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Structure, Orientation and Stability of Lysozyme Confined in layered Materials

Sébastien Balme\textsuperscript{a,b,*}, Régis Guégan\textsuperscript{c,*}, Jean-Marc Janot\textsuperscript{a}, Maguy Jaber\textsuperscript{d}, Mathilde Lepoitevin\textsuperscript{a}, Philippe Dejardin\textsuperscript{a}, Xavier Bourrat\textsuperscript{c}, Mikael Motelica-Heino\textsuperscript{c}

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The confinement of lysozyme in 3 layered materials based on montmorillonite and lamellar double hydroxide exhibiting different surface charges was studied. The protein structure and orientation in these materials were determined by X-Ray diffraction, time resolved fluorescence and fluorescence anisotropy. For both Montmorillonite exchanged with sodium and modified with non-ionic surfactant (tri-ethylene glycol mono n-decyl ether), the lysozyme was found to be located in the interlayer space with “end-on” and “side-on” orientation, respectively. Conversely no lysozyme intercalation was observed with lamellar double hydroxide modified with anionic surfactant (sodium octylsulfate), since protein was adsorbed on the surface of the particles. Fourier transformed infra-red spectroscopy analysis shows that lysozyme confinement in the interlayer space preserves its structure after dehydration, whereas some structural changes were observed for lysozyme adsorbed on the particle surface.

Introduction

The confinement of biomolecules in inorganic porous materials, for the synthesis of novel nanocomposites with particular properties or the manufacture of biosensors, has attracted considerable attention in the past decade.\textsuperscript{1-12} Indeed, many studies have shown the multiple possibilities and applications of these hybrid materials where guest molecules are confined into various inorganic host matrices such as metallic phases and porous alumina or silica.\textsuperscript{9,13-15} Moreover, layered materials are of special interest due to their high internal surface areas, their abilities to confine guest biomolecules within regularly organized interlayer spaces, and to undergo self-orientation at a macroscopic scale. Thus natural and chemically modified clay minerals, i.e. organo-clay materials, have been largely exploited for the intercalation of biological molecules such as amino acids or proteins to build biosensors or to spread clay-protein films on electrodes for electrochemical catalysis.\textsuperscript{1-4,6} The intercalation of biomolecules in layered materials is mainly driven by electrostatic interactions, through the substitution of inorganic ions. The chemical modification of clay minerals or LDHs by intercalation of surfactants leads preferentially to hydrophobic surfaces, which may induce structural modification of confined proteins.

Time resolved fluorescence and anisotropy of fluorescence are powerful techniques for investigating structure and dynamics of biological macromolecules.\textsuperscript{16,17} However these techniques are not frequently used to study the interactions between proteins and inorganic surfaces.\textsuperscript{18-21} The photophysical properties of amino-acids such as tryptophan (Trp) are indeed directly affected by their neighborhood. Investigations of the Trp fluorescence lifetime provide access to fundamental aspects of the proteins structural modifications or protein/ligand interactions. In addition, fluorescence anisotropy, which is based on time dependence of the light depolarization emitted from a fluorophore,\textsuperscript{22,23} can be used to separate the contribution of the entire macromolecule from local dynamic motions.\textsuperscript{24,25}

The aim of this paper is to study the structure modification, orientation and stability of a hard model protein (Lysozyme) induced by its adsorption on different layered materials: sodium homoionic montmorillonite (Na-Mt), montmorillonite with non-ionic surfactant tri-ethylene glycol mono n-decyl ether (C\textsubscript{10}E\textsubscript{12}-Mt), and LDH modified with sodium octylsulfate (SOS-LDH). Lysozyme is a small (MW 14.4 kDa) globular protein. Its crystallographic structure was the first to be established by X-ray diffraction (XRD) methods in 1965.\textsuperscript{26} This discovery opened very quickly the opportunity for researchers to correlate its known structure with its fluorescence properties and thus, nowadays its photophysical behavior is well known.\textsuperscript{27-30}

