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Hydrophobic Collapse in N-Methylacetamide–Water Mixtures

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1 INTRODUCTION

The development of two-dimensional infrared (2D IR) spectroscopy45–47 led to extensive studies of biological and biochemical systems, with structural and dynamical properties of peptides and proteins11,12,16 being at the focus. Amide I (C=O stretch) vibrations were of particular interest due to their abundance in proteins and the resulting potential to access the information on structure and dynamics through their strong mutual coupling.5–12

The mechanism of protein folding13 is an important subject as misfolding of proteins leads to diseases like Alzheimer’s and Parkinson’s disease, type II diabetes, and cancer.14,15 Great progress in the understanding of this complex process has been made in recent years.11,12,16–37 The folding process can be steered by proteins called chaperones,38,39 in the framework model16 have been proposed as the most common mechanisms on a fundamental level for proteins is, however, challenging as sequence-specific effects inevitably play a role as well. The idea of this paper is to study the behavior of the amide groups forming the backbone of protein chains, when mixed with water. We want to eliminate the effect of side chains to address the question to what extent the interactions between water and amide groups drive structural formation.

N-Methylacetamide (NMA) is a convenient model molecule for studying such interactions. NMA is composed of a single peptide bond42–48 (–CONH–), which contains a single amide I mode. Due to this structure, NMA is used as a model compound for the linkage between peptide units.44,49 NMA can both donate and accept hydrogen bonds (HBs)50 which makes it miscible with a wide range of solvents.42–44,46,51,52 This has led to extensive experimental42–45,48,52 and theoretical47,48,53–55 studies of vibrational dynamics of the amide I mode of NMA56 in different solvents.

The studies of NMA diluted in polar and nonpolar solvents42–45,48,51,52,54,56–65 provided important information on the relaxation pathways of the amide I mode and its spectral dynamics. An extensive study of the influence of the solvent on the amide I mode vibrational dynamics was performed by DeCamp et al.48 The amide I lifetime appeared to be weakly dependent on the particular solvent and were reported to be 450 fs in heavy water,44 380 and 430 fs in heavy water and DMSO, respectively52 (with an ~2 ps tail reported in the latter study). In contrast, solvents that form weak HBs lead to slower HB and reorientational dynamics.48,52 Spectral dynamics of NMA molecules diluted in water exhibit bimodal dynamics: a time scale of 50 fs was assigned to water librations, while HB vibration dephasing occurred on a 180 fs time scale.48 Recently, NMA bulk dynamics were studied by optical Kerr effect spectroscopy60 and theoretically and experimentally by 2D IR spectroscopy.67 In the former study, structural relaxation was observed to occur at the time scales of 30 and 180 ps at 300

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K; both times accelerated by a factor of ~30 at 470 K. In the latter paper, an interesting interplay between HB and vibrational coupling was reported and interpreted from the viewpoint of the existence of highly ordered HB chains where the transition dipoles are largely oriented parallel to each other along the chain of NMA molecules.

So far, the NMA dynamics have been studied for both diluted and bulk environments. However, in biological systems neither situation is relevant, as proteins and peptides form HBs with the aqueous surroundings. This situation can be only be mimicked by concentrated NMA–water mixtures. We will study such solutions to understand how the interactions between water and amide groups affect the HB structure and dynamics of the amide groups.

In this paper, we examine the dynamics of mixtures of NMA molecules and water with 2D IR spectroscopy and molecular dynamic simulations combined with spectral simulations. In the 50/50 molar % mixture the amide I lifetime is 450 ± 100 fs which is in line with previous studies.43,44,67 We show that such a mixture leads to clustering of NMA and water molecules, which leads to “frozen” dynamics at the picosecond time scale. The methyl groups in the NMA clusters tend to stick together, suggesting a hydrophobic collapse. However, as evident from the anisotropy measurements, this does not fully prevent the movement of the vibrational excitations between amide I modes typical for bulk NMA, but its extent is limited to the NMA cluster size.

