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Detrimental impact of silica nanoparticles on the nanomechanical properties of *Escherichia coli*, studied by AFM

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**Abstract**

Despite great innovative and technological promises, nanoparticles (NPs) can ultimately exert an antibacterial activity by affecting the cell envelope integrity. This envelope, by conferring the cell its rigidity and protection, is intimately related to the mechanical behavior of the bacterial surface. Depending on their size, surface chemistry, shape, NPs can induce damages to the cell morphology and structure among others, and are therefore expected to alter the overall mechanical properties of bacteria. Although Atomic Force Microscopy (AFM) stands as a powerful tool to study biological systems, with high resolution and in near physiological environment, it has rarely been applied to investigate at the same time both morphological and mechanical degradations of bacteria upon NPs treatment. Consequently, this study aims at quantifying the impact of the silica NPs (SiO 2 NPs) on the mechanical properties of *E. coli* cells after their exposure, and relating it to their toxic activity under a critical diameter. Cell elasticity was calculated by fitting the force curves with the Hertz model, and was correlated with the morphological study. SiO 2 NPs of 100 nm diameter did not trigger any significant change in the Young modulus of *E. coli*, in agreement with the bacterial intact morphology and membrane structure. On the opposite, the 4 nm diameter SiO 2 NPs did induce a significant decrease in *E. coli* Young modulus, mainly associated with the disorganization of lipopolysaccharides in the outer membrane and the permeation of the underlying peptidoglycan layer. The subsequent toxic behavior of these NPs is finally confirmed by the presence of membrane residues, due to cell lysis, exhibiting typical adhesion features.
1. Introduction

The fast development of nanotechnologies has led to both a growing production and a release of nanoparticles (NPs) into the environment. Due to their unique properties, compared to their bulk counterparts, NPs raised a great interest especially for biomedical applications such as theranostics [1-3]. However, they also raised concerns about their potential adverse effects on the health of living systems [4-10]. Notably, when released in air, water and soil, NPs can interact with bacteria necessary for the ecosystem safety, for instance our intestinal or cutaneous microbiota, and damage their integrity [11,12]. Cellular effects of NPs have shown strong dependence on multiple parameters such as their size, charge, shape and chemical nature [13-17], but also differ depending on cell species and environment [18-20]. Consequently, it is as essential as challenging to explore and to understand NPs/cell interactions to provide a safe and innovative development of nanotechnologies in medical or environmental applications, including alternative to antibiotics to specifically and efficiently target bacteria [21].

Much evidence has indicated that the antibacterial activity of NPs takes place through the disruption of bacterial membranes [22,23], the production of Reactive Oxygen Species (ROS) [24,25] and/or the interference with the bacteria metabolism [26,27]. Up to now, the authors have focused on morphological, viability and/or the interference with the bacteria metabolism [26,27]. Below a critical diameter, shape and chemical nature [13-17], but also differ depend on cell species and environment [18-20]. Consequently, it is as essential as challenging to explore and to understand NPs/cell interactions to provide a safe and innovative development of nanotechnologies in medical or environmental applications, including alternative to antibiotics to specifically and efficiently target bacteria [21].

2. Materials and methods

2.1. Materials

Commercial SiO₂ NPs 100 nm and SiO₂ NPs 4 nm were purchased from Biovalley (Marne la Vallée, France) and Alfa Aesar (Schiltigheim, France) respectively. Transmission Electronic Microscopy (TEM) and Diffusion Light Scattering (DLS) measurements were performed to characterize these NPs. Though SiO₂ NPs 4 nm were highly aggregated on TEM cliches, an average hydrodynamic diameter of 3 nm was determined by DLS. For the SiO₂ NPs 100 nm, both technics were in agreement: statistics performed on TEM cliches gave an average diameter of 113 ± 11 nm while the hydrodynamic one was found around 124 nm.

Gram negative E. coli bacteria (MRE 162 strain) were a kind gift from the Centre d'Etudes du Bouchet, DGA (Direction Générale de l’Armement, France). Mica, purchased from Electron Microscopy Sciences (Hatfield, United States), was used as substrate for all AFM measurements. Luria Broth (LB) medium was used as nutritive medium and purchased from Fisher Scientific (Waltham, Massachusetts, United States). Millipore ultrapure water (pH 5.5, 1-2 U.}

help identifying the mechanism by which the particles interact with the bacterial envelope and exert their toxicity [45,46]. In the past decades, atomic force microscopy (AFM) has emerged as a powerful and suited tool to investigate such interactions and to solve many biological issues [45,47-49]. Indeed, not only AFM enables to image, in real time, biological systems with high resolution (10-100 nm scale range) both in air and in physiological environment, but also, with a force spectroscopy mode, it can probe simultaneously their mechanical properties at a molecular level [50-52].

