ring) in the UV spectrum are sensitive to its environment. However, it is difficult to extract structural information from these data. The Franck–Condon (FC) activity on the other hand is more interesting, as it provides information on the structural changes between ground and excited electronic states and, hence, on ring/backbone interactions.

The R2PI spectrum of mass-selected NAPA has been obtained in the region of the origin of the Phe chromophore S₁ → S₀ transition. Because of possible pollution of the NAPA mass channel by fragmented peptide hydrates or peptide dimers, the composition of the species responsible for the weak signals was carefully checked by monitoring the arrival times after the opening of the pulsed valve.

The UV spectrum of NAPA (Figure 2) is rich and, with the help of the IR/UV measurements, three spectral systems labeled A, B, and C can be distinguished. The most intense system A is composed of a low-frequency (18 cm⁻¹) vibrational progression accompanied by much weaker bands. The two other conformers B and C are less populated than the A conformer and are each characterized by a single weak UV band to the blue of the A origin.

Fluorescence excitation spectra carried out under the same conditions exhibit basically the same shape as the R2PI spectrum, apart from the fact that the UV band of species B is missing. Fluorescence decay measurements yield similar lifetime values for the A (62 ± 2 ns) and C (42 ± 2 ns) bands, values comparable to the lifetime of toluene on its origin band (76 ns). The absence of a significant fluorescence signal for the B species indicates a dramatic conformation-dependent quenching effect in which the fluorescence quantum yield of B is lower than that of A or C by at least one order of magnitude.

3.2. IR Spectroscopy and First Structural Assignment. The Amide A Region. Structural information can be obtained from IR spectroscopy because of the sensitivity of band positions to the environment of the NH or CO chromophore. This is especially the case in the amide A region where the frequencies of NH stretch modes typically range from 3420 to 3550 cm⁻¹ depending upon the residue and its environment. With the exception of NH₂ amino groups, NH oscillators on different residues are usually not strongly coupled. A red-shift of an amide A mode will be the fingerprint of an intramolecular H-bond: the stronger the bond, the larger the shift as a general rule. H-bonding to a medium acceptor like O, for example, will cause a splitting of about 120 cm⁻¹.

When one of the H-atoms is involved in a H-bond, the original symmetry and degeneracy are broken, leading to an increased splitting of the modes. This can be described by a simple model of a system of two levels interacting by vibrational coupling. The splitting is then controlled by the strength of the H-bond and the vibrational coupling constant, assumed to be independent of the H-bond. As a result the originally “symmetric” low-frequency component is the most perturbed (Figure 3); its frequency is red-shifted and this red-shift increases with the H-bond strength. Correlated, the frequency of the “antisymmetric” component decreases only slightly as the H-bond strength increases, approaching the

[Figure 2. Mass-selective one-color R2PI spectrum (top) and excitation fluorescence spectrum (bottom) of the model peptide N-Ac-Phe-NH₂ in the origin region of the S₁ → S₀ transition of the phenyl chromophore. Conformational features are identified and labeled according to the isomers deduced from the resonant ion-dip IR experiments.]

[Figure 3. Evolution of the NH stretch frequencies of a H-bonded terminal amide group of NAPA as a function of the H-bond strength (expressed in terms of red-shift) as predicted from a phenomenological vibrational coupling model. When being uncoupled and unperturbed, the two modes are degenerate. They split and one component shifts to the red when the corresponding H atom is involved in H-bonding. Vibrational coupling between the two modes causes splitting into a “symmetric” and an “antisymmetric” component. The parameters of the model (uncoupled frequency and vibrational coupling intensity) have been adjusted to fit the NH stretches of the three conformers of N-Ac-Phe-NH₂. The abscissa provides the strength of the perturbation on the bonded NH.]

Side Chain/Backbone Interactions in Proteins

TABLE 1: Experimental UV Origins and IR Frequencies of the Three Conformers Observed for N-Ac-Phe-NH₂ Assigned by Comparison with the Calculated IR NH and CO Stretch Frequencies (cm⁻¹) and Intensities (Given between Brackets in km/mol) of the Relevant Most Stable Conformations

<table>
<thead>
<tr>
<th>species</th>
<th>UV frequency/shift (cm⁻¹)</th>
<th>IR stretches</th>
<th>IR bends</th>
<th>assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NH₂sym.</td>
<td>NH₂anti.</td>
<td>CO₁</td>
<td>CO₁sym.</td>
</tr>
<tr>
<td>conf. A</td>
<td>3543</td>
<td>3517</td>
<td>1694</td>
<td>1725</td>
</tr>
<tr>
<td>conf. B</td>
<td>+107</td>
<td>3434</td>
<td>3543</td>
<td>1704</td>
</tr>
<tr>
<td>conf. C</td>
<td>+30</td>
<td>not obs.</td>
<td>3348</td>
<td>1680</td>
</tr>
</tbody>
</table>

The third band in the center, corresponding to the NH moiety on Phe, is at 3434 cm⁻¹, slightly red-shifted compared to a free NH stretch, but not compatible with a classical H-bond.

