Molecular Dynamics Simulation of Polyacrylamide Adsorption on Cellulose Nanocrystals

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Abstract: Classical molecular dynamics simulations of polyacrylamide (PAM) adsorption on cellulose nanocrystals (CNC) in a vacuum and a water environment are carried out to interpret the mechanism of the polymer interactions with CNC. The structural behavior of PAM is studied in terms of the radius of gyration, atom–atom radial distribution functions, and number of hydrogen bonds. The structural and dynamical characteristics of the polymer adsorption are investigated. It is established that in water the polymer macromolecules are mainly adsorbed in the form of a coil onto the CNC facets. It is found out that water and PAM sorption on CNC is a competitive process, and water weakens the interaction between the polymer and CNC.

Keywords: cellulose nanocrystals; polyacrylamide; adsorption; molecular dynamics

1. Introduction

Polymer nanocomposites are materials that consist of polymer matrices and nanofillers distributed in them. The most important factor for achieving the enhancement of the bulk physical (transport, thermal, mechanical) properties of nanocomposites is interfacial interactions between the polymer macromolecules and the nanofiller particles. A large number of nanofillers with a variety of polymer matrices have been used to achieve the enhancement of useful properties. The simplified model of a nanofiller reinforced polymer suggests the wrapping of the polymer macromolecules around the nanofiller, providing an intuitive understanding of the mechanism responsible for the improved physical properties. However, large-scale disorder in polymer nanocomposites leads to substantial reduction of physical properties compared to predictions based on idealized filler morphology [1,2]. Computer simulations can address the issues related to the difficulty of characterizing the in situ structure of nanoscale objects [3,4]. Achievements in modeling polymer matrix nanocomposites based on identification of some challenges can enhance our understanding of physicochemical properties of polymer nanocomposites [5]. Specifically, in-depth understanding of structure and dynamics at solid/polymer interfaces at nanometer level plays a key role in designing materials with adjustable properties [6–8].

The features of cellulose nanocrystals (CNC) (a high crystallinity degree, anisotropic shape, high aspect ratio, and large surface area) have important consequences for the interfacial behavior and have attracted great attention of the materials community because of their sustainability, renewability, and biodegradability. Therefore, CNC have widely been used as reinforcing agents in polymeric nanocomposites in recent years [9].

Nowadays, atomistic modeling of cellulose nanocrystals has been used to complement experimental measurements. Computer simulations help to predict self-assembly as well as mechanical, energetic, thermal, and structural features of cellulotic nanomaterials and provide a fundamental understanding of the atomic-scale origins of these characteristics [10–16]. Models have been employed
to predict some CNC properties including the most frequently reported mechanical ones [17,18]. Molecular modeling has been utilized to investigate the properties of amorphous cellulose as opposed to CNC [19–21]. Some studies have applied molecular modeling to investigate the interaction of CNC with a liquid solvent (most solvent studies are focused on water) [22–24] and with polymeric materials [25].

The current work is a continuation of our previous study that was devoted to the investigation of the microscopic mechanism of polyvinylpyrrolidone (PVP) adsorption on a cellulose nanocrystal and the role of water in this process [26]. We have revealed earlier [27] that PVP adsorption onto CNC can block lateral interactions between the CNC and prevents their agglomeration in the lateral direction, i.e., hinders growth of the CNC particles width upon the concentration increase or drying of the composites. Moreover, freezing of CNC suspensions can align rod-like CNC particles in direction of ice crystal growth allowing formation of the CNC aggregates with a high aspect ratio. These aggregates can be broken down easily in water and some organic solvents, providing good dispersibility of the composites.

In this work we focus our attention on the intermolecular interactions in the systems containing polyacrylamide (PAM), CNC and a polar solvent (water). PAM is a water-soluble linear polymer. PAM and its derivatives have different applications in many areas including agricultural, water treatment, medical, and petroleum industries [28,29]. PAM differs from PVP because it contains not only atoms playing the role of proton-acceptors, but also atoms acting as proton-donors. Because of such structure the behavior of PAM in water, as well as its interactions with the solvent and CNC have some features that we describe in terms of structural and dynamic properties by the all atomic molecular dynamics method.