In this study, our purpose was to investigate and probe the mechanical properties of E. coli, at the submicrometric scale, before and after exposure to silica NPs (SiO₂ NPs) with different sizes. On one hand, E. coli cells were chosen because they are widely used systems, either harmless or responsible for serious human diseases. Their cell envelope consists of two lipidic membranes separated by a thin peptidoglycan layer called the cell wall [53]. The outer membrane (OM), which constitutes the interface with the external environment, contains LipoPolySaccharides (LPS) forming a brush like structure at the cell surface. The inner membrane separates a gel like periplasm containing the peptidoglycan from the cytoplasm. In addition, they are ubiquitous in the environment and thus, highly exposed to interactions with NPs [54]. On the other hand, SiO₂ NPs are considered as promising candidates for biomedical imaging, drug delivery, and biosensing applications [55]. Furthermore, despite their easy control in terms of size and surface chemistry, the antibacterial activity of SiO₂ NPs is not well documented and their mechanism of action is still unknown [56-58]. For such NPs, a toxicity towards E. coli was shown to depend on their charge and size [17]. Below a critical diameter \( \Phi_{c} (50\text{ nm} < \Phi_{c} < 80\text{ nm}) \), morphological and membrane structural damages eventually led to cell lysis. To better understand the antibacterial effect of SiO₂ NPs, AFM was here used mainly in force spectroscopy mode and in liquid to measure the Young modulus of E. coli cells, after their treatment with NPs with diameter of either 4 nm or 100 nm (called later SiO₂ NPs 4 nm and SiO₂ NPs 100 nm, respectively). This work aims at quantifying the NPs/bacteria interaction based on the change in the mechanical properties of E. coli upon exposure to NPs. AFM force spectroscopy provides such a quantification, as well as a correlation with the morphological and membrane characteristics of E. coli, obtained with AFM imaging either in air or in liquid, in tapping and contact modes respectively.
resistivity > 18.2 MΩcm) was used for NPs cleaning and bacteria suspensions preparation. Polyelectrolytes, used in this work to immobilize bacteria on mica, are cationic Poly (Allylamine Hydrochloride) (PAH, MW = 56000, pure at more than 95%), and anionic Poly (sodium 4 Styrene Sulfonate) (PSS, MW = 70000, conform infrared spectrum); they were purchased from Sigma Aldrich.

2.2. Bacterial culture and interaction with NPs

_E. coli_ cells were grown on LB agar plates and incubated for 16 h at 37 °C. After scraping, bacteria were suspended in ultrapure water and their concentration was adjusted to 8.10^6 cells/mL, corresponding to an optical density (at 600 nm) of 1.0. Consequently, the studied bacteria are in the stationary phase of their growth. Though many experiments required buffer solution (like Phosphate Buffer Solution, PBS), we chose ultrapure water for convenience reasons (to prevent the formation of PBS crystals for AFM characterizations in air, after the drying step) and because no bacterial mortality was observed in such simple solvent after 2 h.

Then, SiO_2_ NPs (also dispersed in ultrapure water) were introduced in the bacteria suspension at a final concentration of 1 g/L. The co-suspension was mixed and gently manually shaken for homogeneity (4 times every 20 min), then left to interact at room temperature for 2 h. Five μL of the co-suspension were deposited on either bare mica, for AFM observations in air, or on a PolyElectrolyte Multilayer (PEM, see below), for AFM observations in liquid.

2.3. Substrate and sample preparations for AFM observation

With an average roughness of 0.3 nm/μm², mica is perfectly suited for AFM imaging to discriminate any topographic contrast. Bare mica was, then, used in this work for all experiments carried out in air, mainly AFM morphological characterizations. If one wants to image bacteria in a more realistic environment such as a liquid, the sample needs to be fixed on the substrate first, to prevent any motion with the fluid flow. To do so, we successfully applied a sequential deposition of oppositely charged solutions of polyelectrolytes (PAH and PSS) [59]. Polyelectrolytes were dissolved in ultrapure water at a concentration of 0.5 g/L and alternatively deposited through a spin coating process (45 s, 100 rpm, 200 s⁻¹), starting with PAH on bare mica. Consequently, a (PAH PSS)₅ PAH PEM was formed to immobilize the negatively charged _E. coli_ bacteria. This PEM exhibits an average thickness of 4 nm, determined with scratching AFM experiments. Whatever the substrate (bare or functionalized mica), five μL of bacteria suspension, untreated or treated with NPs, were deposited on its surface before imaging.

