Amides as models to study the hydration of proteins and peptides - spectroscopic and theoretical approach on hydration in various temperatures

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Abstract

Interactions with water are one of the key factors which determine protein stability and activity in aqueous solutions. However, the protein hydration is still insufficiently understood. *N*-methylacetamide (NMA) is regarded as a minimal part of the peptide backbone and the relative simplicity of its structure makes it a good model for studies on protein–water interactions.

In this paper, the influence of NMA and N,N-dimethylacetamide (DMA) on surrounding water molecules in a range of temperature (25–75 °C) is studied by means of the FTIR spectroscopy. The results of the difference HDO spectra method are compared with the results of theoretical DFT calculations of NMA and DMA aqueous complexes.

Both NMA and DMA can be regarded as "structure-makers", yet their hydration spheres are different. These molecules exhibit a mixed and mutually dependent types of hydration: hydrophilic and hydrophobic. In the case of a NMA molecule that has one methyl group less than DMA, the type of

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hydrophobic hydration is more important. The DMA hydration sphere is less stable: the interactions between water molecules around the methyl groups are strained. Moreover, the hydration of NMA is much more temperature dependant than in the case of DMA. The source of the differences may be hidden in the N–H···H₂O interaction.

The delicate nature of water interactions with the peptide block models may be cautiously translated into the much more complicated interactions of proteins with their hydration shells.

Keywords: N-methylacetamide, hydration, FT-IR spectroscopy, DFT calculations

1 1. Introduction

Proteins play key role in every biological process. Their functionality
in living organism is determined by their secondary and tertiary structure,
which is formed and preserved by relatively weak interactions called hydrogen
bonds. Hydrogen bonds in protein-water systems consist mostly of intraand intermolecular bonds between protein molecule and water in natural
environment.

Interactions of water and proteins is one of the most important topics when it comes to studying behavior of proteins in their native state. Multiple studies have taken on this topic using various methods [1–12], however, it is still poorly understood. The study of proteins hydration experimentally is problematic due to the high level of complexity of these molecules. Additionally, overall hydration of proteins involves two general types of interactions, namely, hydrophilic hydration and hydrophobic hydration of the non-polar fragments. Those contributions are hard to separate in such complex structures.

N-methylacetamide (NMA) can be regarded as a single peptide bond unit 17 that is used as a model for studying both intra- and intermolecular interac-18 tions that would occur within proteins backbone in their native states. It has 19 been shown that the NMA's hydration sphere is very similar to the hydration 20 sphere of lysozyme. The lysozyme-affected water spectrum mainly consists of 21 the spectral share of water affected by NMA, which indicates the hydration 22 of the protein backbone.[12] Recent studies have confirmed the resemblance 23 of NMA to peptides as it seemed to form chain like structures similar to 24 protein backbone [13]. Existence of NMA dimers and other oligomers have 25 also been proposed [14–17]. Particularly, it was pointed out that NMA dimer 26 might be even more convenient peptide bond model than NMA itself [15]. 27

NMA and its water clusters have been extensively studied both experi-28 mentally [13–15, 18–28] (mainly by means of vibrational spectroscopy) and 20 theoretically [13, 15, 18, 19, 25, 27–35]. It has been numerously investi-30 gated using C=O or N-D stretch as a probe of changes in its surrounding 31 [13, 14, 18, 19, 26, 28–30]. Changes of water OD bands in isotopically diluted 32 systems is also a popular probe for studying hydration of NMA [18, 34, 36]. 33 Using aforementioned approach it has been stated that at 25° C water in 34 NMAs first hydration shell forms stronger hydrogen bonds than bulk water 35 [36]. Recently, these observations have been supported by Salamatova et. al 36 [18], with the results of MD simulations of NMA-water systems. Theoretical 37 studies describing this molecule's hydration have shown that a water molecule 38 forms stronger hydrogen bonds with the C=O group than with the N-H or 39

⁴⁰ N–D [29, 37–39]. Also, it has been observed that C=O····H₂O hydrogen ⁴¹ bonds are stronger than between water molecules. However, computational ⁴² studies of deuterated systems carried out by Yadav *et al.* [29] revealed that ⁴³ bonds formed by the C=O group and water are weaker than water–water ⁴⁴ hydrogen bonds outside NMA first hydration shell.