2. Computational Details

Classical molecular dynamics simulations were carried out using a GPU-accelerated (graphics processing unit) Gromacs-5.0.7 software package [30]. Molecular graphics and visualization were performed using VMD 1.8.6 [31]. The molecular dynamics (MD) simulations were carried out for the NVT ensemble (a constant number of particles N, a volume V, and a temperature T). The reference temperature of 298 K was kept constant using a Nose–Hoover thermostat [32,33] with the coupling constant $\tau = 0.1$ ps. Periodic boundary conditions were applied to all three directions of the simulated cubic box. The Verlet algorithm [34] was adopted to integrate the equations of motion. The modified Ewald summation method [35,36] was used to account for the corrections of the long-range electrostatic interactions with a cutoff radius of 1.5 nm, which was also the cutoff value for the Van der Waals (VDW) interactions. All the bond length constraints were implemented using the LINCS algorithm (LInear Constraint Solver) [37]. In our work for PAM (Figure 1a) we utilized a potential based on OPLSAA (Optimized Potential for Liquid Simulations All Atomic) force field parameters [38]. The initial structure of a PAM macromolecule containing 50 monomer units (with the molecular weight of 3552 g/mol) was constructed by means of Avogadro [39]. This number of monomer units is sufficient to observe the conformational transitions of the polymer and to obtain sufficient statistical data to calculate quantitative characteristics. For cellulose we used GROMOS54a7 force field parameters [40]. The initial structure of CNC was built based on the experimental crystallographic data [41] by a toolkit named Cellulose Builder [42]. The model of the I$_B$ CNC consisted of 14 or 9 glucan chains, and the degree of polymerization of each chain was 10 (Figure 1b,c). The number of chains and degree of polymerization provide a sufficient surface area for effective interactions with the polymer. As known, in the structure of cellulose there is clear segregation into polar (OH) and nonpolar (CH) sites [10]. Because of the hydrophobic properties of the glucopyranose plane, the sheet-like structure of the top and bottom surfaces (200) of CNC (Figure 1c–e) has a predominantly hydrophobic character. CNC with predominantly hydrophilic surfaces (110 and 1–10) (Figure 1h, for example) has a large number of free OH groups (O1H1, O6H4, O3H3, O2H2). In the current study, we consider two types of CNC with large hydrophobic facet (200) and with large hydrophilic facets (110 and 1–10). The systems were solvated by water which was preliminarily equilibrated in $NpT$-ensemble at 298 K and 0.1 MPa.
For water molecules, the SPC/E (extended simple point charge) model \cite{43} was used. After systems energy minimization, we equilibrated the systems for 0.5 ns in the NVT ensemble. The production run simulations were performed in NVT ensemble in a cubic box with periodic boundary conditions for 10 ns for all systems in a vacuum and binary system PAM-water, 15 ns for large systems with a high concentration of PAM. For the systems with a low PAM concentration, the simulation time was chosen as 60 ns to obtain sufficient statistical data. A time step was set of 1 fs. The data for the analysis were collected every 0.1 ps. In Supplementary Material one can find the time dependencies of temperature for all systems, which are evidence that the structures obtained are equilibrated well at all length- and time-scales (Figure S1).

**Figure 1.** Atom numbering in monomer units of polyacrylamide (PAM) (a) and a cellulose nanocrystal (CNC) (b), CNC with a predominantly hydrophobic surface (c—top view; d,e—end-on views) and with a predominantly hydrophilic surface (f—top view; g,h—end-on views). The red spheres denote the oxygen atoms, the gray ones represent the hydrogen atoms, the cyan ones show the carbon atoms, and the blue sphere represents the nitrogen atom. The instant snapshot of System 3 (i): CNC is blue and PAM macromolecules are green, the red dots are the water molecules.

| System | Medium and CNC type | N(PAM) | N(CNC) | N(Water) | t, ns | L, nm |
|--------|---------------------|--------|--------|----------|------|------|
| System 1 | vacuum | 1 | - | - | 10 | 6.0 |
| System 2 | water | 1 | - | 6916 | 10 | 6.0 |
| System 3 | water hydrophobic CNC | 3 | 1 | 31322 | 60 | 9.0 |
| System 4 | vacuum hydrophobic CNC | 3 | 1 | - | 10 | 9.0 |
| System 5 | water hydrophilic CNC | 3 | 1 | 31322 | 60 | 9.0 |
| System 6 | vacuum hydrophilic CNC | 3 | 1 | - | 10 | 9.0 |
Ten systems were simulated; the simulation details are listed in Table 1. The initial configuration of Systems 2 was constructed placing 1 PAM molecule with a nearly extended conformation into a cubic box with 6916 water molecules. Initial Systems 3, 5, 7, and 9 were built by inserting one CNC with predominantly hydrophobic and hydrophilic surfaces, respectively, into the center of the cubic boxes. Then 3 or 64 PAM macromolecules were placed into the cell with CNC, which was followed by the addition of 31322 (System 3 and System 5) or 273740 (System 7 and System 9) water molecules. The size of the water box was chosen so that it ensured the systems had at least a 20 Å solvation shell in all the directions. As an example the instant snapshot of System 3 is presented in Figure 1i. The initial configurations of Systems 4, 6, 8, and 10 were obtained by removing the water molecules from the final configuration of Systems 3, 5, 7, 9, respectively.

| System   | Medium and CNC Type                  | N (PAM) | N (CNC) | N (Water) | t, ns | L, nm |
|----------|-------------------------------------|---------|---------|-----------|------|------|
| System 1 | vacuum                              | 1       | -       | -         | 10   | 6.0  |
| System 2 | water                               | 1       | -       | 6916      | 10   | 6.0  |
| System 3 | water hydrophobic CNC               | 3       | 1       | 31322     | 60   | 9.0  |
| System 4 | vacuum hydrophobic CNC              | 3       | 1       | -         | 10   | 9.0  |
| System 5 | water hydrophilic CNC               | 3       | 1       | 31322     | 60   | 9.0  |
| System 6 | vacuum hydrophilic CNC              | 3       | 1       | -         | 10   | 9.0  |
| System 7 | water hydrophobic CNC               | 64      | 1       | 273740    | 15   | 20.5 |
| System 8 | vacuum hydrophobic CNC              | 64      | 1       | -         | 10   | 20.5 |
| System 9 | water hydrophilic CNC               | 64      | 1       | 273740    | 15   | 20.5 |
| System 10| vacuum hydrophilic CNC              | 64      | 1       | -         | 10   | 20.5 |

