

Osmotic stress and vesiculation as key mechanisms controlling bacterial sensitivity and resistance to TiO₂ nanoparticles

Christophe Pagnout¹, Angelina Razafitianamaharavo^{2,3}, Bénédicte Sohm^{1,3}, Céline Caillet^{2,3}, Audrey Beaussart ², Eva Delatour¹, Isabelle Bihannic ², Marc Offroy ² & Jérôme F. L. Duval ² 

Toxicity mechanisms of metal oxide nanoparticles towards bacteria and underlying roles of membrane composition are still debated. Herein, the response of lipopolysaccharide-truncated *Escherichia coli* K12 mutants to TiO₂ nanoparticles (TiO₂NPs, exposure in dark) is addressed at the molecular, single cell, and population levels by transcriptomics, fluorescence assays, cell nanomechanics and electrohydrodynamics. We show that outer core-free lipopolysaccharides featuring intact inner core increase cell sensitivity to TiO₂NPs. TiO₂NPs operate as membrane strippers, which induce osmotic stress, inactivate cell osmoregulation and initiate lipid peroxidation, which ultimately leads to genesis of membrane vesicles. In itself, truncation of lipopolysaccharide inner core triggers membrane permeabilization/depolarization, lipid peroxidation and hypervesiculation. In turn, it favors the regulation of TiO₂NP-mediated changes in cell Turgor stress and leads to efficient vesicle-facilitated release of damaged membrane components. Remarkably, vesicles further act as electrostatic baits for TiO₂NPs, thereby mitigating TiO₂NPs toxicity. Altogether, we highlight antagonistic lipopolysaccharide-dependent bacterial responses to nanoparticles and we show that the destabilized membrane can generate unexpected resistance phenotype.

¹Université de Lorraine, CNRS, LIEC, Metz, France. ²Université de Lorraine, CNRS, LIEC, Nancy, France. ³These authors contributed equally: Angelina Razafitianamaharavo, Bénédicte Sohm, Céline Caillet. ✉email: jerome.duval@univ-lorraine.fr

Due to their photocatalytic properties, titanium dioxide nanoparticles (TiO₂NPs) are among the NPs that are most produced and used in consumer products¹. The antibacterial activity of TiO₂NPs and of related composite nanomaterials² towards microorganisms has been largely evaluated at the cell population level using a battery of dose–response relationships^{3–7}. This approach, though important for toxicological risk assessment, remains insufficient on its own for addressing NP–cell interactions and the biological implications thereof at a mechanistic nanolevel^{8,9}. In this regard, the development of omics¹⁰ and fluorescence-based bioassays¹¹ has increased our understanding of the molecular processes that underpin the manifestation of adverse effects of TiO₂NPs^{4–6,12–29} and other photocatalytic nanomaterials^{4,12,23,24,28–34} on bacteria. These advances have contributed to unraveling how cell exposure conditions and physicochemical properties of TiO₂NPs determine the toxicity^{4–6,13–29}. In particular, the generation of reactive oxygen species (ROS) by TiO₂NP photocatalysis¹² is commonly described as a key process that leads to cell surface alteration and cell viability loss^{14–21,23–25,30}. However, the genericity of this mechanism is not supported by other reports on the harmful effects of TiO₂NPs on bacteria in the absence of light^{5,13,22,26,27}, and the lack of correlation between ROS production and the toxicity under UV illumination²⁹. In addition, non-ROS-related toxicity has been reported for nanomaterials other than TiO₂NPs but with similar photocatalytic properties such as ZnONPs^{32,35}, MgONPs³¹, CeO₂NPs³³, and fullerenes³⁴. These contrasting findings, together with the occurrence of lipid peroxidation under both light and dark conditions⁵, suggest the existence of a toxicity mechanism that involves non-photocatalytically TiO₂NP-induced ROS^{36,37}. The mode of action of TiO₂NPs in the dark has also been shown to translate into cell osmotic stress¹³ and cell membrane stress as a consequence of the electrostatic attachment of TiO₂NPs and mechanical membrane disruption²⁶. Similarly, Leung et al.²⁹ argued that the toxicity of TiO₂NPs originates from interactions between the nanoparticles and the outer membrane proteins and/or lipopolysaccharides (LPS), resulting in mechanical disruption of the cell membrane and possible entry of the nanoparticles into the cell. However, the genericity of this mechanism has been questioned by Buchman et al.³⁸, who showed the absence of a mechanistic connection between the toxicity of functionalized cationic AuNPs and the extent to which they bind to LPS. The above elements highlight that a comprehensive molecular description of the processes governing TiO₂NP toxicity to bacteria is incomplete.