Materials and methods

Layered materials preparation

The natural clay mineral, montmorillonite (Mt) extracted from the Crook county (Wyoming), was purchased from the Source Clays Repository (Clay mineral Society ref: Srce_Clay_SWy-2). Its cation exchange capacity (CEC) was reported to be about 75meq/100g.\textsuperscript{31} The present work was carried out on clay mineral fraction ≤ 2 µm obtained by dispersion, sedimentation and centrifugation. Cation exchanges were performed in salt solution, i.e. 1 M, made from pure NaCl (Sigma, 99.9%) and de-ionized water. The sample (Na-Mt) preparation and elemental analysis are described in detail elsewhere.\textsuperscript{32}

The organo-Mt modified with nonionic surfactant was synthesized following the procedure of Guegan.\textsuperscript{33} A sodium homoionic Montmorillonite suspension was dispersed in a solution of nonionic surfactant tri-ethylene glycol mono n-decyl ether (C\textsubscript{10}E\textsubscript{12}) (Nikko Chemicals, Inc., Tokyo, Japan) at a concentration above its critical micelle concentration.\textsuperscript{34} In such a way, the resulting nonionic organo-Mt (C\textsubscript{10}E\textsubscript{12}-Mt) preserves Na\textsuperscript{+} cations in its interlayer space for further ion exchange, while
Adsorption of protein on layered materials

Lysozyme from chicken egg white was purchased from Sigma (62971). Protein adsorption experiments were performed in phosphate buffer saline solution (PBS) 50 mM, NaCl 150 mM at pH 8.7 for all layered material with the same experimental protocol. First, 11 mL of protein solutions were prepared at 1 mg mL\(^{-1}\) and 24°C. 1 mL of this solution was used to determine the concentration of protein in solution at time \(t_0\). Then 10 mg of layered material was dispersed in the remaining 10 mL of protein solution. After some chosen time \(t\), 1 mL of the sample was centrifuged for 8 min at 60000 rpm. The concentration of lysozyme in the supernatant was determined by UV absorption at 280 nm (\(\varepsilon_{280} = 36000\) M\(^{-1}\) cm\(^{-1}\)). The concentration of protein adsorbed \(\Gamma(t)\) on hybrid materials (in mg of protein per g of layered material) was determined by the method of supernatant depletion.

\[
\Gamma(t) = \frac{V(C_{t0} - C_{t1})}{m} \tag{eq 1}
\]

Where \(V\) is the volume of the protein solution (mL), \(C_{t0}\) (mg mL\(^{-1}\)) is the initial concentration of protein, \(C_{t1}\) (mg mL\(^{-1}\)) is the supernatant protein concentration at time \(t\), and \(m\) is the mass of the hybrid layered materials (g).

Fourier transform infrared spectroscopy (FTIR)

FTIR spectra, in the range 650-4000 cm\(^{-1}\), were recorded using a Thermo Nicolet 6700 FT spectrometer equipped with a Deuterated Triglycine Sulfate (DTGS) detector and a Nicolet Continuum microscope. The powder samples were spread over an NaCl window of the microscope. The analyzed sample area was a square of side 100 \(\mu\)m chosen under the microscope 15X Infinity Reflexchromat objective. The analyses were performed in transmission mode and each spectrum was the average of 128 scans collected at 2 cm\(^{-1}\) resolution. FTIR was performed on dry samples (25°C under vaccum 2 hours).

X-ray Diffraction (XRD)

XRD patterns of hybrid layered materials were measured using an ARL XTRA Diffractometer (Thermo Electron Corporation, Ecublens, Switzerland) equipped with a Cu anode and an Si solid detector operating at 40 kV and 40 mA. Experimental measurements were carried out at room temperature at 10s counting time per 0.05°(20) step. XRD was performed on dry sample (25°C under vaccum 2 hours) such as FTIR experiments.

UV absorbance and Fluorescence emission spectra

UV absorbance spectra were recorded on a JASCO V-570 UV-Vis spectrophotometer at scanning rate 1 nm s\(^{-1}\). All solutions were filtered on a low protein binding Durapore\textsuperscript{®} membrane (Millex-HV 0.45\(\mu\)m PDMF) to limit the light scattering due to clay microparticles. Moreover, filters were previously flushed with a protein solution to limit the error due to non specific protein adsorption on the filter.