3. Results and Discussion

In order to understand the structural behavior of the polymer chain placed into the water box, we calculated the radius of gyration ($R_g$), which is one of the most important quantities in conformational statistics of polymer chains [44,45]. The radius of gyration was calculated by the following equation:

$$ R_g = \left( \frac{\sum_i \|r_i\|^2 m_i}{\sum_i m_i} \right)^{1/2}, $$

where $m_i$ is the mass of site $i$ and $r_i$ is the position of site $i$ relative to the center of mass of the molecule. As can be seen in Figure 2, $R_g$ differs from the original value in both the vacuum (System 1) and water (System 2). In water, the value of the radius of gyration fluctuates during the simulation process and achieves $\approx 1.5$ nm, while at the beginning of the simulation $R_g$ was 1.7 nm. Therefore, we can conclude that we observe a certain transformation of the polymer chain, which, however, does not lead to a strong folding of the PAM and the chain remains quite expanded, like a coil. In the absence of a polar solvent (water), the polymer chain folds quickly into a dense conformation (polymer globule), and $R_g$ becomes equal to approximately 1.0 nm. The snapshots of the last simulation frame, presenting the PAM coil in water (System 2) and the PAM globule in a vacuum (System 1), are also depicted in Figure 2. It should be noted that in the systems with a high PAM concentration (Systems 7 and 9), the average value $R_g$ slightly decreases compared with the dilute solution containing only 1 PAM macromolecule and equals approximately 1.2 nm. This result is in agreement with the works of Chen and coauthors [46,47], where the value $R_g = 1.20$ nm was obtained for a water solution of a PAM macromolecule with a polymerization degree of 50 at 298 K.

As it may be supposed, changes in the PAM conformation in water or in a vacuum is connected with the PAM ability to form intra- and intermolecular hydrogen bonds (HBs). Using the geometrical criterion of HBs (a donor–acceptor distance less than 0.35 nm, and an acceptor–donor–hydrogen...
angle less than 30°), we calculated the time dependence of PAM–PAM and PAM–water HBs per one monomer unit of PAM in Systems 1 and 2 (Figure 3). Because of PAM folding in the vacuum, the number of HBs increases and at the end of the simulation almost every monomer unit forms one HB with another one. In contrast, in the presence of water, the number of intramolecular HBs between the PAM monomers is significantly smaller and is about 0.3. Although, as we have shown (Figure 2), in water the PAM chain exists in the form of a coil, a certain transformation is observed. The number of intramolecular HBs increases (Figure 3a), and the number of intermolecular PAM–water HBs decreases (Figure 3b). Nevertheless, by the end of the tenth nanosecond, each monomer unit of the polymer forms an average of 2 HBs with the water molecules. Table 2 lists some structural and dynamic characteristics, including average numbers of HBs, for all the systems.

![Figure 2](image-url)  
**Figure 2.** Time dependence of the radius of gyration of PAM (a) in a vacuum (System 1) and water (System 2), the snapshot of the PAM conformations in System 9 at the end of the simulation (b).

![Figure 3](image-url)  
**Figure 3.** Examples of the time dependence of the number of PAM–PAM (a) and PAM–water (b) hydrogen bonds (nHB) per a PAM monomer unit.

The stability of a HB can be characterized by HB lifetimes \( \tau_{HB} \). The continuous mean lifetime of a HB \([48]\), \( \tau_{HB} \), was calculated from the autocorrelation function (ACF) \( C_{HB}(t) \) of the parameter characterizing the HB existence between the \( i \) and \( j \) molecules \( S_{ij}(t) \) using the standard program package:

\[
C_{HB}(t) = \frac{\langle S_{ij}(0)S_{ij}(t) \rangle}{\langle S_{ij}^2(0) \rangle},
\]

(2)

where \( S_{ij}(t) = 1 \) if the criterion of HB existence between the \( i \) and \( j \) molecules was satisfied at the initial moment of time and is satisfied at a current moment of time \( t \), and if the duration of the violated
criterion periods on the time interval from 0 to $t$ did not exceed a predetermined value $t^*$; otherwise, $S(t) = 0$. At $t^* = 0$, the autocorrelation function $C_{HB}(t)$ demonstrates continuous existence of hydrogen bonds. Integration of this function using the standard program package gives the mean HB lifetimes:

$$\tau_{HB} = \int_0^\infty C_{HB}(t)dt,$$

(3)

Table 2. The structural and dynamic characteristics obtained by analyzing the last 10 and 5 ns of the trajectories for water-contained and vacuum systems, respectively.