2.4. AFM observations

AFM experiments were carried out using a Bioscope II operating with the NanoScope V controller. For morphological characterizations, we used the tapping mode in air and in the repulsive domain, in order to prevent any damage to the sample. Images were obtained with commercial cantilevers with a spring constant around 40 N/m, at a scan rate of 0.5–1.0 Hz and with a resolution of 512 × 512 pixels. For imaging in liquid (ultrapure water), we used the contact mode, with an applied force around a few nN to prevent any damage or indentation of the sample. Data were obtained with commercial cantilevers with a spring constant of 0.12–0.20 N/m, at a scan rate of 1.0 Hz. Different areas were systematically imaged and results shown thereafter are representative of the cells in a given set of colonies and of all the tested samples.

When imaging in air, the sample was first left to dry in a desiccator and imaged on the day following the preparation. On the other hand, when imaged in liquid, the sample was prepared and directly imaged.

2.5. AFM force distance curves (FCs) acquisition

In the force spectroscopy mode, the AFM tip is considered as a sensitive sensor, being pressed on the sample surface. Indeed, it is possible thanks to the cantilever to control and measure the applied force and to couple AFM imaging with the mechanical response at the local scale of the studied system [60]. During a FC acquisition, the tip is approached and then retracted from the surface: the former part of the FC can be used to measure the indentation of the sample and its stiffness, while the latter one gives information on the tip sample interactions, like adhesion. For FC measurements, triangular silicon nitride tips, with a spring constant of 0.12–0.20 N/m determined by the thermal noise method, were used. To obtain deflection sensitivity, calibrations were performed in air and then in ultrapure water. FCs were recorded in the central region of different cells, from highly aggregated ones, to avoid any artifact related to edge curvature. To statistically provide robust data, at least three independent experimental sets (bacterial samples from different cultures) were investigated, either treated with NPs or not, and in each set at least 10 different bacteria were studied in different areas of the sample.

After FC acquisition, quantitative measurements of bacterial elasticity can be obtained through three distinct theories: (i) the Hertz model can be applied when adhesion is negligible compared to the load, (ii) the Johnson Kendall Roberts (JKR) model can be applied for large tips and soft samples with large adhesion and (iii) the Derjaguin Muller Toporov (DMT) model can be applied for small tips on stiff material with low adhesion [50]. Despite its limitations, especially concerning the uncertainty in the contact point determination and the assumed homogeneity and linear elasticity of the sample [61], the Hertz model has been one of the simplest and most widely used model to quantify the mechanical properties of biological samples [48]. Consequently, we chose this model to fit our FCs. It gives access to the Young modulus, _E_, of the sample when this last one is indented by a tip with a known geometry (here conical) [62]:

\[
F = \frac{2Etan(\theta)}{\pi}\delta^2
\]

where _F_ if the applied force, _θ_ is the half cone angle of the tip (set to 25°), _U_ is the Poisson ratio (assumed to be 0.5 for soft materials) and _δ_ is the deformation of the cell due to the tip. The Young modulus is also linked to the stiffness of the sample, _k_s_, by the relation:

\[
k_s = \frac{4}{\pi}\delta tan(\theta)
\]

In practice, we obtained curves of the deflection, _d_, according to the tip to sample distance, _z_. In parallel, we developed a Matlab routine to transduce these curves into those expressing the indentation force, _F_, as a function of the deformation, _δ_. These two parameters were calculated according to the following relationships [62]:

\[
\begin{cases}
0 & z \leq z_0 \\
(z-z_0) & \delta > z_0
\end{cases}
\]

\[
F = k(d-d_0)
\]

where (_z_0, _d_0) are the coordinates of the contact point and _k_ is the cantilever stiffness.

Once the curve _F = f (δ)_ was obtained, the Young’s modulus of the sample was calculated by adjusting this curve with the above
formula. Generally, it is estimated that beyond 10% thickness of the cell, the measurement is biased by the intracellular part, which might be modified by the indentation, and potentially the substrate properties [62]. In our case, this thickness corresponds to the height of the bacteria with a maximum value around 700 nm (see below). Consequently, the adjustments made during this work were performed on a maximum distance of 70 nm, though it may also be suitable on a longer distance for some samples. This also limits the effect of the Hertz model assumptions concerning a purely homogeneous, linearly elastic material and an infinite sample thickness.

2.6. Statistical description of AFM data

To determine if statistical variations were present between untreated and treated bacteria (either with SiO$_2$ NPs 100 nm or SiO$_2$ NPs 4 nm), the Mood’s median test, which establishes the equality between medians of independent samples, has been applied to our data. This test was chosen because it is insensitive to exceptional values and does not depend on the distribution shape and variance. If $X^2 > X^2_{a,(n-1)}$ then the null hypothesis is rejected: the n samples are significantly different, at a significant level of $\alpha%$. In our case, $\alpha = 5%$ and $n = 2$. In addition, the Shapiro Wilk test was performed to assess the matching between the Young modulus distributions of treated and untreated bacteria with a normal distribution. Such a test may also be represented with a Q Q plot. The test is significant if the null hypothesis is rejected, $p < \alpha = 5%$, which means data do not follow a normal distribution. Oppositely, if the null hypothesis is accepted, $p > \alpha = 5%$, it does not mean the data follow a normal distribution. Then, to characterize in more details the potential differences obtained on the Young modulus distribution of untreated and treated bacteria, we calculated the skewness, $b_1$, which allows to determine the importance of the asymmetry in a distribution:

$$b_1 = \frac{m_3}{s^3}$$

where $m_3$ is the sample third central moment and $s$ is the sample standard deviation.