Time-resolved fluorescence

Time-resolved fluorescence data were obtained by the time-correlated single-photon counting technique\textsuperscript{24,25}. The excitation wavelength was achieved by using an 82 MHz mode-locked argon ion laser (Spectra Physics model 2030) pumping synchronously a cavity-dumped rhodamine 6G dye laser (Spectra Physics model 375 dye laser and 344 cavity dumper). The UV excitation light, set to vertical polarization, was obtained using a dilute suspension of polystyrene latex (Millex 70 nm diameter) as a scattering solution; it was typically 220 pm FWHM. The emission of fluorescence was detected, after passing through a polarizer set parallel, perpendicular or magic angle (54.73°) relative to vertical excitation, through a double monochromator Jobin–Yvon DH10 on a Hamamatsu H5323 photomultiplier (Cathode S20, TTS 160 ps). The instrumental response function of the equipment was measured by using a dilute suspension of polystyrene nanospheres in water (70 nm diameter) as a scattering solution; it was typically 220-250 ps FWHM. Decays were collected at a maximum count rate of 15 kHz into 2048 channels. This limiting count rate was achieved by dilution of the sample in order to minimize as much as possible the scattering of the particles and after sedimentation of the suspension. The time per channel was set around 18 ps ch\(^{-1}\)

### Table 1 Characteristics of the layered materials

<table>
<thead>
<tr>
<th>Layered Materials</th>
<th>Na-Mt</th>
<th>C\textsubscript{10}E\textsubscript{10}-Mt</th>
<th>SOS-LDH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inorganic host material</td>
<td>((\text{Si}3\text{Al})\cdot(\text{Al}3\text{Fe}3\text{O}10\cdot\text{Mg}2\text{SOS}))\textsuperscript{VI}</td>
<td>((\text{Si}3\text{Al})\cdot(\text{Al}3\text{Fe}3\text{O}10\cdot\text{Mg}2\text{SOS}))\textsuperscript{VI}</td>
<td>((\text{Mg}2\text{SOS})\cdot(\text{Al}3\text{O}10\cdot\text{H}2\text{O}))</td>
</tr>
<tr>
<td>Surfactant</td>
<td>-</td>
<td>(\text{C}<em>{10}\text{E}</em>{10})</td>
<td>SOS</td>
</tr>
<tr>
<td>Organic content (%)</td>
<td>-5</td>
<td>-38</td>
<td>~55</td>
</tr>
<tr>
<td>(d_0) (nm)</td>
<td>1.2</td>
<td>3.6</td>
<td>3.2</td>
</tr>
<tr>
<td>Surfactant arrangement</td>
<td>-</td>
<td>bilayer</td>
<td>bilayer</td>
</tr>
</tbody>
</table>
in order to have a full decay in the experimental time window. All decays were collected in order to have at least 1.5x10^6 counts in total. The excitation wavelength was set at 294 nm and the collected emission at 345 nm for all samples.

Decay analysis was performed using a Levenberg–Marquardt algorithm. For the analysis the fluorescence decay law at the magic angle \( I_M(t) \) and the anisotropy decay law \( r(t) \) were analyzed as a sum of exponentials:

\[
I_M(t) = \frac{1}{2} \sum_{i=1}^{n} a_i e^{-t/\tau_i} \quad \text{(eq 2)}
\]

\[
r(t) = \sum_{j=1}^{n} r_j e^{-t/\phi_j} \quad \text{(eq 3)}
\]

With \( \tau_i \) the fluorescence lifetime, \( a_i \) the pre-exponential factor related to the contribution of each lifetime of component \( i \), \( r_j \) the anisotropy of a component \( j \) for a correlation time \( \phi_j \).