2. MATERIALS AND METHODS

2.1. Sample Preparation. We used NMA with deuterated peptide bonds and deuterated water (D2O) to avoid unwanted overlap of the C==O stretching mode at ~1650 cm⁻¹ with the HOH bending mode (~1645 cm⁻¹). N-Methylaceticamidine-δ1 (NMA-δ1) with isotopic enrichment of >99% was obtained from CDN Isotopes; deuterium oxide (D2O) with isotopic purity of >99.9% of the D atom was purchased from Sigma-Aldrich. Both chemicals were used without any further purification.

NMA-δ1 is a solid at the room temperature (~22 °C), so it was placed on a heating plate of 40 °C until it was completely melted. After that, liquid NMA-δ1 was mixed with D2O in precalculated volumes in order to obtain mixtures with variable mole fractions of NMA-δ1 (hereinafter the mole fraction of NMA in solution is denoted as XNMA, i.e., X = 1 and X = 0 correspond to pure NMA-δ1 and heavy water, respectively). Approximately 1.5 μL of the NMA-δ1/D2O solution was squeezed between two 1 mm thick CaF₂ windows, which were preheated up to T = 40 °C to ensure homogeneous distribution of the sample all over the area. In order to minimize the contact of the samples with air, the assembly was placed in a sealed cylindrical sample holder. The sample thickness (estimated from the absorbance) was ~3 μm assuming that the solution was spread uniformly between the 1 inch diameter CaF₂ windows.

To avoid any contact with air moisture, sample preparation was performed under nitrogen atmosphere. During the experiments, to avoid potential crystallization of NMA-δ1 at high concentration, the temperature of samples were uniformly maintained at temperature of 30.5 ± 1 °C by a thermostat controlled by a thermocouple.

2.2. IR Spectroscopy. An FTIR-spectrometer Vertex-70 purged with dry nitrogen was used for recording the FTIR absorption spectra in the spectral range of 1550–1750 cm⁻¹. The spectral resolution was ~2 cm⁻¹; 32 scans were averaged for every absorption spectrum. The maximal OD in the region of interest was ~0.6.

An interferometer platform used as a base for the collinear 2D IR setup is described elsewhere.70 In short, an IR pulse (~25 μJ of energy, 135 fs in duration, 140 cm⁻¹ fwhm spectral width) centered at 1640 cm⁻¹ was split into two pumps (90% of the total intensity) of equal intensity, probe and reference. The time difference between probe and reference pulses was set at 40 ps to avoid any interference. A wobbler in the probe beam path generated a −π, 0, +π, 0 phase sequence for subsequent pulses (1 kHz repetition rate) to remove sample scattering. Before the sample the polarization of the pump was rotated by a λ/2 waveplate by 45° with respect to the pump polarization. After the sample, either the parallel or the perpendicular polarization component of the probe beam was selected by a polarizer. The probe spectrum was detected by one of the 2 × 32 pixel MCT array rows (Infrared Associates) to provide the ω₁ dimension of the 2D spectra. Detection of the reference beam spectrum was carried out by the second array row of the MCT spectrometer. To increase the signal-to-noise ratio, the probe spectrum was normalized by the reference spectrum for each laser shot. At fixed waiting times T between the probe and the latest pump pulse the delay between two pump pulses, t₁, was scanned up to 2 ps with fast scanning approach.70 The normalized probe spectrum was averaged over the wobbler-generated phase sequence to extract the nonlinear response related to all three pulses (two pumps and the probe), zero-padded to 4 ps, and Fourier-transformed to obtain the ω₁ dimension with the phase correction applied.72 For the pump–probe measurements, one of the pump beams was used.

The interferometer and the sample and detection compartments of the setup were purged with dry air to prevent water peaks in the IR spectrum and to minimize the sample adsorption of the water vapor.