In other works, in which the hydration of this molecule was investigated, 45 it has been pointed out that the dilution of the bulk NMA with water or the 46 increase of temperature leads to the disruption of hydrogen bonded aggre-47 gates present in the bulk NMA at low temperatures [33]. Its hydration leads 48 to the formation of NMA-water clusters in which at least some of NMA dy-49 namics seems to "freeze at a measurable time scale. This means that NMA 50 experiences a kind of hydrophobic collapse while interacting with water [18]. 51 The *ab initio* analysis of $1 \cdot \text{NMA} - 2 \cdot \text{H}_2\text{O}$ clusters and their deuterated deriva-52 tives has allowed to assign normal modes of the aqueous hydrogen bonded 53 NMA [34]. The one water molecule has been hydrogen bonded to C=O group 54 of NMA and another one to the N–H group of those molecule. 55

Yang and Qian [40] conducted MM studies on properties of the isolated 56 cis- and trans-NMA and NMA-water complexes. Calculation of the interac-57 tion energies and cooperative effect have revealed that complex of the trans-58 NMA with two water molecules attached to the carbonyl oxygen is energet-59 ically unfavorable. Buck and Karplus on the basis of MD in a vacuum [39] 60 have found that in the case of such a complex distance between the hydrogen 61 atom of the first water molecule and the carbonyl oxygen atom is 1.71 Å and 62 the second is 1.97 Å. Also, it has been shown that NMA with one hydrogen 63 bonded water molecule on either C=O or N-H prefers a second one interact-64

⁶⁵ ing with the other group [40]. In addition, it has been found that complex
⁶⁶ with 3 water molecules is stable.

Several studies focus on a *cis*-trans equilibrium caused by rotation bar-67 rier around C–N bond in formamide and acetamide derivatives [23, 41–45]. 68 It has been shown that in the case of NMA similarly to proteins *trans* form of 69 NMA is the dominant form in liquid phases. This form also dominates in the 70 gas phase. Theoretical studies on properties of isolated NMA molecule and 71 NMA in aqueous solution revealed also the existence of two other resonance 72 structures in such systems [38, 46–48]. Neutral form of NMA is more pro-73 nounced in gas phase while zwitterionic structure in water. As a result, the 74 elongation of C=O bond and the shortening of N-C bond passing from gas 75 to water are observed. This effect is caused by the stabilization of zwitteri-76 onic structure of NMA by hydrogen bonding with water molecules. Also, the 77 rotation of the amide methyl group as an effect of solvation NMA molecule 78 has been observed. 70

The Born–Oppenheimer molecular dynamics and MD/MM have been used to study NMA in the gas phase and in a box of water molecules [48]. The resultant frequencies of the amide bond vibrations confirmed other theoretical and experimental studies [49, 50].

In this work we have investigated the water structure around NMA in wide temperature range by means of the FTIR spectroscopy. Vibrational spectroscopy is an excellent method for investigation of solute hydration [51, 52]. Isotopic dilution technique of HDO in water allows us to obtain bands which are more convenient to analyze and highly sensitive to changes in a solution. HDO spectra are narrower and less complex than the OH bands

of H_2O . Moreover, the OD vibration is practically free from intramolecular 90 and intermolecular couplings between oscillators. Using difference spectra 91 method [53–55] it is possible to isolate the contribution of HDO affected by 92 the solute from the OD band. Solute-affected HDO spectra can then be 93 compared with HDO spectra of bulk water, thus determining the structural 94 and energetic state of hydrogen bonds of water in the nearest surround-95 ings of solute. In isotopically diluted solutions of NMA there is significant 96 amount of N-deuterated NMA. N–D bond vibrations happen to manifest in 97 the IR spectrum in the same region as HDO bands. Spectra of water af-98 fected by N,N-dimethylacetamide (DMA) have also been obtained to isolate 99 ND stretching vibration from NMA-affected HDO spectra and to compare 100 those compounds in terms of hydration. To confirm the interpretation of 101 spectral results theoretical calculations were performed. 102

¹⁰³ 2. Materials and Methods

104 2.1. Chemicals and Solutions

N-methylacetamide (99+%, Aldrich), N,N-dimethylacetamide (Acros Or-105 ganics, 99.5%) and deuterium oxide (Aldrich, 99.96%) were used to pre-106 pare solutions without purification. Water used was deionized. For each 107 solute (NMA, DMA) stock solution in deionized water of maximum molal-108 ity 1 mol \cdot kg⁻¹ was prepared. Less concentrated solutions were prepared by 109 dilution of weighed amounts of stock solutions using deionized water. Each 110 of the solutions was then divided into two parts in order to prepare sample 111 and reference solutions. Sample solution was prepared by adding D_2O to one 112 of the parts in amount of 4% with respect to H_2O (by weight). Reference 113

solution was prepared by adding the same molar amounts of H_2O to the sec-114 ond part. The amount of used deuterium oxide was confirmed to be enough 115 for reaction $H_2O + D_2O = 2HDO \ (K \approx 4)$ to give an almost quantitative 116 amount of HDO. All solutions were prepared by weight. The prepared so-117 lutions were degassed before density and spectral measurements. Solution 118 densities were measured using the Anton Paar DMA 5000 densitometer at 119 all studied temperatures (25, 35, 45, 55, 65, and 75 °C, with the tolerance of 120 0.001°C). 121