The anisotropy decay, \( r(t) \), is related to the decays collected with vertical, \( I_V(t) \), and horizontal, \( I_H(t) \), polarization, the excitation being vertical:

\[
I_V(t) = I_M(t) \{1 + 2r(t)\} \quad \text{(eq 4)}
\]

\[
I_H(t) = I_M(t) \{1-r(t)\} \quad \text{(eq 5)}
\]

or

\[
r(t) = \frac{I_V(t)-I_H(t)}{I_V(t)+2I_H(t)} \quad \text{(eq 6)}
\]

Fluorescence lifetimes were calculated from data collected at magic angle by iterative adjustment after convolution\(^{37}\) of a pump profile (scattered light) with a sum of exponentials. The anisotropy decay function was determined using the parameters obtained from the analysis of the magic angle decay and simultaneous adjustment of vertical and horizontal data.\(^{24}\) We assumed a Poisson distribution of counts in the calculation of the \( \chi^2 \) criterion; residuals profiles and autocorrelation function as well as Durbin-Watson and skewness factor,\(^{38-40}\) were used in order to estimate the quality of the adjustment. The number of exponentials used for the fit was increased until all the statistical criterions were improved. In order to enhance the fit of the whole decay, the shift that experimentally occurs between the pump profile and the decay was also introduced as a fitting parameter. After adjustment and as a control, the anisotropy decay functions \( r(t) \) calculated from the experimental data and from the extracted parameters were displayed.

Fluorescence experiment was performed phosphate buffer saline solution (PBS) 50mM, NaCl 150 mM at pH 8.7.

**Results and discussion**

**Protein adsorption**

The adsorption of lysozyme was performed below its isoelectric point (IP =10.7) at pH 8.7. Under these conditions, the protein is globally positively charged.\(^{41,42}\) The kinetics of lysozyme adsorption on different layered materials are depicted in Fig. 1. As expected, the kinetics of lysozyme adsorption are fast on Na-Mt. Indeed, this clay mineral is globally negatively charged with exchangeable cations within the interlayer space. Furthermore, the equilibrium state is reached after 90 min and the amount of loaded protein reaches \( \Gamma_{lys} = 791 \text{mg/g Na-Mt} \). Electrostatic interactions between the positively charged lysozyme and the negatively charged clay mineral occurred and resulted in an exchange with the inorganic interlayer cations.\(^{43}\)

After intercalation of C\(_{10}\)E\(_7\) nonionic surfactant, the hydrophobic character of the clay mineral increases. The concentration of adsorbed lysozyme on C\(_{10}\)E\(_7\)-Mt after 450 min is 550 mg/g\( \text{C}_{10}\text{EO}_{13}\)-Mt without reaching equilibrium as for SOS-LDH (300 mg/g\( \text{SOS-LDH} \) at 450 min).

**Protein localization in layered materials in dry state.**

Intercalation of lysozyme into the interlayer space of Na-Mt and C\(_{10}\)E\(_7\)-Mt was characterized by using powder X-ray diffraction. The XRD patterns exhibit an increase of the (001) basal spacing from 1.2 to 5.4 nm and from 3.6 to 4.6 nm for the Na-Mt and C\(_{10}\)E\(_7\)-Mt, respectively (Fig. 2). These increases after intercalation are compatible with the dimension of the lysozyme (4.5 x 3 x 3 nm\(^3\)) and suggest a complete insertion of the protein within the interlayer space. In addition a difference of basal spacing 0.8 nm between both layered materials is observed. The adsorption of lysozyme on Na-Mt and C\(_{10}\)E\(_7\)-Mt involves cation exchange with the sodium counter ions, with complete insertion of the protein within the interlayer space. The adsorption of lysozyme on C\(_{10}\)E\(_7\)-Mt is partly driven by electrostatic interactions as for Na-Mt. In addition to these electrostatic interactions, the protein polar residues interact with the C\(_{10}\)E\(_7\)-Mt hydrophobic surface. The slower and lower lysozyme adsorption on C\(_{10}\)E\(_7\)-Mt than on Na-Mt may be explained by the previous intercalation of C\(_{10}\)E\(_7\) surfactant within the interlayer space which hinders the access to the interaction sites and limits the accessible surface.