2.3. Theory. The molecular modeling simulations were performed with the GROMACS suite 4.6.1 using a combination of the OPLS-AA force field and the SPC/E water model to describe the NMA/water mixture, with variable NMA/water content. The production run of 1 ns at 27 °C was performed at constant volume and the coordinates of all atoms using 1 fs time steps. Snapshots were stored at each 10 fs for analysis and spectral modeling. The temperature was kept constant using the Berendsen thermostat, and a 1.1 nm cutoff was used for both Lennard-Jones and Coulomb interactions. The truncation of the Lennard-Jones potential was compensated by introducing analytical corrections to pressure and potential energy.7 The long-range Coulomb interactions were treated using particle mesh Ewald method, with a grid step of 0.16 nm and a convergence of 10⁻⁵. All bonds were constrained using the LINCS algorithm.

The spectral modeling was performed using the numerical integration of Schrödinger equation (NISE) method.95,80 The time dependent Schrödinger equation was solved numerically for the time-dependent amide I Hamiltonian:

\[
H(t) = \sum_i \omega_i(t) \{B_i^+ B_i - \frac{\Delta}{2} \sum_i B_i^+ B_i B_i + B_i \} + \sum_{i \neq j} J_{ij}(t) B_i^+ B_j^+ B_j - \sum_i \bar{H}_i(t)\bar{E}(t)(B_i^+ + B_i) \tag{1}
\]
where $B^i_{\gamma}$ and $B_i$ are the bosonic creation and annihilation operators, $\alpha_i(t)$ is the time dependent fundamental amide I frequency for the $i$th molecule, and $\vec{\mu}_i(t)$ is the corresponding transition dipole. The anharmonicity, $\Delta_\gamma$, was kept constant at 16 cm$^{-1}$. The amide I site frequencies, $\omega_i(t)$, were calculated using the Jansen electrostatic map, which was reported to work well with the OPLS-AA force field. The long-range couplings between the different amide I units were calculated using the transition charge coupling (TCC) model, where a charge, $q_{\alpha_i}$ and a transition charge, $dq_{\alpha_i}$, and a normal mode coordinate, $\hat{\nu}_{\alpha_i}$, are assigned to each atom of the amide I unit. The TCC model has the form

$$J_\gamma = \frac{1}{4\pi\varepsilon_0} \sum_{n,m} \frac{dq_{\alpha_n}dq_{\alpha_m}}{|\vec{r}_{nm}|} \left( \frac{3q_{\alpha_n}\langle \hat{\nu}_{\alpha_n}\rangle\langle \hat{\nu}_{\alpha_m}\rangle}{|\vec{r}_{nm}|^3} - \frac{dq_{\alpha_n}\hat{\nu}_{\alpha_n}\hat{\nu}_{\alpha_m}q_{\alpha_m}}{|\vec{r}_{nm}|^3} \right)$$

(2)

Here, the subscripts $n$ and $m$ are the number the atoms that belong to different amide I local modes at molecules $i$ and $j$, respectively. The distance vector between two atoms in the involved molecules is given by $\vec{r}_{nm}$. The values of the parameters used are given in ref 18.

3. RESULTS AND DISCUSSION

3.1. Linear Absorption. The experimental and theoretical linear absorption spectra of the NMA-d$_1$/D$_2$O mixtures studied are shown in Figure 1. The experimental and theoretical spectra demonstrate similar trends: with increasing NMA-d$_1$ concentration, the spectra are blue-shifted, become broader at intermediate concentrations, and narrow down in diluted NMA solutions. Both pure NMA spectra exhibit a high-frequency shoulder, which was earlier shown to have mixed HB–vibrational coupling origin.