| System          | 1     | 2     | 3     | 4     | 5     | 6     | 7     | 8     | 9     | 10    |
|-----------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| $<n_{HB}>$ PAM–PAM | 0.76  | 0.30  | 0.35  | 0.66  | 0.40  | 0.76  | 0.31  | 1.00  | 0.32  | 1.03  |
| PAM–Water       | -     | 2.01  | 1.80  | -     | 1.83  | -     | 1.93  | -     | 1.94  | -     |
| PAM–CNC         | -     | -     | 0.08  | 0.32  | 0.033 | 0.22  | 0.02  | 0.76  | 0.04  | 1.03  |
| CNC–Water       | -     | -     | 2.91  | -     | 3.22  | -     | 2.90  | -     | 3.24  | -     |
| CNC–CNC         | -     | -     | 3.61  | 4.26  | 3.31  | 3.88  | 3.66  | 4.3   | 3.40  | 3.86  |
| Water–Water     | -     | 3.57  | 3.54  | 3.56  | -     | -     | -     | -     | 3.57  | -     |
| $\tau$, ps PAM–PAM | 1.71  | 1.47  | 1.52  | 1.58  | 1.62  | 1.61  | 1.45  | 1.63  | 1.44  | 1.67  |
| PAM–Water       | -     | 2.06  | 1.79  | -     | 2.14  | -     | 1.95  | -     | 1.94  | -     |
| PAM–CNC         | -     | -     | 2.40  | 2.85  | -     | 2.83  | 2.37  | 3.07  | 2.14  | 2.81  |
| CNC–Water       | -     | -     | 3.19  | -     | 3.87  | -     | 3.21  | -     | 3.86  | -     |
| CNC–CNC         | -     | -     | 6.50  | 6.62  | 6.87  | 5.64  | 7.09  | 6.88  | 8.01  | 6.87  |
| Water–Water     | -     | 2.81  | 2.77  | -     | 2.77  | -     | -     | -     | 2.77  | -     |
| $R_g$, nm PAM   | 1.00  | 1.48  | 1.28  | 1.19  | 1.34  | 1.30  | 1.21  | 0.95  | 1.22  | 0.96  |
| $R_{e-t-e}$ nm  | 0.99  | 3.56  | 3.55  | 3.04  | 3.56  | 3.75  | 2.78  | 1.75  | 2.79  | 1.81  |
| $<N_c>$ PAM–CNC | -     | 199   | 668   | 5     | 262   | 66   | 1519  | 73    | 1387  |       |
| $<r_{min}>$ nm  | PAM–CNC | -     | 2.07  | 1.67  | 2.19  | 2.17  | 2.06  | 1.62  | 1.97  | 1.63  |

$<n_{HB}>$ and $\tau$ are the average number of hydrogen bonds and lifetime of PAM–PAM and PAM–water hydrogen bonds (per a PAM monomer unit), PAM–CNC, CNC–water, and CNC–CNC ones (per a CNC monomer unit), and water–water ones (per a water molecule); $R_g$ and $R_{e-t-e}$ are the radius of gyration and the end-to-end distance of PAM; $<N_c>$ and $<r_{min}>$ are the average number of close contacts and average minimum distance between any pair of atoms of PAM and CNC within the limits of 0.5 nm.

Table 2 shows that the PAM–water HB lifetime is longer than the PAM–PAM $\tau_{HB}$. Moreover, in water the lifetime of PAM–PAM HBs is shorter than in a vacuum.

In order to understand which sites of molecules take part in HB formation, we calculated the atom–atom radial distribution functions (RDFs). Figure 4 shows the radial distribution functions of oxygen Ow and hydrogen Hw water atoms around the polymer atoms in System 2. Figure 4a compares five intermolecular RDFs for pairs of PAM atoms with water molecules, including H6-Ow, H7-Ow, O4-Hw, N5-Ow, and O4-Ow, where H6, H7, N5, and O4 represent hydrogen, nitrogen, and oxygen atoms of the PAM, respectively (Figure 1a). In Figure 4a, two peaks appear in the vicinity of the carbonyl oxygen, O4, for $g_{O4-Hw}(r)$ and $g_{O4-Ow}(r)$. These peaks are positioned at approximately 0.19, 0.33 nm and 0.29, 0.54 nm for $g_{O4-Hw}(r)$ and $g_{O4-Ow}(r)$, respectively, and represent two hydration shell structures forming around O4. The first peak at 0.19 nm in the $g_{O4-Hw}(r)$ indicates a high probability of the HB formation between O4 and water molecules. At the same time the pre-peaks at 0.24 nm and 0.25 nm in the $g_{H6-Ow}(r)$ and $g_{H7-Ow}(r)$ (which also satisfy the HB geometric criterion $R_{H-O} \leq 0.26$ nm) confirm hydrogen bond formation. However, the height and position of the peaks allow us to suppose the existence of weaker H6-Ow and H7-Ow HBs compared with O4-Hw. Besides, the height of the first peak of $g_{O4-Ow}(r)$ is greater than that of $g_{N5-Ow}(r)$, which also means that hydration of oxygen is stronger than that of nitrogen. The behavior of the C1-Ow, C2-Ow, H89-Ow, and H11-Ow RDFs, namely, the existence of small shoulder peaks at long distances, can be attributed to a highly hydrophobic nature of the PAM hydrocarbon chain. Thus, a comparison of the RDFs confirms that the HBs are
predominantly formed through the PAM carbonyl oxygen and only a few of them are formed through the amide group hydrogen atoms. In Singh’s work [49], the DFT computations were performed on various acrylamide-water clusters in the gas phase in order to explore the microsolvation. In clusters consisting of one acrylamide molecule and 5–15 water molecules, the length of the Ow—H-N HB was ≈ 0.21 and the length of the Hw—O=C HB varied between 0.18 and 0.19 nm. This observation is in agreement with our molecular dynamics results. Also the authors of [49] calculated the binding energy of the acrylamide–acrylamide and acrylamide–water hydrogen-bonded complexes and obtained the values of 6.52 and 12.48 kcal/mol per an acrylamide molecule, respectively. This result is consistent with our findings for the HB lifetime of PAM–PAM and PAM–Water interactions, and explains why PAM macromolecules tend to form intermolecular HBs with water molecules rather than intramolecular HBs.