On the contrary, the X-ray diffraction patterns (not shown) for the SOS-LDH did not display any particular differences with or without protein; the (001) reflections remained at a similar 2\( \theta \) angle; this suggest that the adsorption is only occurring on the external surface of the particles. This result is not surprising since no cation exchange can occur between the positively charged protein and the LDH. Indeed, the SOS-LDH is characterized by a hydrophobic surface and a weak anionic exchange capacity since carbonate can easily be re-intercalated. In this case, the adsorption process may not be driven by the electrostatic interactions. Indeed, the positively charged protein can only be
adsorbed on the external surface of the SOS-LDH and cannot be exchanged with the carbonates located in the interlayer space.

![Graph](image)

**Fig. 2:** XRD patterns for (a) Na-Mt in black line, (b) Na-Mt with lysozyme in black dash line, (c) C10E1-Mt synthesized by using C10E1 surfactant in gray line and (d) C10E1-Mt with lysozyme in gray dash line. Large peaks for the layered materials without lysozyme (a), (c) and with lysozyme (b), (d) correspond to d001 reflections at 1.2 or 3.6 nm, and at 5.4 and 4.6 nm.

**Stability of lysozyme in layered materials in dry state.**

The confinement of proteins into the interlayer space of lamellar materials can induce an increase in their stability. One consequence can be that they keep their conformational structure even under lack of water. FTIR is a powerful technique for the determination of the conformational changes of proteins in interaction with surfaces. Indeed it is sensitive to modification of ratio α-helix versus β-sheet. Infrared spectra of dry samples of lysozyme loaded on layered materials are reported in Fig. 3. The insert highlights the amide bands I and II of the peptide backbones. These bands are very sensitive to conformational changes of proteins and are used to probe their modification of structures. These carbonyl vibration modes in the amide bands I and II usually are observed for lysozyme in buffer solution at 1600 and 1525 cm⁻¹, respectively. For intercalated lysozyme in Na-Mt and C10E1-Mt, these vibration modes remained at the same frequency which means a constant ratio α-helix versus β-sheet with respect to solution.

These results suggest that no strong conformational change is occurring when the water is removed. This stabilization of the structure of the protein is induced by the confinement. Indeed, hard proteins are often stabilized when they are confined in porous materials. Thus it can be assumed that lysozyme almost keeps its geometry, even if small conformational changes cannot be dismissed. Regarding XRD experiments (Fig 2), which are performed under same condition than FTIR, the difference of basal spacing 0.8 nm obtained in between Na-Mt and C10E1-Mt could be interpreted as the difference in between the model of “end-on” and “side-on” orientations (Fig. 4a and b).

In contrast, the FTIR spectra of lysozyme adsorbed on external surface of SOS-LDH show noticeable shifts in frequencies (between 5 and 10 cm⁻¹) for the peptide backbone vibration in the amide bands. This indicates a modification of the conformation of the adsorbed proteins. As previously discussed, lysozyme adsorption on the external surface (Fig 4c) of SOS-LDH does not seem to induce this unfolding. Thus the dehydration may induce partial structural changes of lysozyme because the protein is not protected inside the interlayer space.

![Graph](image)

**Fig. 3** Infrared spectra of lysozyme confined in (a) Na-Mt, (b) C10E1-Mt, and (c) adsorbed on SOS-LDH particles obtained after dehydration compared to native lysozyme (Lyz). The insert represent a zoom of the lysozyme carbonyl vibration modes, associated with the peptide backbone vibration in the amide bands.
Protein conformation and orientation in layered materials in buffer suspension.

Fluorescence properties of the lysozyme tryptophans:

Background.