122 2.2. FTIR Measurements

FTIR spectra of prepared solutions were recorded on Thermo Electron 123 Co. Nicolet 8700 spectrometer with resolution of 4 cm⁻¹. Each spectrum 124 was an average of 128 independent scans. A liquid cell (model A145, Bruker 125 Optics) with CaF_2 windows separated PTFE spacers was used. The path 126 length was determined interferometrically and was 29.0 μ m. The temperature 127 of measurements was kept at 25, 35, 45, 55, 65, and 75 °C with a tolerance 128 of $\pm 0.1 \,^{\circ}$ C and monitored using electronic thermometer with thermocouples 129 placed in the sample. The temperature of 75 $^{\circ}$ C has been chosen as an upper 130 temperature limit being the highest possible temperature due to technical 131 capabilities of the experimental setup. In the case of DMA solutions, the 132 maximum temperature was limited to 55 °C due to decomposition of solution 133 components to gaseous products at higher temperatures. The spectrometer 134 was purged with dry nitrogen to eliminate the influence of air components 135 on recorded spectra, namely, CO₂ and water vapor. 136

137 2.3. Spectral data analysis

Procedures of recording and analyzing spectra were handled using the
commercial PC software: OMNIC (Thermo Electron Co.), GRAMS/32 (Galactic Industries Corporation, Salem, NH) and RAZOR (Spectrum Square Associates, Inc., Ithaca, NY).

To obtain the desired information the difference spectra method [56] was 142 applied. This method assumes that water in a solution can be divided into 143 two additive contributions: the "bulk" water—identical to pure water—and 144 "affected" water—under the influence of solute. Therefore information about 145 solute-affected water can be isolated from solutions IR spectrum. The de-146 tailed procedure of spectral data analysis toward extraction of the solute-147 affected water spectra has been described in section S1 of Supplementary 148 Data. 149

¹⁵⁰ 2.4. Interpretation of the solute-affected water spectra

To interpret given solute-affected spectra the Badger-Bauer rule is used 151 [57]. This empirical rule states that energy of hydrogen bonds is proportional 152 to the shift of the OD band position, thus it is a convenient method of com-153 paring energetic state of solute-affected water with bulk water and investigate 154 the changes of this characteristics with temperature [54, 55]. The shape of an 155 OD bands contain information about hydrogen bond energy distribution in 156 water. With the existence of empirical curves that link the oxygen-oxygen 157 intermolecular distance (R_{OO}) with the OD stretching wavenumber (ν_{OD}) 158 it is possible to transform the spectral band shapes to the oxygen-oxygen 159 distance distribution function $P(R_{OO})$ [56, 58]. 160

161 2.5. Theoretical calculations

All calculations involving optimization of NMA or DMA complexes with 162 water molecules were performed with the Gaussian 09 v.D1 software [59] 163 available at the Academic Computer Center in Gdansk (TASK). The analy-164 sis of resulting wavefunction files involving electron density difference calcu-165 lations and the reduced density gradient (RDG) method [60] was performed 166 with the Multiwfn software v.3.3.9. [61]. Structures of NMA or DMA com-167 plexes with water molecules were optimized similarly to the procedure pre-168 sented in our previous papers [16, 62]. Final structures were optimized at 160 the density functional theory (DFT) level with B3LYP hybrid exchange-170 correlation functional [63, 64] and 6-311++G(d,p) basis set [65] within the 171 conductor-like polarizable continuum model (CPCM) of water as a solvent 172 [66, 67]. All calculations were performed with the D3 version of Grimme's 173 empirical dispersion correction including Becke–Johnson damping [68]. Such 174 a selection gives a good consistency of theoretical and experimental results. 175 Differences in electron densities caused by NMA or DMA interactions 176 with water molecules were calculated and visualized according to section 177 4.5.5 of the Multiwfn manual. Shortly, electron densities were calculated 178 for the water complex with NMA or DMA and separately for their isolated 179 fragments: NMA or DMA, and water molecules, and then appropriately 180 subtracted from each other. 181

182 3. RESULTS AND DISCUSSION

183 3.1. FTIR investigation of water structure

3.1.1. Characteristic of solute-affected water spectra and "bulk" water spectra 184 Figure 1 presents the NMA-affected HDO spectra (Figure 1a), DMA-185 affected HDO spectra (Figure 1b), and the "bulk" water spectra (Figure 1c) 186 as a function of temperature. Band shapes of these spectra were transformed 187 into the oxygen–oxygen distance distribution function $P(R_{OO})$ of the water 188 molecules. The obtained distance probability distributions are shown in Figs. 189 2a and 2b for solute-affected water, $P^a(R_{OO})$, and in Figure 2c for "bulk" 190 water, $P^{b}(R_{OO})$. The band parameters for affected HDO bands, together 191 with the bulk HDO bands, for measured temperatures are presented in Table 192 1, along with intermolecular oxygen-oxygen distances, R_{OO} . 193