The IR spectra of species B and C (Figure 4 top panel, Table 1) differ from that of A by the presence of a red-shifted broad band in the 3340–3350 cm⁻¹ energy range and by having a much weaker red-shift of the blue most band (3520 cm⁻¹ region). The splitting between the outermost lines in the spectrum of structure B and C of 174 and 169 cm⁻¹, respectively, together with the overall red-shift is indicative for a terminal amino group engaged in significant intramolecular H-bonding. In B, a third band is located around 3438 cm⁻¹. For C, a mode corresponding to the third NH oscillator is not observed and most likely located in the gap of the OPO tuning range.

When one compares the above results to the H-bonding analysis in Figure 1, one can conclude that the two minor conformers B and C have their terminal NH₂ engaged in a H-bond in a folded C₇ backbone with a γ turnlike structure. The two conformers could differ by the type of turn (inverse or direct) or by the orientation of their side chain. In contrast, the major conformer A does not have any strong NH → CO H-bond and consequently no γ-like structure.

The Amide I and II Regions. The 5–8-μm region contains the CO stretch and NH in-plane bending modes. The NH bend vibrations are often tightly coupled with CH bends and C=–N stretches, and the assignment of the spectra in this region is generally more complicated. However, because these couplings are often conformation-dependent, the spectral patterns observed turn out to be useful information for assignment.

The 5–8-μm IR spectrum of conformer A (Figure 5, Table 1) exhibits four main bands, two in the 1620–1700 cm⁻¹ range (amide I) and two in the 1350–1600 cm⁻¹ region (amide II). The IR spectra of species B and C (Figure 5, Table 1) are very similar and resemble that of A, especially in the amide I region, apart from a change in the band intensity pattern. In the amide II region, one band (~1500 cm⁻¹) is present in the spectrum of all three species whereas the second most intense in A, at 1393 cm⁻¹, does not show up in the spectra of B and C.
because of the equatorial position of the residue compared to the backbone fold), and, (iii) the folded $\gamma_D(g^-)$ form, related to the direct $\gamma$-turns in proteins (referred to as the $C_{1ax}$ conformation, because of the axial position of Phe compared to the backbone fold). This last form is about 1 kcal/mol higher in energy than the four other forms (Table 2), but it has nevertheless been considered in our analysis as it represents the other low-energy topological basin ($\gamma_D$) able to account for intramolecular H-bonding. The other forms and basins that are either even higher in energy\cite{32} or not able to account for H-bonding (or both) have not been considered.

Harmonic frequencies and IR absorption intensities (DFT B3LYP/6-31+G(d)) have been obtained for each conformation and are compared to the experiment.\cite{32} To match the experiment, scaling factors had to be introduced. In agreement with what is anticipated from qualitative spectroscopic arguments alone, the amide A data of the A conformer can only be explained by the $\beta_L(a)$ extended conformation. The data for the B conformer, in particular the NH band at 3438 cm$^{-1}$, can only be reproduced by the $\gamma_1(g^+)$ species. Starting from this assignment, different scaling factors are used for amide A, I, and II spectral regions. Figures 4 and 5 show that a fair agreement is found between experimental and theoretical frequencies for A and B in these three regions. Discrepancies are less than 10 cm$^{-1}$ in all cases.

This good agreement suggests using theoretical spectra together with the above-mentioned scaling factors to assign the structure of the minor conformer C. Figure 4 show that the best agreement between experiment and theory is with either the $\gamma_1(g^-)$ or a $\gamma_D(g^-)$ conformation. At this stage, the amide I and II data appear as a decisive source of information, since the band intensity pattern of the $\gamma_D(g^-)$ form (Figure 5) does not fit the experiment in contrast to $\gamma_1(g^-)$ or $\gamma_1(a)$. We can thus assign the C form to the $\gamma_1(g^-)$ conformation of NAPA.