The amino acid sequence of lysozyme contains six tryptophan residues; three of them are localized in the active site, Trp62 and Trp63 on one side and Trp108 (Fig 5) on the opposite side of the ligand pocket, two others, Trp28 and Trp111, in the hydrophobic matrix region of the protein, the last Trp123 apart from all the others on the other side of the protein. Concerning the fluorescence properties of the protein, it has been shown that Trp63 and Trp123 do not participate to the total emission of fluorescence; the presence in their neighborhood of disulphide linkages (Cys(76)-Cys(94) and Cys(6)-Cys(127) respectively) induces a quenching of their emission.\(^{28}\) Nishimoto et al.\(^{50,51}\) in a study of two mutants of lysozyme (W62Y and W108Y) where Trp62 and Trp108 are respectively replaced by tyrosine residues, have resolved the emission spectrum of the whole protein in solution. They showed that the emission of the protein can be describe as two regions, one characteristic of a hydrophobic environment at short emission wavelength with Trp28 and Trp111 (partially resolved) in the range 320-330 nm, the other of a polar environment at longer wavelength with Trp62 and Trp108 at 352 and 342 nm respectively. In their study the contributions of Trp28 and Trp111 have been estimated to be 38\% and 23\% of the total emission for the W62Y and W108Y mutants respectively.\(^{50}\) These results imply that at least 70\% of the total emission in the wild type lysozyme is coming from Trp62 and Trp108. This proportion agrees with previous works where a contribution of 80 \% was already estimated.\(^{27}\)

In our experiments, the lysozyme conformation and orientation in solution and in layered materials have been studied by following the time resolved fluorescence and anisotropy of fluorescence of the tryptophan residues. In all our experiments data were collected at 345 nm. This emission wavelength was chosen far from the excitation wavelength in order to avoid problems of scattering light and close enough from the maximum of emission in order to get a high count rate. At 345 nm the contributions of Trp62 and Trp108 are almost equal and can be estimated from Nishimoto et al\(^{50,51}\) to be around 92\% of the total emission of the protein; the other 18\% are contributions of Trp28 and Trp111, which have a maximum emission at longer wavelength in the blue side of the spectrum.
Lysozyme in buffer solution.

A typical experiment of time-resolved fluorescence and time-resolved depolarization of lysozyme in solution is illustrated in Fig. 6. Results of the analysis are summarized in Table 2. Four fluorescence lifetimes has been necessary in order to describe adequately the decays of the lysozyme in solution. This multi-exponential behavior is frequently observed even for proteins with a single tryptophan; this can be explained by the complex photophysical properties of the indole chromophore. Two overlapping transition dipoles $1^{L_a}$ and $1^{L_b}$, nearly perpendicular to each other are involved in the absorption and emission process of the indole chromophore. The possibility of different rotamers in the ground state and in these two nearly degenerate electronic states may explain the multi-exponential behavior of the fluorescence decay.\textsuperscript{29,52,53} During the analysis, the introduction of a fourth component was necessary in order to improve significantly the adjustment of the decays. This calculation has led to the appearance of a short component of around 10 ps. As we used a double monochromator and are 50 nm away from the excitation wavelength, a contribution of scattering light can be rejected. The possibility of an artifact in the convolution process is also rejected since we did not observe similar behavior for other molecules. Although even if this component is at the limit of our resolution, which means that, we cannot resolve it, we think we are still able to evidence it. A similar component with a maximum weight at shorter wavelength has already been observed; it was suggested by the authors to be a contribution of Trp28 and Trp111.\textsuperscript{50} Apart from this short component three other fluorescence lifetimes, 3.79 ns, 1.78 ns and 0.62 ns, were necessary in the analysis. The corresponding average lifetime is estimated to be 2.05 ns. The anisotropy decay of the lysozyme in solution was best analyzed using two correlation times: $\phi_1 = 3.98$ ns and $\phi_2 = 0.15$ ns. These results are in accordance with previous works.\textsuperscript{54,55}