A comparison of the values of the mean oxygen–oxygen distances (R_{OO}^g) 194 for solute-affected water and for "bulk" water (Table 1) points out that water-195 water hydrogen bonds are shorter in the presence of these solutes. The 196 shift of the values of the gravity center of bands, ν_{OD}^{g} , (related to the mean 197 energy of water hydrogen bonds) towards lower values with respect to the 198 ones corresponding to pure water at a given temperature (Table 1) suggests 199 that water affected by NMA and DMA forms on average stronger H-bonds 200 than pure water in the whole temperature range. In addition, the hydrogen 201 bonds of water molecules around NMA are stronger and shorter than those in 202 water affected by DMA. The above results indicate that both amides enhance 203 the water structure in their nearest surrounding in the whole temperature 204 range, and can be classified as "structure-making" solutes. This statement is 205 supported by the analysis of radial distribution functions obtained by means 206

of MD simulations at 27°C [18]. Analyzing the effect of temperature variation 207 on the affected HDO spectra, it can be seen that solute-affected water spectra 208 at lower temperature are characterized by stronger hydrogen bonds than at 209 higher temperatures. A similar relationship is shown by the "bulk" water 210 spectrum. This is obvious because at higher temperatures the distances 211 between water molecules increase and hydrogen bonds weaken. Furthermore, 212 the number of moles of affected water molecules (N values from Table 1)213 around NMA decreases as the temperature increases, and as a result at the 214 highest temperature, the two water molecules are affected by NMA. Water 215 molecules in the hydration sphere of DMA are not susceptible to temperature 216 variations, which is reflected the constant value of number of affected water 217 molecules as a function of temperature. 218

219 3.1.2. Differences in oxygen-oxygen distance distributions between "affected 220 water" and "bulk" water

To provide greater insight into the difference in intermolecular distances 221 between affected water and "bulk" water the following procedure was used: 222 the distance distribution function for "bulk" water, $P^b(R_{OO})$, (Figure 2c) 223 was subtracted from the distribution function of water affected by solute, 224 $P^{a}(R_{OO})$, (Figures 2a and 2b) at a given temperature. The results of the 225 subtraction, $\Delta P(R_{OO})$, each analyzed temperature are shown in Figures 3a 226 and 3b for NMA and DMA, respectively. This operation allows to observe 227 subtle changes in the population of hydrogen bonds of water affected by 228 solute relative to "bulk" water at a given temperature. 229

The analysis of distance differences (Figure 3b) clearly shows that the hydration sphere of DMA contains two populations of hydrogen bonds of

water for all studied temperatures: the first one corresponds to the weak 232 hydrogen bonds ($R_{OO} \approx 2.9$ Å), while the second one corresponds to the 233 strong water-water hydrogen bonds ($R_{OO} \approx 2.73$ Å). The distances assigned 234 to the second population is close to the distance typical to the water in the ice 235 phase $(R_{OO} = 2.76 \text{ Å})$ [69]. The creation of the above-mentioned populations 236 takes place mainly at the expense of very weak hydrogen bonds ($R_{OO} \ge 3.0$ 237 Å) and to a lesser extent of hydrogen bonds with mean energy (the population 238 of water-water hydrogen bonds only slightly longer than and equal to the 239 most probable distance in bulk water, value $R_{OO} = 2.826$ Å at 25 °C, see 240 Table 1). A slightly larger population of strong water hydrogen bonds relative 241 to the weak ones causes that in the surrounding of DMA the water structure 242 is strengthened with respect to the "bulk" water. Furthermore, it can be 243 seen that the differences between DMA-affected water and "bulk" water are 244 practically the same at all temperatures. This means that DMA enhances 245 to the same extent the water structure in the whole temperature range, and 246 thus demonstrates the high stability of the DMA hydration shell. 247

A common feature of the differences in intermolecular distances distribu-248 tion for NMA, $\Delta P(R_{OO})$, (Fig. 3a) is the reduction of population of very 249 weak hydrogen bonds of water $(R_{OO} \ge 3.0 \text{ Å})$ and simultaneous distinct 250 increase the strong ones $(R_{OO} \approx 2.73 \text{ Å})$, in comparison to "bulk" water. 251 The first population is practically unchanged with increasing temperature. 252 The most significant changes relate to the population of water molecules with 253 mean energy and weak hydrogen bonds of water ($R_{OO} \approx 2.9$ Å). At the lowest 254 temperatures $(25 \, {}^{\circ}\text{C} \text{ and } 35 \, {}^{\circ}\text{C})$, it may be seen that population of such hy-255 drogen bonds of water around NMA decreases, relative to the "bulk" water. 256

²⁵⁷ However, starting from the temperature of 45 °C, the discussed population ²⁵⁸ of hydrogen bonds increases with the temperature increase.