If $b_1 = 0$ the distribution is symmetric. If $b_1 > 0$, the distribution is left dissymmetric whereas if $b_1 < 0$, the distribution is right dissymmetric.

3. Results

3.1. Untreated E. coli: Soft rod like cells

Fixed on a PEM and imaged in ultrapure water, E. coli cells form films, more or less dense depending on the rinsing step (Fig. 1a and b). They have the typical well known rod shape, previously observed in air and liquid [63] (Fig. 1c and d). Their length and width are similar to those estimated in air, around 2 $\mu$m and 1 $\mu$m respectively. Because of their hydrated state, the only difference lies in their height: 700 nm at maximum, corresponding to about twice as much as the estimated height for E. coli observed in air. Such characteristics, in agreement with previous studies [63], are representative of healthy bacteria. Noteworthy is the more homogeneous aspect (still rough and irregular) of the E. coli surface in liquid (Fig. 1c) in comparison to the ripples structure, observed in air and characterized by a specific relief associated to the presence of LPS in its OM (Fig. 1d) [59]. This phenomenon is likely due to the hydration state of the OM allowing a more or less significant extension of LPS, even if a screening effect due to the liquid environment cannot be excluded.

To achieve the mechanical study of E. coli surface, FCs were performed on highly aggregated cells as well as more isolated ones. As expected, during the approach, FCs performed on the central part of E. coli cells exhibit the specific features of those obtained with...
a soft sample (Fig. 2a): a baseline (I) in the absence of interaction, a linear part (III) following the repulsive interactions between the tip and the sample and, in between, a nonlinear zone (II) due to the indentation of the sample. Such a response is very different from the substrate one. Indeed, due to the very small thickness of the PEM (average thickness of 4 nm), the FC obtained on this layer exhibit the typical behavior of stiff material (abrupt increase in the deflection signal after contact) such as the mica substrate underneath: no indentation is observed at the observation scale (Fig. 2a and b). To account for any artifact due to the curvature of *E. coli*, we also performed FCs in various areas of the same cell (data not shown). Qualitatively, results show similar features, as described above, implying a global homogeneous soft structure of the bacteria, regardless of the scanned zone. Discrepancies between bacteria are also observed, in the non linear and linear parts, confirming other studies based on force mapping of bacteria [41,64]. They are likely due to different molecular organizations and/or compositions leading to stiffer and softer domains present in the cell wall, which involve different mechanical responses between peripheral and apical regions [65,66]. These observations justify our choice to perform FCs only in the central region of the cell; this also prevents any artifact due to edge effect of both cell and tip. Among various experimental sets of *E. coli* and exploring different zones (highly aggregated cells to more isolated ones), bacteria showed an overall homogeneity in their response (Fig. 2c).

Quantitatively, we calculated the Young modulus of about 30 bacteria over 4 experimental sets (i.e. independent samples from different bacterial cultures), by adjusting their FC with the Hertz model, according to the procedure described in the Materials and Methods. In one experimental set, the Young modulus can vary by one order of magnitude. The apparently slight discrepancies observed on the curves turn out to hide a relatively high dispersity in the elasticity of the cells (Fig. 2d). Such a heterogeneity is also highlighted among the different experimental sets of bacteria. These results reflect the limitations associated with live cell experiments, where the intrinsic mechanical behavior is potentially influenced by the physiological state of cells [64,67] or their adhesion to the substrate [68,69]. A median value for the Young modulus of *E. coli* was estimated to be 1.2 MPa, corresponding to a stiffness of 0.05 N/m. The interquartile range (1st quartile/3rd quartile) is [0.5, 2.3 MPa] (Fig. 2d inset).

This is in good agreement with previous studies on *E. coli* (Table 1). The observed differences can be ascribed to the diverse methods used in immobilizing bacteria that lead to different adhesion forces between cells and substrate [70] or to different solvents and therefore different osmotic and turgor pressures: the higher the turgor pressure the higher the stiffness [39,71].