5. Discussion

5.1. Energetics of the NAPA Forms. The most abundant form of NAPA in the experiment is obviously the extended $\beta_L(a)$ form, followed by the $\gamma_1(g^+)$ and $\gamma_1(g^-)$ forms. Despite the numerous minima in the potential energy landscape of NAPA,\cite{32} only these three conformations are observed under jet conditions, which illustrates the efficiency of the structural relaxation processes during the expansion. The coexistence of the three structures should be considered as a consequence of very similar energies for these three species together with relatively low isomerization barriers (which can be crossed easily and for a significant period of time during the expansion).

The comparison with relative energies obtained from quantum chemistry calculations\cite{32} provides a stringent test of the methods and level of theory used (Table 2). DFT B3LYP/6-31+G(d) and MP2/6-31+G(d) calculations corrected for the zero-point vibrational energy (ZPE) seem to provide results that are in fair agreement with experimental abundances. Both methods show the same trends: the $\beta_L(a)$ conformation is the most stable one, followed by the three $\gamma_1$ conformations, which lie within a 1 kcal/mol energy range, starting typically 0.5 kcal/mol higher in energy (Table 2). The lowest energy $\gamma_D$ structure is significantly higher in energy, certainly higher than the lowest $\gamma_1$ conformation (Table 2) as well as the other $\beta_L$ conformations ($g^+$ and $g^-$).\cite{32} The differences in relative stabilities and ordering within the $\gamma_L$ family when comparing the two types of calculations give an indication of the order of magnitude of the precision achieved by the theoretical methods.

Comparison with calculated energies of related species, which do not exhibit an aromatic/side chain interaction such as N-Ac-

![Figure 5](image-url) Resonant ion-dip IR spectra of the three conformers of N-Ac-Phe-NH$_2$, in the amide I (CO and acetyl stretches) and II (NH bends) spectral regions together with the stick spectra of the five lowest energy conformations of NAPA, as obtained from DFT B3LYP/6-31+G(d) calculations.\cite{32} The frequencies in the stick spectra are scaled by a factor of 0.980 and 0.970 in the amide I and II regions, respectively.

**TABLE 2: Energetics of the Lowest Conformations of the N-Ac-Phe-NH$_2$, N-Ac-Gly-NH$_2$, and N-Ac-Ala-NH$_2$.**

<table>
<thead>
<tr>
<th>Form</th>
<th>$\Delta E_{B3LYP}$</th>
<th>$\Delta E_{MP2}$</th>
<th>$\Delta E_{B3LYP} + \Delta ZPE$</th>
<th>$\Delta E_{MP2} + \Delta ZPE$</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-Ac-Phe-NH$_2$</td>
<td>$\beta_L(a)$</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td></td>
<td>$\gamma_1(g^-)$</td>
<td>0.35</td>
<td>0.72</td>
<td>0.56</td>
</tr>
<tr>
<td></td>
<td>$\gamma_1(g^-)$</td>
<td>1.74</td>
<td>1.20</td>
<td>2.24</td>
</tr>
<tr>
<td>N-Ac-Gly-NH$_2$</td>
<td>$\gamma_{1D}$</td>
<td>-0.92</td>
<td>-2.23</td>
<td>-0.88</td>
</tr>
<tr>
<td></td>
<td>$\beta_L$</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>N-Ac-Ala-NH$_2$</td>
<td>$\gamma_L$</td>
<td>-1.19</td>
<td>-1.72</td>
<td>-0.74</td>
</tr>
<tr>
<td></td>
<td>$\beta_L$</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
</tbody>
</table>

4. Refined Assignment

The IR data collected for the three species observed are now compared to the results from calculations (Tables 1 and 2) reported in the accompanying paper\cite{32} to derive a more precise structure assignment.