The experimental spectra at $X = 0.5$–0.7 also show a noticeable high-frequency sub-band, while its appearance in the theoretical spectra is not as noticeable except for the $X = 0.3$ case. The discrepancy most probably originates from the fact that the applied force fields overestimate the NMA clustering, which leads to more heterogeneity of the solution. Therefore, in what follows we have chosen to compare the experimental spectra of the $X = 0.5$ mixture with $X = 0.3$ of the simulated mixture (Figure 2) as representing the most similar heterogeneous case.

![Figure 2. Linear FTIR spectrum of the NMA-d$_1$/D$_2$O solution (wine, $X = 0.5$) in comparison with the simulated coupled (blue) and uncoupled (magenta) spectra at $X = 0.3$.](image)

To disentangle the origin of the main NMA band and the high-frequency sub-band, the linear spectrum of the exact same system was calculated with the vibrational coupling between amide I modes set to zero (Figure 2). The spectra with zero coupling (hereafter denoted as uncoupled) are narrower and blue-shifted by $\sim$13 cm$^{-1}$ in comparison with the coupled spectra. This indicates a strong effect of vibrational coupling on the spectral line shape as was previously reported for bulk NMA. The high-frequency sub-band vanishes, which suggests that the sub-band originates from the vibrational coupling.

To verify this conclusion, all possible types of HB between NMA and water were extracted from the MD simulations. Geometric criteria were used to determine if HBs were present between the molecules. For water–water and water–NMA HBs a common criterion defined for water was used, while the criterion described in ref 67 was used for NMA–NMA HBs. In total, there are 21 types of HB species (see the Supporting Information, section 1), which can be further categorized into three main groups according to the number of accepted HBs. Group I includes the free NMA (f-NMA) and NMA donating one HB (1d-NMA). The species in group II either accept one HB (1a-NMA) or donate one and accept one HB (1a-1d-NMA). Group III is characterized by two accepted HBs (2a-NMA) or two accepted and one donated HBs (2a-1d-NMA). No difference with respect to the origin of the HB donor (water or NMA) is made in this group categorization. The fraction of each group is depicted in Figure 3; group II is by far the most populated one.

To estimate the effect of the intermolecular coupling to the spectral features, the linear absorption spectra for each group were calculated in the static approximation for both coupled and uncoupled cases (Figure 4). The central frequency of group I is the most blue-shifted because of similarity of the effect of
HB donation and no HB on the frequency of the C=O oscillator. In contrast, for group III the central frequency is the most red-shifted due to two accepted HBs. The dominant single HB accepting species (group II) absorbs at a frequency in the middle of the band. Of course, this is true only for the central frequencies: due to the broadness of the spectral distributions, there are double HB accepting complexes (group III) of which the frequency is blue-shifted as compared to free NMAs in group I and vice versa.

For each group the vibrational coupling leads to broadening of the spectra due to additional inhomogeneity in the coupling strengths (compare Figure 4a and Figure 4b), similar to the observation in bulk NMA.67 However, the red shift of the coupled spectra does not exceed 4−5 cm\(^{-1}\), which is substantially lower than the 13 cm\(^{-1}\) red shift seen in the full dynamical calculations (Figure 2). This is because when calculating the static spectra for the different groups, the coupling between species belonging to different groups is neglected. Therefore, the similarity of the static structures tells us that the coupling between molecules belonging to different groups determines the shift in the dynamics spectra, and therefore the molecules belonging to the same group are dispersed throughout the liquid. A comparison between full static spectra and linear absorption spectra simulated with the NISE approach (i.e., accounting for motional narrowing effects) shows that the HB dynamics lead to narrowing of both the coupled and uncoupled spectra, hereby proving the importance of the motional narrowing effects.

In principle, the uncoupled spectra in Figure 2 could be obtained experimentally by using isotopically diluted NMA where the carbonyl carbon or oxygen is replaced by \(^{13}\)C and/or \(^{18}\)O which leads to red-shifting of the amide I absorption by 20−60 cm\(^{-1}\).\(^{34}\) This is comparable to typical values of vibrational couplings of <20 cm\(^{-1}\),\(^{35}\) so the intermolecular coupling is not fully suppressed.