3.2. Morphology of *E. coli* treated with *SiO<sub>2</sub>* NPs

In a previous work, we showed the existence of a critical diameter, below which *SiO<sub>2</sub>* NPs are toxic for *E. coli*: 50 nm < Φc < 80 nm [17]. Above this diameter, NPs do not change the overall morphology of *E. coli* or its membrane structure. On the opposite, below this diameter, NPs induce (i) a transition from a rod to a more spherical shape (Fig. 3a), (ii) the formation of pore like lesions/membrane invaginations (arrows in Fig. 3a) and (iii) the reorganization of the LPS molecules of the OM of *E. coli* from...
extended ripples to more aggregated ones (Fig. 3b to be compared to Fig. 1d). Such effects seem to illustrate the disruption of E. coli OM through breaks into the LPS organization and the destructuration of its peptidoglycan layer involved in the cell shape. Eventually, cell lysis and subsequent leakage of intracellular components might occur (Fig. 3c).

3.3. Change in E. coli elasticity after interaction with SiO₂ NPs

Taking into account the existence of this NP critical diameter with respect to NPs antibacterial activity, we will focus here on the effect of two SiO₂ NPs with diameters well below and well above \( U_c \), on the bacterial membrane properties, at the submicrometric scale.

\( U_c > U_c \): case of SiO₂-NPs 100 nm – Fig. 4a shows typical FCs obtained on E. coli cells treated by SiO₂ NPs 100 nm. In most cases, FCs exhibit the same features as those obtained with untreated E. coli. The non linear transition which is due to the membrane elasticity can exhibit a more or less important width. At the same time, the linear part presents a more or less steep slope. However, FCs obtained following such treatment by SiO₂ NPs 100 nm are often included in the area of the heterogeneous responses of untreated bacteria (Fig. 4a dashed area).

Like in the case of untreated E. coli, dispersity is observed in the Young modulus of treated cells (Fig. 4b). The estimated median elastic modulus of 1.4 MPa, corresponding to a stiffness of 0.06 N/m, suggests that no significant difference exists between these treated E. coli and the healthy ones (Mood’s test: \( \chi^2 \leq \chi^2_{0.05, 1} \)). Nevertheless, the interquartile range \((Q1 1st quartile/Q3 3rd quartile)\) of \([1.1, 3.3 \text{ MPa}]\) shows that there is a slight shift towards a higher value of the Young modulus.

\( U_c < U_c \): case of SiO₂-NPs 4 nm – Treatment of E. coli cells using SiO₂ NPs 4 nm eventually triggers the morphological transition from a rod shape to a spherical one (Figs. 3a and 5a). Such a treatment also induces the formation of an unusual homogeneous and porous film on the substrate (Fig. 5a arrows), which will be described afterwards.

On Fig. 5a and b, bacterium (1) shows a height (530 nm) similar to those observed in healthy bacteria though it exhibits a spherical shape. In contrast, bacterium (2) has a lower height section (270 nm), suggesting its collapse, while it exhibits the expected rod shape. Consequently, treated bacteria can not only exhibit a different shape but also a different height, characteristics that are not necessarily correlated for all cells. For both bacteria FCs have features mainly typical of those obtained with a soft sample with different indentations (Fig. 5c). More specifically, the FC on the “spherical” bacterium (1) indicates a higher rigidity than that of

### Table 1

| Fixation          | Solvent     | \( k_S \) (N/m) | Ref. |
|-------------------|-------------|-----------------|------|
| Polyelectrolyte   | Water       | 0.05 ± 0.01     | This work |
| Polys-1-lysine    | Water       | 0.04 ± 0.01     | [36] |
| Glutaraldehyde    | Buffer      | 0.194           | [72] |
| Glutaraldehyde    | Buffer      | 0.23 ± 0.01     | [73] |
| EDC/NHS           | Buffer      | 0.10 ± 0.01     | [73] |
| EDC/NHS           | Tris buffer | 0.044 ± 0.014   | [74] |
| Glutaraldehyde    | Tris buffer | 0.13 ± 0.01     | [74] |

Fig. 3. AFM images, obtained in air, illustrating the antibacterial activity of SiO₂-NPs 4 nm towards E. coli. (a) 3D topography image showing the transition from a rod to a more spherical cell (compare with Fig. 1d) and the invaginations (red arrows). (b) Phase image emphasizing the reorganization of LPS molecules in the OM (compare with Fig. 1d). (c) Amplitude image illustrating the lysis of E. coli with the presence of debris (arrow) and empty cells. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
untreated bacteria, with an elastic modulus of 7 MPa. On the other hand, the FC on the collapsed bacterium (2) exhibits a softer behavior with a Young modulus of 0.4 MPa, which is in favor of an alteration of its membrane integrity.

Statistics performed on other bacteria exhibiting similar FCs show that, again, a large dispersity is obtained for *E. coli* elasticity (Fig. 5d). The estimated median elastic modulus of 0.5 MPa, corresponding to a stiffness of 0.02 N/m, suggests that SiO$_2$ NPs 4 nm induce a significant increase in the elasticity of bacteria (Mood’s test: $X^2 \geq X^2_{0.05}$). Also, the interquartile range (Q1-Q3) of [0.4, 1.6 MPa] shows that there is a shift towards the lowest values of the Young modulus, while the extreme values (minimal and maximal) are higher by one order of magnitude than in the case of untreated cells.