To prove the existence or absence of the interaction between PAM and CNC in Systems 3–10, we calculated the time evolution of the number of contacts (Figure 5) between any pair of PAM and CNC atoms within a given distance (0.5 nm). In Systems 3, 7, and 9, the adsorption process was successful, and all the three PAM macromolecules in System 3, four PAM molecules in System 7, and one PAM molecule in System 9 were adsorbed on the CNC. In System 3 after 5 ns of the modeling, the number of close contacts increased and at the end of the simulation reached ≈200 (Figure 5a and Table 2). At the same time in System 5, where the CNC surface possesses a predominantly hydrophilic character with a large number of OH-groups, only a few rare episodes of interactions between PAM and CNC were observed (Figure 5a and Table 2). On the contrary, in a vacuum (Figure 5b,d) the PAM molecules were adsorbed on both the hydrophobic (Systems 4, 8) and hydrophilic facets (Systems 6 and 10) of CNC. The number of close contacts in a vacuum was significantly bigger than that in water. This implies that there were competitive interactions in Systems 3, 5, 7, and 9, since the molecules of water and PAM were able to form hydrogen bonds with the oxygen and hydrogen atoms of CNC. Because of the high hydrophilicity of CNC in Systems 5 and 9, almost all the free OH-groups of cellulose were involved in HB formation with the water molecules. Table 2 shows that an average number of HBs for CNC–water was bigger and their lifetime was longer in Systems 5 and 9 than in Systems 3 and 7. Moreover, the CNC–water hydrogen bonds were slightly more stable than those between the water molecules. In general, if we analyze the hydrogen bond lifetimes in all the systems, then we can arrange HBs in the order of increasing stability as follows: $\tau_{HB(PAM–PAM)} < \tau_{HB(PAM–Water)} < \tau_{HB(PAM–CNC)} < \tau_{HB(Water–Water)} < \tau_{HB(CNC–Water)} < \tau_{HB(CNC–CNC)}$. Cellulose chains had a strong affinity with each other and with materials containing hydroxyl groups. Therefore, the lifetimes of the CNC–CNC and CNC–water HBs were higher than the lifetimes of other HBs in the systems. In the work of Chen et al. [50], it was found that water-cellulose HBs are on
average stronger than water–water ones. In particular, they found that $E_{\text{HB}}(\text{Water–Water}) \approx 4.2 \text{ kcal/mol}$, while $E_{\text{HB}}(\text{Water–cellulose}) \approx 5.4 \text{ kcal/mol}$. Having compared $\tau_{\text{HB}}(\text{PAM–CNC})$ for Systems 3 and 4, we can suppose that water weakens the interaction of the polymer with the cellulose nanoparticles.

Figure 5. Time dependence of the number of contacts between any pair of atoms of PAM and CNC within a given distance of 0.5 nm. For System 5 when $\text{NC} = 0$, the red line coincides with the abscissa and therefore is not visible in the figure. (a) Systems 3 and 5, (b) Systems 4 and 6, (c) Systems 7 and 9, (d) Systems 8 and 10.

In order to visualize the process of the polymer adsorption on cellulose, we presented snapshots of the modeling cells for Systems 3–10 at the end of the simulation (Figure 6). To highlight CNC and PAM, all the water molecules from Systems 3, 5, 7, and 9 were removed. In System 3, two PAM macromolecules were adsorbed on the top hydrophobic side of CNC and one macromolecule was adsorbed on the bottom hydrophobic side of CNC. The same picture was observed for System 4, i.e., for PAM–CNC composite in a vacuum. The losses of CNC–water intermolecular hydrogen bonds were compensated with new hydrogen bonds with PAM, and we observed a significant increase in $<n_{\text{HB}}>$ for PAM–CNC (Table 2). On the other hand, although PAM molecules interacted with a hydrophilic facet of CNC more actively in a vacuum than in water, a large part of the CNC surface remained free, and the number of PAM–CNC close contacts in System 6 was three times less than in System 4. Besides, the average minimal distance of the PAM contact with a CNC hydrophilic facet, $<r_{\text{min}}>$, was approximately 1.3 times longer than with a hydrophobic one (Table 2). At the same time, in the systems with a high polymer concentration (Systems 7–10), the difference between the average numbers of closest contacts was not so noticeable for hydrophobic and hydrophilic CNC, i.e., the probability to find PAM near CNC significantly increased and almost all the accessible HB centers of CNC were occupied by PAM. Table 2 clearly shows that the average numbers of PAM–CNC HBs per one CNC monomer in Systems 8 and 10 were equal to 0.76 and 1.03, respectively, which is $\approx 2.4$ and $\approx 4.7$ times higher than in the systems with a deficiency of PAM molecules. The difference between the $<N_{\text{C}}>$ values for Systems 8 and 10 (1519 and 1387, respectively) is due to the fact that the area of the available surface for the polymer is smaller for hydrophilic CNC. Additionally, with gmx_sasa tool from Gromacs-5.0.7 we calculated the total accessible solvent surface area (SASA) for both CNC types: SASA (hydrophobic CNC) = 73.1 nm$^2$ and SASA (hydrophilic CNC) = 58.4 nm$^2$.

Figure 7 shows the places of the polymer direct contact with the cellulose in the systems. It can be seen that in the presence of the solvent the contact areas between the polymer and cellulose are much smaller than in the vacuum. Most of the CNC solvation shell in System 3 is water molecules (Figure 7).
Figure 6. Snapshots of the last frame for Systems 3–10. The water molecules from Systems 3, 5, 7, and 9 were removed to highlight CNC (orange) and PAM. For better understanding, the hydrophobic (200) and hydrophilic (110, 1–10) facets are marked.