### 3.2. Pump−Probe

To extract the amide I mode lifetime, the frequency-dependent isotropic component of the pump−probe signal was extracted from the experimental spectra as

\[
I_{\text{iso}}(t) = \frac{I_{\|}(t) + 2I_{\perp}(t)}{3}
\]

where \(I_{\|}(t)\) and \(I_{\perp}(t)\) are the parallel and perpendicularly polarized components, respectively, of pump−probe signal with respect to the pump. The isotropic transient spectra for an NMA-\(d_{1}/D_{2}O\) mixture with \(X = 0.5\) recorded at several pump−probe delays are shown in Figure 5. The transient spectra consist of a positive bleaching component (at \(~1650\) cm\(^{-1}\)) and a negative induced absorption component (at \(~1615\) cm\(^{-1}\)). The transient spectrum at 5 ps is drastically different from those at shorter times. This is ascribed to thermal effects\(^{86}\) that begin to dominate the pump−probe signal at longer times.

To determine the time scales of the population relaxation and thermalization, we used the isotropic pump−probe transients at the extreme points of bleaching/stimulated emission (\(~1650\) cm\(^{-1}\)) and induced absorption (\(~1615\) cm\(^{-1}\)) frequencies, along with the transient near the compensation point at \(~1625\) cm\(^{-1}\) (Figure 6). The latter represents the thermalization signal that is the least affected by the population dynamics. Its setup time of 4.2 \(\pm\) 0.3 ps is apparently much longer than the population relaxation time, which hints that a four-level model\(^{87}\) (with the ground, excited, intermediate, and hot-ground states) should be used to describe the transients. The lifetime of amide I mode calculated from such a model amounted to \(~450 \pm 100\) fs. While the obtained lifetime of the amide I mode corroborates the previous results, it might be affected by the presence of HB configurations with different lifetimes and mixing up frequencies of excited-state absorption and bleaching/stimulated emission of these species. Nonetheless, the value of 450 fs is sufficiently reliable for estimating the time window provided for 2D IR experiments (\(~1.5\) ps).

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**Figure 3.** Fraction of each NMA-\(d_{1}/D_{2}O\) group. The number of accepted−donated HBs in the group is shown in the bar. Above the bar, representative corresponding structures are depicted.

**Figure 4.** Linear absorption spectra in a static approximation of each group for coupled (a) and uncoupled (b) cases. For comparison, the full dynamic spectra from Figure 2 are also shown by the circles.

**Figure 5.** Isotropic transient spectra of \(X = 0.5\) NMA-\(d_{1}/D_{2}O\) mixture at different delay times (as shown in the legend).
3.3. 2D IR Spectroscopy. The linear spectrum does not contain any information about the NMA-d$_{1}$/D$_2$O dynamics. Such information can be extracted from nonlinear 2D spectra (Figure 7). All spectra are strongly elongated along the diagonal, which suggests “frozen” dynamics. However, more careful inspection reveals the presence of two peaks with central frequencies at $\sim$1635 cm$^{-1}$ and $\sim$1660 cm$^{-1}$ (see the linear spectra in Figure 2 for comparison). Therefore, the diagonal elongation might result from a simple combination of the two peaks each of which is not necessarily elongated diagonally (providing there is no chemical exchange between the peaks; see the Supporting Information, section 2). Hence, we decided to perform the center line slope (CLS)$^{88,89}$ analysis in two separate regions which correspond to the two peaks in Figure 2.

In the low-frequency region (Figure 8a), the experimental CLS values at short times are about 0.55 which indicates a substantial contribution of the homogeneous broadening, in accordance with the theoretical calculations. Most probably, this broadening originates from fast bath modes, such as the water librations. In contrast, the CLS values at short times in the high-frequency region (Figure 8a) are close to unity, indicating that this region of the spectrum is primarily inhomogeneously broadened (see Figure 4 and the discussion around it). The small rise at short waiting times most probably originates from the pulse overlap region and/or nonresonant (instantaneous) background response.