259 3.1.3. Differences between $\Delta P(R_{OO})$ vs. temperature 25°C

Changes in populations of hydrogen bonds in the hydration sphere of 260 NMA, with respect to the temperature of 25 °C as a reference point, are 261 shown in Figure 4. As can be seen, the population of water molecules with 262 mean and weak energy of hydrogen bonds is increased in comparison to 263 $\Delta P(R_{OO})$ at 25 °C. This population seems to be responsible for the weakening 264 of the hydrogen bond network around NMA when the temperature increases, 265 relative to the course of $\Delta P(R_{OO})$ at 25 °C. The NMA hydration sphere at 266 35 °C is additionally characterized by a slightly larger population of strong 267 hydrogen bonds (ca. 2.65 Å) relative to the temperature of 25 °C. Probably, 268 such a population can be caused by the formation of NMA–NMA dimers 269 [14–16] at this temperature, which may break up at higher temperatures. 270 Such an interpretation is however speculative because the presence of dimers 271 in aqueous solutions is difficult to observe. 272

273 3.2. DFT calculations: sources of differences in NMA and DMA hydration

A series of DFT calculations was performed to try to explain the differences in strong and weak populations of hydrogen bonds recognized on the basis of NMA- ad DMA-affected water spectra. Undoubtedly, the difference between NMA and DMA hydration lies in the presence of additional methyl group. Thus, we created a closed ring of water molecules encircling those compounds in the closest proximity of the amine proton or methyl group in the case of NMA and DMA, respectively, by a gradual addition of water

molecules to optimized NMA and DMA structures (see Figure 5). The water 281 ring is a model of the full hydration shell, which could be too demanding 282 to calculate for the selected set of method and basis set. In each case, the 283 minimal number of water molecules needed to create a closed circle of water 284 molecules was equal to eight. Animations showing optimized structures of 285 these complexes (in two variants: with overlaid differences in electron density 286 or with visualized places of weak interactions determined by RDG method) 287 are included in Supplementary data. 288

289 3.2.1. Steric effect

For both NMA and DMA the ring of water molecules is skewed (Figure 290 5, top view). The skewness seems to be connected with the steric effect 291 that nitrogen-bounded methyl group exerts on water molecules approaching 292 the carbonyl oxygen atom. The starting dihedral angle between C-H and 293 C=O bonds in NMA or DMA isolated molecules is close to 0°. This result is 294 in accordance with ref. [37]. The situation changes when interactions with 295 water molecules occur. When water molecules approach the carbonyl bond of 296 NMA the C–H bond easily adopts the dihedral angle to ca. 30° to fit the water 297 ring encompassing the molecule (Figure 5, front view), hence the skewness 298 of the water ring plane. The rotation of the amide metyl group of NMA, 299 when going from gas phase to solution, has also been found by Mennuci *et.al* 300 [37]. However, the examined clusters have been smaller (they contained up 301 to 3 water molecules) and the value of the dihedral angle was different. The 302 weak steric interaction between carbonyl group and methyl proton (marked 303 with red arrow in Figure 6a) also shifts to one side and makes place for a 304 water molecule to take the optimal location. The position of methyl proton 305

is not as fixed as in the case of DMA, where an additional steric interaction is placed between both amine methyl groups (large green/red patch under carbonyl group in Figure 6a). Moreover, carbonyl group (C=O) and water molecules interacting with it lies in one plane and the water ring is highly symmetrical (Figure 5, front view).

Interactions between water molecules or between water molecules and the 311 carbonyl group of NMA or DMA are enhanced with respect to the "bulk" 312 structure ($R_{OO,bulk,av} = 2.817$ Å, see Fig. S5 in Supplementary Data), thus, 313 an enhanced population is expected to be present in the R_{OO} distance dis-314 tribution. In fact, both compounds exhibit such an enhancement (Figure 3). 315 The population origins not only in water–carbonyl interactions but can be 316 the effect of the hydrophobic hydration, in which van der Waals interactions 317 have a dominant role (green/olive patches around methyl groups in Fig. 6a). 318 In the case of formamide, i.e. a molecule that does not contain hydrophobic 319 groups in its structure, we observe only an increasing population of weak 320 hydrogen bonds of water, in comparison to the "bulk" water (see Fig. 8a 321 in ref. [36]) This indicates that in the presence of formamide, the hydrogen 322 bonds between water molecules and the oxygen atom of carbonyl group and 323 those between water molecules are weaker. 324

The interaction of carbonyl group of NMA with a single water molecule does not seem to be distinguished, and its energetics is similar to the one between water molecules [16, 29]. However, the closing of the water ring around NMA, involving weak hydrophobic interactions (green/olive patches in Fig. 6a), enhances significantly $C=O\cdots H_2O$ binding. In such a situation, two water molecules interact strongly with carbonyl oxygen and almost to the same degree ($R_{OO} = 2.710$ Å and 2.728 Å). Similar length (2.73 Å) beetween carbonyl oxygen and water oxygen has been obtained by means of MD simulations for a larger system in ref. [38]. The shape of the electron density differences caused by the interaction of NMA and the water ring confirms high similarity, or symmetry, in both water–carbonyl interactions.