![Fig. 4.](image1.png) (a) FCs in approach representative of the different cases obtained following the treatment of *E. coli* by SiO$_2$-NPs 100 nm. The dashed area represents the heterogeneous responses of untreated *E. coli*. (b) Box chart representation of the Young modulus (E) of *E. coli* bacteria untreated and treated with SiO$_2$-NPs 100 nm. Percentiles are indicated in the case of *E. coli* and are the same for treated cells.

![Fig. 5.](image2.png) (a) AFM height image, obtained in ultrapure water, of *E. coli* cells treated with SiO$_2$-NPs 4 nm. (b) Height profiles of the two bacteria imaged in (a). (c) FCs in approach realized on the same two bacteria. The dashed area represents the heterogeneous responses of untreated *E. coli*. (d) Box chart representation of the Young modulus E of *E. coli* bacteria untreated and treated with SiO$_2$-NPs 4 nm. Percentiles are indicated in the case of *E. coli* and are the same for treated cells.
Moreover, in a few cases, FCs exhibited a three-stage trend: (i) a non-linear zone characteristic of a soft sample, (ii) a plateau over a few tens of nanometers where the deflection is null and the force constant, and (iii) a linear part (Fig. 6). This particular FC is associated with a collapsed bacterium, similar to bacterium (2) in Fig. 5. As untreated bacteria did not show such a behavior, it could be explained by the effect of SiO2 NPs 4 nm on the organization of the bacterial envelope and its subsequent degradation. A hypothetical scheme, explaining such a FC, is presented thereafter. First (A), the tip indents the bacterium surface with an elastic behavior. Only an elastic deformation occurs, without any perforation. At some point (B), it encounters a structure/defect within the bacterial envelope where it slides and penetrates without any resistance, resulting in a null deflection. Finally, once this damaged structure is crossed, the tip probes again the local elastic properties of the bacterium (C) coupled with the properties of the substrate (D).

This particular profile highly suggests that SiO2 NPs 4 nm induce a structural modification of the bacterial envelope, or even of the plasma membrane, and supports a transition in the organization of LPS and membrane invaginations already observed in air (Fig. 3). Furthermore, the higher slope of the linear part (D) is also in favor of cell flattening and degradation, as the tip responds to a greater extent and earlier to the substrate than for healthy untreated E. coli.

Finally, the treatment of E. coli with SiO2 NPs 4 nm also induces on the substrate the formation of an unusual homogeneous and porous film, called bacterial “fingerprints” later on, besides the expected dense film (Fig. 7a). These “fingerprints” lower than the PEM substrate are encircled by an accumulation of materials (arrows in Fig. 7a) with thickness from tens to about one hundred nanometers (Fig. 7b). Given the height profiles of the borders, it is unlikely that this material is made of NPs. Also, as the entire area is not covered by this accumulation, polyelectrolytes are probably not involved since they rather should be homogeneously distributed on the substrate due to the spin coating deposition. In

![Fig. 6. Comparison of FCs performed on healthy E. coli and on those treated with SiO2-NPs 4 nm bearing a degraded envelope.](image)

![Fig. 7. (a) AFM topography image, obtained in ultrapure water, of E. coli cells treated with SiO2-NPs 4 nm showing a zone devoid of healthy cells, where only “fingerprints” remain. (b) Height profile of the “fingerprint” indicated in (a) (dashed line) showing an accumulation of materials at its edges (white arrows). At the bottom, a scheme illustrating the height image and profile. The grey area represents the PEM substrate, the blue area the aggregates at the edges and the red area the debris present in the fingerprint. c) Representative FCs in retract performed on these “fingerprints”. (d) Schemes explaining the adhesion jumps observed on (c) by the presence of bacterial residues within the “fingerprints”. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)](image)
addition, areas inside the “fingerprints”, close by and far from the edges show different interactions with the AFM tip, as emphasized by friction images (data not shown). Consequently, these three areas (Fig. 7a and b) are probably made of diverse materials organized differently explaining different mechanical properties.

FCs realized on the bacterial “fingerprints” exhibit different types of behavior with typical ones presented in Fig. 7c: (i) a baseline (almost zero deflection) followed by a purely linear part with a slope close to 1, as for the PEM film (grey curve). This baseline shows many discrepancies: contrary to the PEM substrate, “jumps” in force are observed, which are more or less wide and numerous (red curve). These phenomena reveal the presence of a structure generating adhesion events attracting the tip to the sample. Such an adhesion could be due to the presence of crosslinked cell debris within these “fingerprints”; these debris act like “glue” by attaching to the tip and induce a negative cantilever deflection despite the sample removal (Fig. 7d). Combined with the morphological study, these results confirm the toxicity of SiO$_2$ NPs 4 nm towards E. coli and strongly argue in favor of their potential bacteriolytic activity towards some cells in the bacterial population.