The theoretical CLS values in the low-frequency region capture the experimental ones rather well (Figure 8a). However, the theory predicts more homogeneity in the high-frequency region (Figure 8b); this may in part be due to the overestimation of vibrational couplings$^{81,90}$ (which is supported by the fact that the CLS values are higher when the coupling is set to zero; see Figure 8) and overestimation of frequency fluctuations of the free species. The latter likely arises as the molecular polarizability in the point charge based force fields used here are accounted for by scaled point charges to match the average dipole moment in the liquid and configurations far from this average are described less accurately.$^{54,75}$ We also note that the coexistence of free and clustered NMA molecules might lead to a vibrational lifetime in the low-frequency region that differs from the lifetime in the high-frequency region (see discussion in section 3.2).

At the main peak position the CLS values for the uncoupled simulations are generally higher than for the simulations accounting for the coupling (Figure 8a). In the high-frequency

Figure 6. Representative pump–probe transients of $X = 0.5$ NMA-d$_{1}$/D$_2$O mixture at the maximum of bleaching/stimulated emission (black symbols), induced absorption (red symbols), and the compensation point (blue symbols). The fits obtained from the four-level thermalization model are shown by the solid lines; the respective relaxation times (the excited state lifetime $T_1$ and the thermalization time $T_{\text{int}}$) are given in the legend. The sign of the 1615 cm$^{-1}$ transient was inverted to allow easy comparison with the other transients.

Figure 7. Normalized 2D IR spectra of $X = 0.5$ NMA-d$_{1}$/D$_2$O mixture at several representative waiting times $T$. The experimental (the left-hand columns) and the simulated coupled and uncoupled (the middle and right-hand columns, respectively) spectra are shown for parallel and orthogonal polarization arrangement. Red and blue colors represent the bleaching/stimulated emission and induced absorption, respectively. The equidistant contours are drawn with 10% steps from the maximal amplitude. The thick black lines show the results of the CLS analysis in the following frequency regions: experiment $1613 < \omega_1 < 1667$ cm$^{-1}$, theory (coupled) $1617 < \omega_1 < 1660$ cm$^{-1}$, and theory (uncoupled) $1624 < \omega_1 < 1660$ cm$^{-1}$.
region the CLS values are much more similar (Figure 8b). This suggests that the couplings are more important for the NMA molecules involved in HB. One must of course keep in mind that when the coupling is included, the excitations are no longer localized on a specific HB type.

For all waiting times, the experimental and theoretical CLS values do not change substantially in both frequency regions, which suggests “frozen” dynamics of the amide I mode at the picosecond time scale. Such “frozen” dynamics have been reported earlier for water in reversed micelles and around hydrophobic groups. This effect was explained as breakage of the 3D HB network among the water molecules. As we concluded from the analysis of the static spectra (Figure 4), different types of HB species are dispersed in the mixture. From 2D spectroscopy, we inferred that the dispersed structures are static at least at the time scale of a few picoseconds. This situation is visualized in Figure 9 where a snapshot from MD simulations clearly shows the phase separation of the mixture onto NMA clusters (in the center) surrounded by the water molecules (at the periphery).