Further analysis shows that under the influence of water molecules the 336 electron density increases on the carbonyl oxygen. This is accompanied with 337 the decrease of electron density along the C=O bond. Also, the decrease 338 of electron on the amide hydrogen and its increase along the N-H bond 339 towards the nitrogen atom can be observed. In addition, electrons of N–C 340 bond are pushed towards the carbon atom. This results are in agreement 341 with the analysis of Wannier Centers average positions for NMA molecule in 342 gas phase and in water given in ref. [38] 343

A different picture emerges from the analysis of an analogous DMA data. 344 Such a water ring closure in the DMA system is less favorable. The van der 345 Waals interactions are less organized and differences in $C=O\cdots H_2O$ inter-346 actions are larger than in the case of NMA. The introduction of additional 347 methyl group simply inhibits the rotation of other two methyl groups in 348 DMA molecule. The nitrogen-bonded methyl group of DMA cannot adopt 349 an optimal angle to fit the water ring as in NMA system. This way, only one 350 water molecule of water can easily interact with the carbonyl group while the 351 second one has to find another—non-optimal with respect to NMA—place 352 at the expense of the interaction energy with the carbonyl group. One water 353 molecule is closer to the carbonyl oxygen atom ($R_{OO} = 2.705$ Å), while the 354 second one interacts poorly ($R_{OO} = 2.886$ Å, higher even than the average 355

"bulk" distance). Such an inequality may contribute to the presence of two 356 distinct water hydrogen bond populations (stronger and weaker relative to 357 the "bulk" water) in the vicinity of DMA molecule, as in experimental results 358 presented in Figure 3. It should be stressed that the population of strong 359 hydrogen bonds includes also interactions between water molecules around 360 methyl groups. The symmetry of water-carbonyl bond-water double inter-361 action is broken. Such a division is also clearly visible in the electron density 362 difference (Figure 5, top view), where the weakly interacting water molecule 363 is pushed away from the plane of the water ring. The change in electron den-364 sity for the first water molecule is highly similar to the case of NMA, while 365 the second one is non-symmetrical and exhibits a weaker electron density 366 transfer. 367

According to the above discussion, the differences in the hydration prop-368 erties of both molecules are related to the formation of a water cage around 369 them. Both NMA and DMA due to the presence of hydrophilic and hy-370 drophobic groups in their molecule are characterized by two types of hy-371 dration, which are mutually dependent. Moreover, hydrogen bonds between 372 water molecules involved in the cage formation and the interaction of water 373 molecules with the carbonyl oxygen atom are cooperative. The formation 374 of a water cage around hydrophobic groups affects the equality/inequality 375 of interactions of water molecules with the carbonyl group. In the NMA 376 molecule, the hydrophobic type of hydration is more important, although 377 it has one methyl group less than DMA. This is due to a worse fit of the 378 "ice-like" water around methyl groups of DMA to the molecular shape of the 379 molecule. 380

381 3.2.2. Importance of $N-H \cdots H_2O$ interaction

The interaction of water molecule with hydrogen atom of amino group of 382 NMA ensures the stability of the hydration sphere around NMA. It acts like 383 a "clip" stabilizing the hydration cage around NMA. At lower temperatures, 384 this interaction is incorporated in the hydration cage [36]. When the tem-385 perature increases, the hydrogen bonds around NMA weaken and eventually 386 break, and as a result, the water molecule interacting with the hydrogen 387 atom of amino group starts to lose its contact with other water molecules. 388 The relatively weak $N-H\cdots$ water interaction (the distance between the ni-389 trogen atom of NMA and oxygen atom of water is 2.890 Å), freed from other 390 interactions, contributes to the increase of the population of weak hydrogen 391 bonds in the presence of NMA at higher temperatures (see Fig.3a). 392

The hydration shell around DMA, represented with the modeled ring of eight water molecules, is already less stable than in the case of NMA at ambient temperatures (see Figure 6a, the van der Waals interactions are scattered much more than in the case of NMA). The ring of water molecules around DMA is stretched from the very beginning due to the relatively large methyl group in the place of amine proton. In consequence, the already weakened hydration shell may be less prone to the temperature increase.