The expression of the anisotropy $r(t)$ can be rewritten as:

$$r(t) = \left[ r_{Trp} e^{-t/\phi_{Trp}} + r_p \right] e^{-t/\phi_p}$$

(eq 7)

where $\phi_p$ and $\phi_{Trp}$ correspond respectively to the correlation times for the global motion of the protein and to the local motion of the fluorophores inside the protein. The long component $\phi_1$, consistent with the rotational correlation time for the whole-body of lysozyme calculated by the Einstein-Stokes relationship, is associated to $\phi_p$ and the short component $\phi_2$ related to the fast depolarization motion $\phi_{Trp}$ of the tryptophans inside the protein. Using $\phi_1$ and assuming a spherical geometry for the protein, the hydrodynamic volume of the lysozyme has been estimated to $1.69 \times 10^{-26}$ m$^3$ (hydrodynamic diameter 3.2 nm), which is in agreement with previously reported values.\textsuperscript{54,55}
The intrinsic anisotropy $r(0)$ is a function of the relative orientations of the absorption and emission transition dipole moments. Its maximum and minimum values are 0.4 and -0.2, which are reached when both absorption and emission transition dipoles are parallel or perpendicular, respectively. The absorption of the tryptophan originates from two transition dipoles, $I_a$ and $I_e$, which are nearly perpendicular to each other with different initial anisotropy (0.33, -0.165 respectively); the resulting global depolarization $r(0)$ being an average value of these two contributions. Recent studies show that ultrafast internal conversion (80-150 fs) occurs between these two electronic states as well as solvation dynamics from the environment (picosecond scale for proteins). Resonance energy transfer has been demonstrated to occur in the same ultrafast range. All these ultrafast phenomena contribute to decrease the observable initial anisotropy $r(0)$ at the available fluorescence timescale. For the lysozyme in solution we found $r(0) = 0.25 (r_1 + r_2)$ which is in agreement with usual values using similar techniques.

**Lysozyme loaded on layered materials**

As for the lysozyme in solution, a four lifetime adjustment was necessary for the layered materials. The analysis for the lysozyme incorporated in Na-Mt and C$_{10}$E$_7$-Mt (Table 2) show a slight decrease with respect to the lysozyme in solution for most lifetimes except the longer one; the average fluorescence lifetime decreases, 1.77 ns and 1.89 ns for Na-Mt and C$_{10}$E$_7$-Mt respectively. On the contrary, lifetimes and average lifetime (2.11 ns) for the SOS-LDH remain almost similar to those of the lysozyme in buffer. The emission spectra of the suspension of all the adsorbed samples have been recorded; they remain identical to the one of the protein in solution (not shown). As the emission spectrum of the fully denatured protein is known to exhibit a red shift of about 14 nm, which is not evidenced for any of our samples, a denaturation of the adsorbed protein is very unlikely. This result is in accordance with the known “hard protein behavior” of the lysozyme. The decrease of the average lifetime for the Na-Mt and C$_{10}$E$_7$-Mt samples is in agreement with a modification of the polarity of the surrounding medium of the tryptophans observed at 345 nm. This can be correlated to the confinement of the protein in the interlayer space of the material as already suggested by the XRD experiments. On the contrary, the unchanged average lifetime for the SOS-LDH suggests a different behavior for this sample. This different behavior was already shown in the XRD experiments where no intercalation was evidenced. Anisotropy decays of lysozyme adsorbed on Na-Mt, C$_{10}$E$_7$-Mt and SOS-LDH are presented in Fig. 7. Data for lysozyme confined in Na-Mt and C$_{10}$E$_7$-Mt can be fitted with two correlation times (Table 2) i) a long one around 40 ns ii) and a shorter one of 0.18 and 0.31 ns, respectively. For the lysozyme loaded on SOS-LDH, only one correlation time of 48 ns was needed. No correlation time around 4 ns, which is characteristic of the free lysozyme in solution, has been revealed by the analysis; this suggests that the adsorption process is probably complete and irreversible for all samples. The uncertainty in the determination of very long correlation times is very high in time resolved anisotropy experiments since the observation time window (<37 ns in these experiments) is limited by the fluorescence lifetime of the sample.
The correlation times around 40 ns are probably under estimated; they are associated to the global rotation of the whole entity “protein-material”.