The most stable forms calculated that are expected to be populated under jet cooling conditions (Figure 4) are listed below, labeled according to the Ramachandran terminology for the backbone conformations as well as to the classical (a, g+, and g−, for anti, gauche+, and gauche−) nomenclature for the side chain orientation (C(O)−C$^\alpha$−C$^\beta$−C$^\gamma$ side chain dihedral angle): (i) the extended $\beta_L(a)$ conformer related to the $\beta$ strandlike conformation in proteins, (ii) the three-folded $\gamma_L$ conformations, related to the so-called inverse $\gamma$-turns in proteins (also referred to as the $C_{7eq}$ conformation in the literature,
Ala-NH$_2$ and N-Ac-Gly-NH$_2$, is quite informative (Table 2). For these molecules, the folded conformations ($\gamma_L$ and $\gamma$, respectively) are more stable than the $\beta_L$ (a) conformation by typically 1 kcal/mol, in agreement with the experimentally detected conformer on N-Ac-Ala-NHMe.$^2$ This marked contrast with the NAPA case, together with the lack of stability of the $\beta_L$ (g+) and $\beta_L$ (g-) in NAPA.$^{22}$ demonstrates that the increase in stability of the NAPA $\beta_L$ (a) structure is essentially caused by the side chain. Of particular importance is the occurrence of Ar(i)$\cdots$NH(i+1) interactions, which are only possible for an anti conformation of the side chain. The corresponding stabilization energy, estimated from the change in relative energy between the $\beta_L$ and $\gamma_L$ structures when going from N-Ac-Ala-NH$_2$ to NAPA, is about 2 kcal/mol.

In the same spirit, the difference in energies of the three $\gamma_L$ conformations of NAPA of about 1 kcal/mol stems from differences in $\gamma_L$ backbone/side chain Ar(i)$\cdots$NH(i) interactions (which turn out to be weaker than the Ar(i)$\cdots$NH(i+1) interaction). The $\gamma_L$ (g+) conformation exhibits the strongest Ar(i)$\cdots$NH(i) interaction, in contrast to $\gamma_L$ (a), where this interaction cannot occur (Figure 4).

Finally, one will notice that the absence of the $\gamma_D$ conformer in the jet illustrates the efficiency of isomerization toward the $\gamma_L$ or $\beta_L$ topological basins during the expansion.

5.2. Structure and Spectroscopy of the Most Stable NAPA Conformers. The $\beta_L$ (a) Conformation (A). The calculated $\beta_L$ conformation exhibits an extended backbone structure, with two close contacts, corresponding to the NH(i+1)$\cdots$C$^\alpha$(Phe) (NH$\cdots$C$^\alpha$) and the NH(i)$\cdots$CO(i) C$_3$ interactions (DFT interatomic distances of 2.64 and 2.16 Å, respectively). The latter, although being characterized by a rather short H$\cdots$O distance, cannot be considered as a H-bond, in particular because of its nonlinear character (NH$\cdots$O angle of 107°). IR spectroscopy does exhibit the fingerprint of these weak interactions. Figure 6 shows that the stretching frequency of the NH group engaged in a C$_3$ bond is in the 3430 cm$^{-1}$ region for Gly, Ala, or Phe, independent of the residue considered and thus red-shifted by $\sim$50 cm$^{-1}$ from the free NH region (3460–3485 cm$^{-1}$), as obtained in the calculations of the $\gamma$ conformations (Figure 4). Similarly, the simultaneous red-shift of the two components of the terminal amino group, when going from the Gly or Ala residue to Phe (Figure 6), corresponds to the spectroscopic fingerprint of a weak perturbation of the terminal amino group, namely, here by the interaction of the trans NH of the amide with the C$^\alpha$ carbon of the phenyl ring. This effect on the frequencies is accounted for by the vibrational coupling model presented in Figure 3 assuming a H-bond strength of about 20 cm$^{-1}$. Such a red-shift would be expected for a methyalted amide group on the C terminus.$^{19}$

Among the three $\gamma$ conformations, only the most stable one, $\gamma_L$ (g+), exhibits a NH(i)$\cdots$C$^\alpha$(Phe) interaction. Experimentally, this leads to a red-shift of the frequency of $\sim$30 cm$^{-1}$ relative to the free NH mode for conformer B, and the DFT-optimized structure yields a H$\cdots$C$^\alpha$ distance of 2.46 Å. In the $\gamma_L$ (a) conformation, the anti orientation of the Phe side chain clearly forbids such an interaction. In the $\gamma_L$ (g-) conformation, the H$\cdots$C$_3$ DFT (B3LYP/6-31+G(d)) distance of 2.83 Å suggests a much less intense interaction than for $\gamma_L$ (g+). This is confirmed by both the absence of a red-shifted NH$_\text{NHMe}$ stretch band in the IR spectrum of conformer C (Figure 4) and the relative ordering of the MP2 energies within the $\gamma_L$ manifold (Table 2), which indicates that the energetic contribution of the NH(i)$\cdots$C$^\alpha$(Phe) interaction is about 0.8 kcal/mol.