400 4. Conclusions

The effect of NMA and DMA on the water structure as a function of temperature has been studied by means of FTIR spectroscopy, supported with DFT calculations. The results of theoretical calculations helped to resolve spectral results. The agreement between experimental and computational results was satisfactory and gave a consistent picture of the amides hydration
in aqueous solution.

The results revealed that hydrogen bonds of water molecules around these 407 solutes are stronger and shorter than those in pure water in the whole tem-408 perature range. Although both solutes strengthened the water structure, 409 significant differences in the characteristics of their hydration shells could be 410 noticed. The essential difference in the hydration of NMA and DMA is the 411 way they create the water cage around them, which is the result of both 412 hydrophilic and hydrophobic hydration. These two types of hydration are 413 mutually dependent. Thus, the increase in the population of strong hydrogen 414 bonds of water in the surroundings of these solutes is the effect of the inter-415 actions of water molecules with the oxygen atom of carbonyl group and those 416 between water molecules around hydrophobic groups. The hydration sphere 417 of formamide, a molecule that does not contain any hydrophobic group, is 418 characterized by an increased population of weak hydrogen bonds of water, 410 with respect to the "bulk" water ref. [36], without the strengthening ob-420 served in the case of NMA or DMA. This suggest that the presence of a 421 hydrophobic element in the molecule is a factor conditioning the enhanced 422 network of hydrogen bonds around the solute. 423

The DMA molecule is additionally surrounded by a population of water molecules that forms weak hydrogen bonds with respect to the "bulk" water (at all given temperatures). The presence of this population results from the poor fit of the water cage to the solute's geometry. Such a disadvantageous fit turn out to be less susceptible to the temperature change because the hydration cage is already distorted at lower temperatures, in comparison to 430 NMA.

On the other hand, the NMA water cage is better organized than in the 431 case of DMA. The hydrophobic type of hydration is more important in the 432 case of NMA molecule, although it has one methyl group less than DMA. 433 The interaction of water molecule with the hydrogen atom of amino group is 434 significant for the stability of the hydration sphere of NMA. Thus, at lower 435 temperatures the hydration sphere of NMA contains only the population of 436 stronger hydrogen bonds, in comparison to the "bulk" water. As the temper-437 ature increases, the additional population of water molecules, characterized 438 with weak hydrogen bonds, is observed in the water population affected by 439 NMA. It results from the weak $N-H\cdots$ water interaction. At lower tempera-440 tures, this interaction is incorporated into the hydration cage, while at higher 441 temperatures it starts to be exposed when water molecule interacting with 442 the hydrogen atom of amino group loses its contact with the cage around the 443 molecule. 444

Although both NMA and DMA are simple molecules, their similarity 445 to the basic building block of proteins makes them convenient models of 446 protein–water interactions. The obtained results clearly show that the con-447 tributions to the hydration sphere of the solute are not additive due to the 448 number of non-polar groups. A stable hydration sphere depends on the pos-449 sibility of building a cooperatively reinforced network of hydrogen bonding 450 water molecules around the solutes. The changes in the hydration of solutes 451 caused by the temperature increase and their explanation may be cautiously 452 translated into phenomena accompanying the process of protein denatura-453 tion. 454

455 5. Acknowledgements

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6. Declaration of interest

⁴⁶⁰ The authors declare that they have no conflict of interest.

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Table 1: The parameters of HDO bands of water affected by NMA (Figure 1a), water affected by DMA (Figure 1b), "bulk" water (Figure 1c), and the respective intermolecular oxygen–oxygen distances. R_{OO} errors have been estimated on the basis of the HDO bands position errors.