The second correlation time and its amplitude of depolarization for the lysozyme loaded on Na-Mt (\( \phi_2 = 0.18 \) ns, \( r_2 = 0.02 \)) are very similar to those in the buffer solution (\( \phi_2 = 0.15 \) ns, \( r_2 = 0.02 \)); for the C10E7-Mt sample, results are also very similar, with a slight increase of \( \phi_2 \) (\( \phi_2 = 0.31 \) ns, \( r_2 = 0.02 \)). These results suggest that the observed tryptophan at 345 nm have a similar behavior in lysozyme adsorbed on Na-Mt than in solution; the protein located in the interlayer space is surrounded by a medium of lower polarity and its structure remains mainly unchanged. In the interlayer space, the protein is still hydrated. As the \( ^1L_0 \) transition is more sensitive to solvation than the \( ^1L_4 \) transition, any modification of polarity will induce a modification in the average value of the initial anisotropy \( r_{(0)} \). This effect as well as other ultrafast processes involving the surrounding medium, as mentioned before, could be responsible for the lower value of \( r_{(0)} \approx 0.17 \) observed for the Na-Mt sample. A same behavior is observed for the C10E7-Mt sample where the initial anisotropy \( r_{(0)} \) decreases to 0.16. The data for the lysozyme adsorbed on C10E7-Mt present a slower motion of a similar amplitude (\( \phi_2 = 0.31 \) ns, \( r_2 = 0.02 \)) comparatively to the sample loaded on Na-Mt or in solution. The protein located in the interlayer space is as before surrounded by a medium of lower polarity (the lifetimes are shorter) with a possible steric obstruction due to the presence of nonionic surfactant (C10E3) or a stronger interaction with the surface of the material than in the case of sample Na-Mt. From Nishimoto et al.\(^{30,51} \) it is known that Trp62, in the lysozyme in solution, has a higher degree of freedom and is more exposed to the bulk water than Trp108 (Fig. 5). From this result, the increase of the shorter correlation time \( \phi_2 \) may be interpreted as a decrease of the mobility of Trp62. This behaviour suggests a stronger interaction in the neighbourhood of Trp62 in the C10E7-Mt sample than in the Na-Mt sample. As the results for the Na-Mt and the C10E7-Mt samples in suspension are very similar (lifetimes, yields, average lifetimes, emission spectra) apart for this small difference in the internal correlation time \( \phi_2 \), it is reasonable to think that the structures of the lysozyme in both cases are very similar and that the main difference between the two samples is the position of the protein in the interlayer. These observations are compatible with an orientation “side-on”, which was already suggested by the interlayer space determined in XRD experiments, where the Trp62 is expected to be in a higher interaction with the material in the C10E7-Mt sample. For the Na-Mt sample, where the shorter correlation time is almost identical to the one in solution, which means not perturbed by its environment, an orientation “end-on” is more adequate.

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For lysozyme loaded on SOS-LDH, the lack of the short correlation time suggests a loss of the motion of the tryptophans observed at 345 nm. As both lifetimes (and average lifetime) and emission spectrum are very similar to those of the protein in solution, a denaturation of the protein can probably be rejected. The low value of \( r_{(0)} = 0.12 \) obtained from the anisotropy measurements suggests a strong modification of the neighborhood of the tryptophans. According to the XRD experiment, where it was shown that the lysozyme cannot be intercalated inside the interlayer space, a possible adsorption of the lysozyme on SOS-LDH, with an orientation of the protein compatible with an interaction of the tryptophan with the surface is suggested. Such a model is illustrated in Fig. 3c.

20 **Conclusion**

Lysozyme adsorption was studied on three different layered materials: Na-Mt, C\(_{10}E_{10}\)-Mt and SOS-LDH. For all materials the adsorption process does not induce large structural changes of the protein. It was shown that the protein is confined inside the interlayer space of Na-Mt and C\(_{10}E_{10}\)-Mt with a possible “end-on” and “side-on” orientations, respectively. On the contrary, for SOS-LDH the protein is located on the surface of particles since no cation exchange can occur due to the positive charge of LDH. After dehydration, strong structural changes have only been observed for protein adsorbed on the surface of particles of SOS-LDH.

**Notes and references**

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