Though they deeply illustrate and enlighten the interaction between E. coli and SiO$_2$ NPs 4 nm, FCs showing uncommon features (different from those of soft samples) (Figs. 6 and 7) could not be processed by the Matlab routine and, as a consequence, the Young modulus was not calculated in these cases.

3.4. Comparative analysis

The set of results obtained on E. coli exposed or not to SiO$_2$ NPs is shown in Fig. 8a. These histograms allow an easier overview and comparison between the elasticity of the untreated and treated E. coli, than the unique values of medians reported above.

As seen above, whatever the treatment, the distribution of the Young modulus (in log) is relatively wide, probably due to the intrinsic inhomogeneity of the cellular population. Indeed, cells are probably not in the same growth stage as bacteria are not synchronously dividing. Nonetheless, depending on the NPs treatment, different shapes in the Young modulus distributions can be observed, which are compared to a normal distribution in Fig. 8b. As long as the treatment by SiO$_2$ NPs 100 nm is concerned, according to the Shapiro Wilk test, both untreated and treated E. coli might follow a normal distribution. Another way to characterize such distributions is the Q Q plot, which is a graphical method to compare distributions by plotting their quantiles against each other (Fig. 8b). In each case, the Q Q plot does not follow the 45° line y = x (slight inflection points are observed), suggesting that the distributions obtained for E. coli and E. coli treated with SiO$_2$ NPs 100 nm do not actually follow the normal distribution. Indeed, the calculated skewness of 0.5 and 0.4 respectively, confirm that these distributions are not symmetric and, consequently, are not equal to a normal distribution. In addition, the distribution obtained on E. coli treated with SiO$_2$ NPs 100 nm con firms the shift towards the highest Young modulus, previously observed on the box chart (Fig. 4b). Bins of low Young modulus values are depopulated (Q1 = 0.5 MPa for untreated cells vs Q1 = 1.1 MPa for treated cells) while bins of high Young modulus are equally populated (Q3 = 2.3 MPa for untreated cells vs Q3 = 3.3 MPa for treated cells). Despite this shift, SiO$_2$ NPs 100 nm do not seem to have a major impact on the stiffness of E. coli (equivalent median), in agreement with their harmless effect on the cell morphology and membrane structure.

As long as the treatment by SiO$_2$ NPs 4 nm is concerned, the Shapiro Wilk test and the Q Q plot both show that the distribution is far from following a normal distribution. A skewness as high as 1.5 confirms a significant shift towards the lowest value of the Young modulus, as observed in the box chart (Fig. 5d). Then, such a treatment induces the opposite effect of that observed for SiO$_2$ NPs 100 nm (Q1 = 0.4 MPa, Q3 = 1.6 MPa). SiO$_2$ NPs 4 nm induced a significant decrease in cell elasticity, in good agreement with the morphological and membrane damage previously shown [17].

4. Discussion

Our study aims are not only to correlate the morphological changes observed after NPs treatment to the mechanical properties...
of these structures, but also to quantify the mechanical interactions involved in these processes. The case of SiO$_2$ NPs 100 nm will not be discussed thereafter because no significant modifications, neither on the morphology nor on the elasticity of cells, were observed, implying their harmless behavior towards *E. coli*.

We will then focus on the case of SiO$_2$ NPs 4 nm treatment. Among the previous studies involving mechanical properties of bacteria, most of them were dealing with the effect of antibacterial compounds, and provided elements to emit hypotheses on the bacterial tercidal action of SiO$_2$ NPs 4 nm. First, a significant decrease in the Young modulus of *E. coli* has been shown. Such a decrease was also reported upon treatment with chitosan and lysostaphin (enzyme degrading peptidoglycan) on *Staphylococcus aureus* [75,76]. In this study, the decrease in stiffness was correlated with a global weakening of the cell wall, potentially accompanied by cell lysis. Despite a different mechanism of action, β-lactam and aminoglycoside also led to the disruption of the peptidoglycan layer and a decrease of the Young modulus of *Pseudomonas aeruginosa* cells [77]. Other detailed studies on the action of antimicrobial peptides also correlated such a decrease in cell elasticity with cell envelope modifications associated with higher surface roughness, cell flattening and pore formation [78,80]. Indeed, Quiles *et al.* made the link between the decrease in *Pseudomonas fluorescens* elasticity and the permeabilization of the bacterial membrane after the action of a linear cationic peptide [80]. All these cases are relevant to organic and biochemical molecules, and to our knowledge it is the first time that such a quantitative study on bacteria exposed to such small size metal oxide NPs is undertaken. In a previous paper silver NPs were twice as big [81]. We also outline and quantify in this work, the similar damages induced by tiny nanoparticle cles when interacting with bacteria. Indeed, in the case of SiO$_2$ NPs 4 nm, it is reasonable to assume that the decrease in Young modulus in *E. coli* is associated to the modifications of, at least, the envelope structure because the NPs are small enough to interact deeply with the outer membrane.