Figure 7. Snapshots of solvation shells at a distance of 0.5 nm from CNC (yellow) for System 3, System 4, System 6, System 7, System 9, System 8, System 10. The ice blue surface denotes the water shell and the red surface denotes the places of the PAM direct contact with CNC. For better understanding, the hydrophobic (200) and hydrophilic (110, 1–10) facets are marked.
It should be noted that in our previous work devoted to studying the adsorption of PVP on CNC, we also observed the crucial role of water in the interaction between PVP and CNC [26]. In particular, we showed that the presence of water makes the interaction between the polymer and cellulose weaker than in a vacuum, and the polymer and cellulose mainly interact through their solvation shells. Unlike PVP, which only has a hydrogen acceptor atom, PAM is able to act as a proton donor because of the amide group hydrogen atoms, which increases the probability of HB formation with the cellulose atoms. To characterize the interaction between PAM and CNC atoms in greater detail, we also analyzed RDFs for Systems 3–6 (Figures 8–10). The RDFs for Systems 7–10 demonstrate the same behavior; that is why they are not shown here. As in the case of PAM–water interactions, the carbonyl oxygen O4 takes the most active part in the HB formation with the cellulose H1, H2, H3 atoms (Figure 8). The typical peaks, which can be found on the RDF curves $g_{H7O1}(r)$, $g_{H7O2}(r)$ at $\approx 0.23$–$0.24$ nm, indicate that the H7 atom also actively interacts with CNC through the O1 and O2 atoms (Figure 8).

**Figure 8.** Atom–atom radial distribution functions for PAM–CNC in System 3.

**Figure 9.** Atom–atom radial distribution functions for PAM–CNC in System 4.
In the absence of water (System 4) all the peaks on the RDF curves become more pronounced and increased compared with those for System 3, which confirms that more HBs are formed between the PAM and CNC in a vacuum. A dramatic increase in the peak heights on \( g_{H7O2}(r), g_{H7O3}(r), g_{O4H2}(r), g_{O4H3}(r), g_{O4H4}(r) \) (Figure 9) is observed. It means that in water most of the cellulose hydroxyl groups O2H2 and O3H3 are involved in the HBs with the solvent molecules rather than the polymer ones. As for the amide hydrogen atoms, the probability of the H7 atom participation in HB formation is higher than for the H6 atom, which is located closer to the carbonyl oxygen O4 and may have a steric hindrance to the HB formation. System 6 does not have such preferences for HB formation through cellulose hydroxyl groups O2H2, O3H3, O1H1 (Figure 10), with the exception of the ended O6H4 hydroxyl group (Figure 1b).

4. Conclusions

Classical molecular dynamics simulations of polyacrylamide (PAM) adsorption on a cellulose nanocrystal (CNC) were carried out. Ten different systems containing PAM and CNC in a vacuum or in water were simulated. The average number of hydrogen bonds and lifetime of PAM–PAM, PAM–water, PAM–CNC, CNC–water, CNC–CNC, and water–water hydrogen bonds were calculated. The PAM radius of gyration, end-to-end distance, and the average number of close contacts and average minimum distance between any pair of atoms of PAM and CNC were determined. The atom–atom radial distribution functions for PAM, PAM–CNC in a vacuum and water were analyzed. It was confirmed that hydrogen bonds are predominantly formed through carbonyl oxygen of PAM and the \(-\text{NH}_2\) group is the less active participant of the HB-formation process in the systems. It was noted that PAM macromolecules tend to form intermolecular hydrogen bonds with water molecules rather than intramolecular ones. Moreover, even in the systems with a high PAM concentration the number of PAM–CNC hydrogen bonds is low, despite the fact that such HBs are more stable than PAM–PAM and PAM–water HBs. Based on the analysis of the PAM radius of gyration it can be concluded that the polymer spends more time in a more extended configuration in water and on the CNC surface (\(R_g \geq 1.19\) nm) than in a vacuum at some distance from the CNC (\(R_g \leq 1.0\) nm). The crucial role of
water in the interaction between PAM and CNC was highlighted. It was found out that water and PAM sorption on CNC is a competitive process, and water weakens the interaction between the polymer and CNC.

Supplementary Materials: The following are available online at http://www.mdpi.com/2079-4991/10/7/1256/s1, Figure S1. Time dependence of the system temperature. Figure S2. The instant snapshots of System 6 at 7 ns (a) and 7.8 ns (b). These snapshots explain the jump in NC(t) at ≈ 7.5 ns.