The $\gamma_D$ Conformations. The $\gamma_D$ conformation considered, $\gamma_D$ (g-), exhibits the C$_7$ H-bond between the NH(i+1) and the CO(i-1) group, with a short H$\cdots$O interaction distance of 1.94 Å (B3LYP/6-31+G(d)) and a linearity similar to the $\gamma_L$ geometries. The axial geometry of the residue relative to the C$_7$ ring causes a significant destabilization, which explains the higher energy of this conformation with respect to the $\gamma_L$ basin. No other interaction is noticed, as illustrated by the calculated amide A IR spectra.

5.3. Comparison with Similar Dipeptide Analouges. The present results can be compared to those obtained by the same technique on the analogous system NAPMA and on model dipeptides in which Phe is replaced by another more extended aromatic side chain, such as tryptophan. For NAPMA, Gerhards and co-workers found experimentally four conformations, of which three of them were assigned to $\beta_L$(a), $\gamma_L$(g+), and...
γL(g− or a).23 This study, in qualitative agreement with the present observation of the βL(a), γL(g+), and γL(g−) conformations on NAPA, emphasizes the negligible role of the methyl capping group on the C terminal amide on the relative conformational stabilities of the system. In both NATA and NATMA, Zwier and co-workers found that βL and γL conformations also do compete: both structures are observed simultaneously in the jet and exhibit very similar DFT energies.19 A careful analysis of the reported experimental data and DFT calculations indicates that, as in the NAPA case, IR spectroscopy of these molecules in the amide A region exhibits the spectral fingerprints, specific of the weak NH—π interactions, identified in the present work. The convergence of the NAPA, NAPMA, and NATA or NATMA data therefore provides a consistent set of data that provides evidence for the role of backbone/aromatic side chain interactions in the stabilization of extended backbone structures.

5.4. Conformer-Selective Fluorescence Yield. The dramatic dependence of the fluorescence quantum yield upon the structure requires us to examine the issue of fluorescence quenching. We showed that the conformer for which the low fluorescence yield is observed is in a γL (g+) conformation, which contains a NH(i) → π interaction. This suggests that this interaction is responsible for the fluorescence quenching. Such issues have been widely addressed in the condensed phase. In particular, the existence of conformers has, for instance, been evoked to explain biexponential fluorescence behaviors of the tyrosine (Tyr) and tryptophan amino acid derivatives and dipeptides.47 Charge transfer with the amide group as an electron acceptor has also been proposed as the mechanism responsible for the quenching. In such a case, however, solvation is thought to be the fundamental process for stabilizing charge-transfer states, and the charged terminus (NH2+ or COO−) when one of the termini is not protected is also expected to play a role as it influences the electron density. In the present gas-phase experiments, none of these effects can be effective and other origins have to be considered. One should also notice that the quantum yields in the condensed phase are rather high for Trp and Tyr (~0.2) but much lower for Phe (<0.04),38 showing that the processes responsible for quenching in Phe are probably different. Dramatic phenomena in fluorescence were reported by Zwier and co-workers for the hydrogen-bonded complexes benzene—HCl and toluene—HCl.48 Two-color two-photon ionization experiments, in which the delay between the S1 → S0 laser and the ionization laser was continuously scanned, gave evidence that the hydrogen-bonding interactions lead to enhanced intersystem crossing to the triplet state. As the present dipeptide has a similar interaction as the π-type H-bond in toluene—HCl, the same interpretation in terms of enhanced intersystem crossing seems to be plausible here.

6. Conclusion

The conformational preferences on a model dipeptide in the gas phase deduced from the comparison of IR experimental and theoretical considerations enable us to provide chemical physics grounds, on the basis of characterization of local interactions, for the observed conformational distributions of residues in proteins in biological systems. The most abundant forms observed in biological systems correspond to the most stable forms observed in the modified gas-phase peptides. This suggests that simple energetic reasoning, free of any entropic, solvent, or environment effects, may be significant and relevant to biological systems.

The IR/UV double-resonance laser techniques provide spectroscopists with important information about the interactions that play a role in the system, including weakly polar ones such as NH → π interactions. The experimental information collected, which was interpreted with the help of theory, also enabled us to validate the energetic side of these calculations, providing a good example of synergy between experiment and theory. This encouraging result suggests that the study of larger model peptide systems, which is already on its way,23,50,51 should provide a fruitful approach for our understanding of the competition between local conformational preferences of each residue and the more global secondary structures in the peptide chain, like β-turns.

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References and Notes