To quantify the clustering effect, we calculated the joint angular–radial distribution functions (JARDFs) of the NMA/water mixture and compared them with JARDFs of bulk NMA (Figure 10). The initial peak at ~0.48 nm corresponds to HBs between amide units in the first solvation shell. After this, the JARDF for NMA contains apparent recurrences, in both the radial and angular dependences, which are also presented in less pronounced form in JARDF of the mixture. The features of the JARDF revealing the liquid structure are highlighted in Figure 11 where two particular cuts of the JARDFs are depicted, one in the radial direction at a fixed angle and another one at an angle direction at a fixed distance. The main peak of the bulk NMA JARDF has a higher amplitude than its mixture counterpart, which reflects more ordering among HBs between amide units already in the first solvation shell. Furthermore, several recurrences follow that correspond to HBs between amide units in the second and third solvation shells. This is not surprising given the previous findings that NMA in bulk forms chain-like structures with well-defined orientation of the HBs between amide units. These features are less pronounced in the NMA–water mixture, lending additional support for the system heterogeneity. The radial cut shows a similar trend: a relatively narrow (~50°) distribution of C=O angles for bulk NMA and a much broader (~65°) distribution for the NMA–water mixture. These observations lead to the conclusion that the NMA–water mixture is not as well-organized as bulk NMA and that the formation of the HB chains of NMA molecules is largely destroyed by water molecules.

The nanoscopic phase separation in the mixture can also be interpreted from the perspective of water molecules (Figure 12). The radial distribution function (RDF) for O···O distances of the NMA–water mixture has a higher amplitude initial peak and stronger recurrences than the RDF of bulk water. This indicates that the water molecules in the mixture form extended clusters, with a better defined water structure than in the bulk. This is well in line with many reported studies of water at molecular interfaces. To further quantify the clustering of NMA molecules, the RDF between methyl groups (C···C distance) of the NMA molecules were calculated (Figure 13) and compared with that of bulk NMA. The probability of finding two NMA methyl groups at a distance of 0.38 nm distance is higher in the mixture compared to bulk NMA. This makes us conclude that in the

Figure 8. CLS analysis of the 2D spectra (parallel polarization) in the amide I mode region performed at the low (experiment, coupled, 1620 < \( \omega_1 < 1636 \) cm\(^{-1}\); uncoupled, 1633 < \( \omega_1 < 1648 \) cm\(^{-1}\)) and high (experiment, coupled, 1644 < \( \omega_1 < 1659 \) cm\(^{-1}\); uncoupled, 1656 < \( \omega_1 < 1670 \) cm\(^{-1}\)) frequencies. The region of the CLS analysis is shown in the inset as a shaded area. Theoretical values are computed for the coupled case (blue circles) and uncoupled (magenta diamonds).

Figure 9. Snapshot from the MD simulation of the NMA–water mixture. Oxygen atoms are shown in red, hydrogen atoms in white, carbon atoms in cyan, and nitrogen atoms in blue.
mixture, the NMA molecules tend to form clusters in which the hydrophobic methyl groups stick together. This, thus, resembles the hydrophobic collapse observed in the protein folding process, but in this case it is solely driven by the presence of the small methyl groups in the NMA molecule instead of the large hydrophobic side chains present in real proteins.

To verify the destabilizing effect of water on the structural order of NMA, we measured and calculated the anisotropy decay. Both the experimental and calculated coupled anisotropies decay much faster than the rotational correlation function (RCF) but with the same rate as the population transfer regardless the spectral region. This is in a sharp contrast with bulk NMA where the anisotropy decays faster than RCF but slower than the population transfer. The latter was interpreted as vibrational excitation moving through the transition dipole network ordered by HBs. Apparently, the current situation is very different: the vibrational excitation is still mobile (the anisotropy decays faster than RCF), but the orientations of the transient dipole moments are not mutually aligned so that the anisotropy is quickly scrambled. The vibrational excitation in the mixture is delocalized over 8.6 NMA molecules on average as compared to 42 in the bulk (as determined using the inverse participation ratio). In the mixture, the delocalization is thus strongly reduced as it has to be localized inside the NMA cluster. All these observations confirm our conclusion that water induces a hydrophobic collapse in NMA destabilizing the intermolecular HBs and lowering the structural order.