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References

1. Schaefer, D.W.; Justice, R.S. How nano are Nanocomposites? Macromolecules 2007, 40, 8501–8517. [CrossRef]
2. Kumar, S.K.; Krishnamoorti, R. Nanocomposites: Structure, phase behavior, and properties. Annu. Rev. Chem. Biomol. Eng. 2010, 1, 37–58. [CrossRef] [PubMed]
3. Moniruzzaman, M.; Winey, K.I. Polymer Nanocomposites containing carbon nanotubes. Macromolecules 2006, 39, 5194–5205. [CrossRef]
4. Ganesan, V. Some issues in polymer nanocomposites: Theoretical and modeling opportunities for polymer physics. J. Polym. Sci. Part B 2008, 46, 2666–2671. [CrossRef]
5. Vogiatzis, G.G.; Theodorou, D.N. Multiscale molecular simulations of polymer-matrix Nanocomposites or what molecular simulations have taught us about the fascinating Nanoworld. Arch. Comput. Methods Eng. 2018, 25, 591–645. [CrossRef]
6. Sgouros, A.P.; Vogiatzis, G.G.; Megariotis, G.; Tzoumanekas, C.; Theodorou, D.N. Multiscale simulations of graphite-capped polyethylene melts: Brownian dynamics/kinetic Monte Carlo compared to atomistic calculations and experiment. Macromolecules 2019, 52, 7503–7523. [CrossRef]
7. Sgouros, A.P.; Vogiatzis, G.G.; Kritikos, G.; Boziki, A.; Nikolakopoulou, A.; Liveris, D.; Theodorou, D.N. Molecular simulations of free and graphite capped polyethylene films: Estimation of the interfacial free energies. Macromolecules 2017, 50, 8827–8844. [CrossRef]
8. Theodorou, D.N.; Vogiatzis, G.G.; Kritikos, G. Self-consistent-field study of adsorption and desorption kinetics of polyethylene melts on graphite and comparison with atomistic simulations. Macromolecules 2014, 47, 6964–6981. [CrossRef]
9. Julkapli, N.M.; Bagheri, S. Progress on nanocrystalline cellulose biocomposites. React. Funct. Polym. 2017, 112, 9–21. [CrossRef]
10. Moon, R.J.; Martini, A.; Nairn, J.; Simonsen, J.; Youngblood, J. Cellulose nanomaterials review: Structure, properties and nanocomposites. Chem. Soc. Rev. 2011, 40, 3941–3994. [CrossRef]
11. Rojas, O.J.; Lokanathan, A.R.; Kontturi, E.; Laine, J.; Bock, H. The unusual interactions between polymer grafted cellulose nanocrystal aggregates. Soft Matter 2013, 9, 8965–8973. [CrossRef]
12. Zhu, C.; Monti, S.; Mathew, A.P. Evaluation of nanocellulose interaction with water pollutants using nanocellulose colloidal probes and molecular dynamic simulations. Carbohyd. Polym. 2020, 229, 115510. [CrossRef]
13. Mehandzhiyski, A.Y.; Rolland, N.; Garg, M.; Wohlert, J.; Linares, M.; Zozoulenko, I. A novel supra coarse-grained model for cellulose. Cellulose 2020, 27, 4221–4234. [CrossRef]
14. Rolland, N.; Mehandzhiyski, A.Y.; Garg, M.; Linares, M.; Zozoulenko, I.V. New patchy particle model with anisotropic patches for molecular dynamics simulations: Application to a coarse-grained model of cellulose Nanocrystal. J. Chem. Theory Comput. 2020, 16, 3699–3711. [CrossRef] [PubMed]
15. Bergensträhle, M. Crystalline Cellulose in Bulk And at Interfaces as Studied by Atomistic Computer Simulations. Doctoral Thesis, KTH School of Chemical Science and Engineering, Stockholm, Sweden, 2008.
16. Qiao, Q.; Li, X.; Huang, L. Crystalline cellulose under pyrolysis conditions: The structure–property evolution via reactive molecular dynamics simulations. J. Chem. Eng. Data 2020, 65, 360–372. [CrossRef]

17. Muthoka, R.M.; Kim, H.C.; Kim, J.W.; Zhai, L.; Pooja, S.; Panicker, P.S.; Kim, J. Steered pull simulation to determine Nanomechanical properties of cellulose Nanofiber. Materials 2020, 13, 710. [CrossRef]

18. Sinko, R.; Keten, S. Traction–separation laws and stick–slip shear phenomenon of interfaces between cellulose nanocrystals. J. Mech. Phys. Solids 2015, 78, 526–539. [CrossRef]

19. Yu, Y.; Tyrikos-Ergas, T.; Zhu, Y.; Fittolani, G.; Bordoni, V.; Singhal, A.; Fair, R.J.; Grafmüller, A.; Seeberger, P.H.; Delbianco, M. Systematic hydrogen bond manipulations to establish polysaccharide structure-property correlations. Angew. Chem. Int. Ed. 2019, 131, 13261–13266. [CrossRef]

20. Mazeau, K.; Heux, L. Molecular dynamics simulations of bulk native crystalline and amorphous structures of cellulose. J. Phys. Chem. B 2003, 107, 2394–2403. [CrossRef]

21. Kulasinski, K.; Keten, S.; Churakov, S.V.; Derome, D.; Carmeliet, J. A comparative molecular dynamics study of crystalline, paracrystalline and amorphous states of cellulose. Cellulose 2014, 21, 1103–1116. [CrossRef]

22. Crawford, B.; Ismail, A.E. Insight into cellulose dissolution with the tetrabutylphosphonium chloride–water mixture using molecular dynamics simulations. Polymers 2020, 12, 627. [CrossRef] [PubMed]

23. Sahputra, I.H.; Alexiadis, A.; Adams, M.J. Effects of moisture on the mechanical properties of microcrystalline cellulose and the mobility of the water molecules as studied by the hybrid molecular mechanics–molecular dynamics simulation method. J. Polym. Sci. Pol. Phys. 2019, 57, 454–464. [CrossRef]

24. Khazraji, A.C.; Robert, S. Interaction effects between cellulose and water in nanocrystalline and amorphous regions: A novel approach using molecular modeling. J. Nanomater. 2013, 409676. [CrossRef]

25. Krishna, S.; Patel, C.M. Computational and experimental study of mechanical properties of Nylon 6 nanocomposites reinforced with nanomilled cellulose. Mech. Mater. 2020, 143, 103318. [CrossRef]

26. Gurina, D.; Surov, O.; Voronova, M.; Zakharov, A.; Kiselev, M. Water effects on molecular adsorption of Poly(N-vinyl-2-pyrrolidone) on cellulose Nanocrystals surfaces: Molecular dynamics simulations. Materials 2019, 12, 2155. [CrossRef]