T^{a}	$N^{\ b}$	$\nu^o{}_{OD}$ c	${\nu^g}_{OD}$ d	$fwhh \ ^{e}$	I^{f}	$R^o{}_{OO}{}^g$	$R^{g}{}_{OO}{}^{h}$
NMA-affected water spectrum							
25	$3.0{\pm}0.5$	2505 ± 2	2486 ± 2	177 ± 4	13120	$2.821 {\pm} 0.003$	$2.821 {\pm} 0.003$
35	2.8 ± 0.5	2517 ± 2	2492 ± 2	178 ± 4	11836	$2.821 {\pm} 0.003$	$2.828 {\pm} 0.003$
45	$2.6{\pm}0.5$	2518 ± 2	2495 ± 2	167 ± 4	10640	$2.836 {\pm} 0.003$	$2.833 {\pm} 0.003$
55	2.5 ± 0.5	2526 ± 2	2501 ± 2	175 ± 4	10059	$2.836 {\pm} 0.003$	$2.838 {\pm} 0.003$
65	$2.4{\pm}0.5$	2526 ± 2	2505 ± 2	175 ± 4	9400	$2.841 {\pm} 0.003$	$2.844{\pm}0.003$
75	2.2 ± 0.5	2534 ± 2	2511 ± 2	169 ± 4	8447	$2.851 {\pm} 0.003$	$2.849 {\pm} 0.003$
DMA-affected water spectrum							
25	2.5 ± 0.5	2517 ± 2	2497 ± 2	165 ± 4	16169	$2.836 {\pm} 0.003$	$2.836 {\pm} 0.003$
35	2.5 ± 0.5	2524 ± 2	2503 ± 2	167 ± 4	14666	$2.836 {\pm} 0.003$	$2.836 {\pm} 0.003$
45	$2.6{\pm}0.5$	2528 ± 2	2505 ± 2	167 ± 4	13467	$2.841 {\pm} 0.003$	$2.841 {\pm} 0.003$
55	2.5 ± 0.5	2534 ± 2	2511 ± 2	166 ± 4	12066	$2.851 {\pm} 0.003$	$2.849 {\pm} 0.003$
"bulk" water spectrum							
25	-	2509 ± 2	2505 ± 2	162 ± 4	10434	$2.826 {\pm} 0.003$	$2.844{\pm}0.003$
35	-	2513 ± 2	2509 ± 2	164 ± 4	10022	$2.836 {\pm} 0.003$	$2.849 {\pm} 0.003$
45	-	2519 ± 2	2513 ± 2	164 ± 4	9670	$2.836 {\pm} 0.003$	$2.854{\pm}0.003$
55	-	2522 ± 2	2517 ± 2	167 ± 4	9281	$2.838 {\pm} 0.003$	$2.859 {\pm} 0.003$
65	-	2528 ± 2	2521 ± 2	170 ± 4	8965	$2.844{\pm}0.003$	$2.861 {\pm} 0.003$
75	-	2532 ± 2	2524 ± 2	169 ± 4	8612	$2.849 {\pm} 0.003$	$2.867 {\pm} 0.003$

^{*a*} Temperature (°C). ^{*b*} Affected number, equal to the number of moles of water affected by one mole of solute. ^{*c*} Band position at maximum (cm⁻¹). ^{*d*} Band position at gravity center (cm⁻¹). ^{*e*} Full width at half-height (cm⁻¹). ^{*f*} Integrated intensity (dm³ · mol⁻¹ · cm⁻¹). ^{*g*} The most probable O···O distance (Å). ^{*h*} Mean O···O distance (Å).



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Figure 1: Solute-affected spectra in the OD stretching region for (a) NMA, (b) DMA, and (c) "bulk" HDO spectra as a function of temperature.



Figure 2: Interatomic oxygen-oxygen distance distributions function derived from the HDO spectra affected by (a) NMA, (b) DMA, and (c) "bulk" water spectra (Figure 1) for all studied temperatures. The vertical dashed line corresponds to the value of the most probable oxygen-oxygen distance in bulk water at 25 °C (2.826 Å, see Table 1).



Figure 3: Differences between interatomic oxygen–oxygen distance distribution function of solute-affected water, $P^a(R_{OO})$, and the "bulk" water, $P^b(R_{OO})$, (Figure 2) at given temperature for (a) NMA and (b) DMA. The vertical dashed line corresponds to the value of the most probable oxygen-oxygen distance in bulk water at 25 °C (2.826 Å, see Table 1).



Figure 4: The result of subtraction of the oxygen-oxygen distance function at 25 °C, $\Delta P(R_{OO})_{25 \ ^{\circ}C}$, (blue line in Figure 3a) from the distance function at different temperatures, $\Delta P(R_{OO})_T$, obtained for NMA. The vertical dashed line shown to the position of the most probable oxygen-oxygen distance in bulk water at 25 °C (2.826 Å, see Table 1).







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Figure 5: Differences in electron density of NMA and DMA complexes with water molecules caused by interactions of a central molecule with the water ring. Red regions correspond to increase of the electron density, blue regions mark its decrease. Additional movie files with the same structures are available in Supplementary Data.



Figure 6: (a) Visualization of weak interactions in NMA and DMA complexes with water molecules. Blue disks correspond to hydrogen bonds, green/olive patches correspond to weak van der Waals interactions, red (marked with red arrows) correspond to steric repulsion. Selected water molecules are made transparent for clarity. Hydrogen bonds are marked with dashed black lines. Numbers indicate $O \cdots O$ pairs sorted according to their length (as in Figure b). Additional movie files with the same structures are available in Supplementary Data. (b) Distribution of oxygen \cdots oxygen distances in NMA (red) and DMA (white) complexes with water. Only distances shorter than 3 Å are shown, and pairs corresponding to interaction of water molecules with the carbonyl oxygen are marked with arrows. The $O \cdots O$ pairs in red, green, or blue frames correspond to strengthened, "bulk-like", and weakened hydrogen bonds, respectively.