4. CONCLUSIONS

In this paper, we have studied NMA–water mixtures with linear and nonlinear IR spectroscopy. The lifetime of the amide I mode has been found to be similar to the previous reports of dilute solutions of NMA. We revealed that adding water to NMA leads to disruption of the HB chains of NMA molecules, which were recently reported to be present in bulk NMA. Instead, solvation results in the formation of water and NMA clusters where the amide I dynamics are “frozen”, at least at the picosecond time scale. Curiously, only the frequency and

Figure 10. Joint angular–radial distribution functions of (a) NMA–water mixture and (b) bulk NMA. Θ is the angle between the pairs of C=O bond vectors (cos(Θ) = 1 and cos(Θ) = -1 for the parallel and antiparallel bond vectors, respectively), and r = r_{C=O} is the distance between the centers of pairs of C=O bonds. The color map corresponds to the occurrence of formation of the pair of C=O bonds: the red color shows the high occurrence, while the blue is used for low occurrence. Both distributions are normalized to unity at large r.

Figure 11. (a) Radial distribution function of NMA mixture and bulk NMA at cos(Θ) = 1 and (b) angular distribution function of dissolved and bulk NMA at r = 0.48 nm.

Figure 12. Radial distribution functions of water oxygen for NMA–water mixture (red) and bulk water (black). Both distributions are normalized to unity at large r.

Figure 13. Methyl–methyl RDF of NMA–water mixture (red) in comparison with bulk NMA (black). Both distributions are normalized to unity at long distances r.
rotational dynamics are frozen while the anisotropy is not and does decay at the ~1 ps time scale. This is because the excitation is mobile and therefore samples a few amide I modes with similar frequencies but different orientations. The former leads to constant CLS values (after the initial drop), as the excitations tend to be shared between vibrations with similar frequency, while the latter results in the anisotropy decay. It would be interesting to investigate if the water dynamics are frozen, too (similar to the recently reported “mayonnaise effect”\(^{100}\)); these experiments are underway.

In our opinion, the results obtained here could form a basis for understanding the hydration structure and (vibrational) dynamics of more complex biological systems than the NMA−water system studied in the current work. Real proteins may, for example, have charged groups at the surface, a complication that could be avoided using simpler systems such as NMA as a testing ground. We clearly find that even without an actual backbone or side chains the amide unit is experiencing a kind of hydrophobic collapse when exposed to water. This leads to disruption of the long-range ordered HB network between NMA molecules observed in bulk NMA\(^ {67}\) and to clustering of both NMA and water molecules. The resulting spectral dynamics reflecting the HB dynamics, at least in the NMA phase, are “frozen” as compared to the dynamics in the bulk. This suggests that the hydrophobic collapse is an intrinsic property of the amino acid backbone units, which may of course be enhanced by hydrophobic side chains or suppressed by hydrophilic ones.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpca.8b00276.

All possible types of hydrogen bonding between NMA and water and their bundling to the groups; 2D IR spectra of experimental NMA-\(d_1/D_2\)O and theoretical NMA−water mixture for both coupled and uncoupled cases; CLS analysis of 2D spectra of the bleaching part (PDF)

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**Figure 14.** Anisotropy decay of the main (a) and high-frequency (b) absorption peaks for experimental and theoretical spectra. The anisotropy values were calculated from the integration of the 2D spectra over the region shown in the corresponding inset; for the uncoupled case, the spectral area around the main peak at ~1650 cm\(^{-1}\) (Figure 2) was taken. (c) Rotational correlation function (RCF) and population transfer function obtained from MD simulations. Note that the uncoupled case produces the anisotropy decay similar to the RCF as expected from the localized (immobile) vibrational excitation.

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**REFERENCES**


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**ASSOCIATED CONTENT**

- Supporting Information

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**ABBREVIATIONS**

NMA, N-Methylacetamide; HB, hydrogen bond; 2D IR, two-dimensional infrared; FTIR, Fourier transform infrared; fwhm, full width at half-maximum; CLS, center line slope; JARDF, joint angular−radial distribution function; RCF, rotational correlation function; RDF, radial distribution function