27. Voronova, M.; Rubleva, N.; Kochkina, N.; Afineevskii, A.; Zakharov, A.; Surov, O. Preparation and characterization of Poly(nvinylpyrrolidone)/Cellulose nanocrystals. Nanomaterials 2018, 8, 1011. [CrossRef]

28. Schuberth, S.; Münstedt, H. Transient elongational viscosities of aqueous polycrystalline solutions measured with an optical rheometer. Rheol. Acta 2008, 47, 139–147. [CrossRef]

29. Xu, L.N. Viscoelasticity of mixed polycrystalline solution. J. Cent. South Univ. Technol. 2008, 15, 443–446. [CrossRef]

30. Van Der Spoel, D.; Lindahl, E.; Hess, B.; Groenhof, G.; Mark, A.E.; Berendsen, H.J.C. Gromacs: Fast, flexible, and free. J. Comput. Chem. 2005, 26, 1701–1718. [CrossRef] [PubMed]

31. Humphrey, W.; Dalke, A.; Schulten, K. VMD: Visual molecular dynamics. J. Mol. Graph. 1996, 14, 33–38. [CrossRef]

32. Nose, S. A Molecular dynamics method for simulations in the canonical ensemble. Mol. Phys. 1984, 52, 255–268. [CrossRef]

33. Hoover, W.G. Canonical dynamics: Equilibrium phase-space distributions. Phys. Rev. A 1985, 31, 1695–1697. [CrossRef] [PubMed]

34. Allen, M.P.; Tildesley, D.J. Computer Simulation of Liquids; Clarendon Press: London, UK, 1987; 385p.

35. Darden, T.; York, D.; Pedersen, L. Particle mesh Ewald: An N-log(N) method for Ewald sums in large systems. J. Chem. Phys. 1993, 98, 10089–10092. [CrossRef]

36. Essmann, U.; Perera, L.; Berkowitz, M.L.; Darden, T.; Lee, H.; Pedersen, L.G. A smooth particle mesh Ewald method. J. Chem. Phys. 1995, 103, 8577–8592. [CrossRef]

37. Hess, B.; Bekker, H.; Berendsen, H.J.; Fraaije, J.G. LINCS: A linear constraint solver for molecular simulations. J. Comput. Chem. 1997, 18, 1463–1472. [CrossRef]

38. Jorgensen, W.L.; Tirado-Rives, J. The OPLS potential functions for proteins, energy minimizations for crystals of cyclic peptides and crombin. J. Am. Chem. Soc. 1988, 110, 1657–1666. [CrossRef]

39. Hanwell, M.D.; Curtis, D.E.; Lonie, D.C.; Vandermeersch, T.; Zurek, E.; Hutchison, G.R. Avogadro: An advanced semantic chemical editor, visualization, and analysis platform. J. Cheminform. 2012, 4, 17. [CrossRef]

40. Malde, A.K.; Zuo, L.; Breeze, M.; Stroet, M.; Poger, D.; Nair, P.C.; Oostenbrink, C.; Mark, A.E. An automated force field topology builder (ATB) and repository: Version 1.0. J. Chem. Theory Comput. 2011, 7, 4026–4037. [CrossRef]
41. Nishiyama, Y.; Langan, P.; Chanzy, H. Crystal structure and hydrogen-bonding system in cellulose Iβ from synchrotron X-ray and neutron fiber diffraction. *J. Am. Chem. Soc.* 2002, 124, 9074–9082. [CrossRef]

42. Gomes, T.C.F.; Skaf, M.S. Cellulose-Builder: A toolkit for building crystalline structures of cellulose. *J. Comput. Chem.* 2012, 33, 1338–1346. [CrossRef]

43. Berendsen, H.J.C.; Grigera, J.R.; Straatsma, T.P. The missing term in effective pair potentials. *J. Phys. Chem.* 1987, 91, 6269–6271. [CrossRef]

44. Fixman, M. Radius of gyration of polymer chains. *J. Chem. Phys.* 1962, 36, 306–310. [CrossRef]

45. Cooper, A.I. Polymer synthesis and processing using supercritical carbon dioxide. *J. Mater. Chem.* 2000, 10, 207–234. [CrossRef]

46. Chen, P.; Yao, L.; Liu, Y.; Luo, J.; Zhou, G.; Jiang, B. Experimental and theoretical study of dilute polyacrylamide solutions: Effect of salt concentration. *J. Mol. Model.* 2012, 18, 3153–3160. [CrossRef]

47. Liu, Y.Y.; Chen, P.K.; Luo, J.H.; Zhou, G.; Jiang, B. Molecular simulation of dilute Polyacrylamide solutions. *Acta Phys. Chim. Sin.* 2010, 26, 2907–2914. [CrossRef]

48. Rapaport, D.C. Hydrogen bonds in water: Network organization and lifetimes. *Mol. Phys.* 1983, 50, 1151–1162. [CrossRef]

49. Singh, S.; Srivastava, S.K.; Singh, D.K. Hydrogen bonding patterns in different acrylamide–water clusters: Microsolvation probed by micro Raman spectroscopy and DFT calculations. *RSI Adv.* 2014, 4, 1761–1774. [CrossRef]

50. Chen, M.; Coasne, B.; Guyer, R.; Derome, D. Role of hydrogen bonding in hysteresis observed in sorption-induced swelling of soft nanoporous polymers. *Nat. Commun.* 2018, 9, 3507. [CrossRef]