Moreover, these NPs also generate important morphological and membrane damages (Figs. 3 and 6). Cell lysis is further emphasized by the residual structures, “fingerprints” that have similar width and length as untreated bacteria and exhibit edges of tens to hundreds of nanometers in height, probably bacterial residues. To our knowledge, such “fingerprints” have been reported only a couple of times, notably in the study of the antimicrobial peptide PGa action on *E. coli* [78] and the work on *E. coli* incubation in egg white at bactericidal temperature [82]. Adhesion events observed in these “fingerprints” show evidence of lysis debris still attached to the substrate resulting from the rupture and the loss of cytoplasmic materials. Previous studies on the interaction between diverse biolayers and substrates reported such adhesion events on FCS: desorption forces were estimated around 40 200 pN for polyelectrolytes depending on their chain length and concentration [83], and between 10 and 100 pN for peptides depending on the substrate functionalization [84]. In our case, we estimated a desorption force around 500 pN to 1 nN. These high values are likely the result of a “mixture” of complex composition including polyelectrolytes, lipids and proteins of bacterial membranes. In particular, the diversity of lipids in terms of chain lengths and polar head group composition probably explains the higher desorption forces determined here. Finally, the very rare studies exploring the elastic modification of bacteria following their treatment with bactericidal silver NPs also underlined a decrease in the Young modulus caused by extensive damage to the cell membrane (composition and structure) [81].

In summary, the previous studies on bio organic molecule impacts and our results strengthen the idea that SiO$_2$ NPs 4 nm, as organic species, probably interact with *E. coli* according to the following mechanism. First, likely due to their small size and chemical nature, they induce a disorganization in the close packing of the LPS molecules, and change the surface structure of *E. coli* (Fig. 3b). Doing so, they induce the formation of pore like lesions in the OM (Figs. 3a and 6), which is then permeabilized and the peptidoglycan potentially compromised. This probably induces a transition from a rod to a spherical shape (Fig. 5a) and an increase in the cell wall elasticity (Fig. 5d). Finally, as a direct or indirect consequence of their presence, a cell lysis occurs and cytoplasmic elements are released (Figs. 3c and 7). Though this hypothetical mechanism is proposed, many questions remain and still need to be addressed, especially in the chronology of the observed damages. In addition, heterogeneous responses were observed after treatment with SiO$_2$ NPs 4 nm, among others high values of the Young modulus. They could find their origin in the potential change of the cell shape without being associated to the cell flattening, leading to higher surface tension.

5. Conclusion

Many studies have focused on the well known antibacterial activity of metal oxide and silver NPs, coping with the charge and size dependency of their interactions without addressing their impact on the mechanical properties of cells [16,28,81]. In our previous work, we had shown the existence of a critical diameter (50 nm < Φ < 80 nm) below which SiO$_2$ NPs induce morphological and membrane structural damages eventually leading to the *E. coli* cell lysis [17]. Here, by combining AFM imaging with spectroscopy mode, we also quantitatively demonstrate that such a toxicity is statistically correlated with modifications in the *E. coli* elasticity. Indeed, as opposed to SiO$_2$ NPs 100 nm which do not have a major impact on the cell elasticity, the SiO$_2$ NPs 4 nm do induce a significant decrease in the cell Young modulus. This, along with the morphological study, provides clues to the SiO$_2$ NPs potential bactericidal mechanism of action: by first damaging the bacterial OM (LPS reorganization, invaginations of the envelope), such small NPs induce the deconstruction of the peptidoglycan layer and, subsequently, lead to the cell lysis. Such a statistical analysis and a mechanical quantification of the NPs/bacteria interaction has only been reported once, for metallic silver NPs twice as big as those reported here [81]. In the case reported here, it should be mentioned that the study only concerns the mechanical cell/NP interaction, since no leaching of any kind is supposed to take place as in the case of Ag$^+$ for silver nanoparticles.

Up to now, interactions between bacteria and SiO$_2$ NPs have not been well documented although such NPs are considered as promising tools for diverse biomedical applications, including antibacterial agent, because of their harmless behavior towards eucaryotic cells and their easy surface modifications [21,55]. Consequently, we have, here, addressed the intrinsic effects of SiO$_2$ NPs on *E. coli* membrane, from both morphological and mechanical points of view. Other parameters, like the charge and shape of NPs, play a critical role in their deleterious impact on cells [13]. They should now be investigated in light of the great promise offered by the combination of AFM imaging and spectroscopy modes, both in air and in liquid.

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