Surprisingly, studies on the toxicity of metal oxide nanoparticles towards Gram-negative bacteria have neglected the possible production of membrane vesicles (MVs), despite the essential defense function they play in mitigating osmotic and oxidative stress^{39,40}. In addition, whilst the toxicity of TiO₂NPs in relation to their surface chemistry has been extensively studied^{5,6,26}, the role of cell surface composition has received far less attention. Accordingly, the current work has the following objectives: (i) decipher the processes that govern the toxicity of TiO₂NPs towards bacteria with controlled LPS surface phenotype, (ii) evaluate and inter-connect the cell responses probed at the gene, single-cell, and population scales over a broad range of TiO₂NP concentration conditions, and (iii) identify and explain the cell resistance and sensitivity patterns. Herein, we thus analyze the modes of action of TiO₂NPs on *Escherichia coli* deep rough mutants⁴¹ at the molecular, single-cell, and population levels. Exposures are performed in the dark and under hypotonic conditions that limit the initial aggregation of the TiO₂NPs. Targeted transcriptomics and single-cell nanomechanics, assessed by multiparametric atomic force microscopy (AFM), highlight that TiO₂NPs actuate osmotic stress as a consequence of cell

surface abrasion. This effect is found to be operational even at low TiO₂NP doses. Remarkably, dysregulated expressions of genes involved in osmotic stress tolerance are found to match non-monotonous variations in cell membrane elasticity and cell Turgor pressure with increasing TiO₂NP concentration. Additional fluorescence-based assays consistently support the observed TiO₂NP-mediated changes in membrane permeability and cell Turgor pressure, as well as oxidative cell damage triggered by the osmotic stress at sufficiently high TiO₂NP concentrations. The TiO₂NP modes of action are shown to intimately depend on the molecular composition of the LPS. In particular, TiO₂NP-induced vesiculation is evidenced for only the most sensitive mutant that harbors an unaltered LPS inner core. Direct and indirect defense functions of secreted MVs against TiO₂NPs are further highlighted. Overall, the results show that osmotic stress and cell vesiculation are associated with either TiO₂NP resistance or sensitivity depending on the LPS phenotype.

Results

***rfaG* mutation in *E. coli* leads to hypersensitivity to TiO₂NPs.** The selected *E. coli* K12 *rfa*-mutants express O-antigen-free LPS with distinct inner or outer core compositions (Fig. 1a). Of particular interest are the deep rough mutants JW3606 ($\Delta rfaG$) and JW3596 ($\Delta rfaC$), which lack the outer core LPS component and differ according to the presence or absence of heptose (hep) units in the inner core, respectively (Fig. 1a)⁴¹. In the following, JW3606 ($\Delta rfaG$) and JW3596 ($\Delta rfaC$) are thus referred to as JW3606 (hep+) and JW3596 (hep–). Selected TiO₂NPs (21 nm pristine radius) display a predominantly anatase structure and are positively charged under the adopted exposure conditions (Supplementary Fig. 1a, b). Briefly, cells were exposed in the dark for 20 h to—unless otherwise specified—0–50 mg/L TiO₂NPs at pH~5.5–6 under agitation conditions (see “Methods”). Preliminary measurements of the colony-forming units (CFUs) on cells exposed to a high TiO₂NP dose (100 mg/L) reveal that JW3606 (hep+) is the most sensitive to TiO₂NPs of all *rfa*-mutants tested (~3 log units difference, Fig. 1b). This finding underscores a connection between the toxicity of TiO₂NPs and the LPS inner core composition. Given that the responses of the wild type (WT), JW3601 ($\Delta rfaI$), JW3603 ($\Delta rfaB$), and JW3605 ($\Delta rfaP$) were similar to that of JW3596 (hep–) (see Fig. 1b), the latter mutant is chosen below for a detailed comparison with JW3606 (hep+). This choice is motivated by the deep rough phenotypes of JW3596 (hep–) and JW3606 (hep+) and their comparable sensitivity to, e.g., antibiotics or detergents⁴², which strikingly contrasts with their relative sensitivity to TiO₂NPs (Fig. 1). Figure 2a confirms the marked sensitivity of JW3606 (hep+) as compared to that of JW3596 (hep–) at TiO₂NP concentrations higher than 5 mg/L.

Flow cytometry analysis reveals a significant impact of TiO₂NPs on cell membrane potential (Fig. 2b), membrane permeability (Fig. 2c), and lipid peroxidation (Fig. 2d) for JW3606 (hep+) at TiO₂NP concentrations >1 mg/L. Data further support an oxidative stress (Fig. 2e) at concentrations >2 mg/L. All of these proxies depend on the TiO₂NP concentration according to clear dose–response relationships. The respective rates of change in lipid peroxidation (Fig. 2d) and oxidative stress levels (Fig. 2e) with increasing TiO₂NP concentration suggest that oxidative stress alone cannot explain lipid peroxidation. Interestingly, Fig. 2 shows a marked offset between the CFU-based response of JW3606 (hep+) with increasing TiO₂NP concentration (Fig. 2a) and that inferred from flow cytometry (Fig. 2b–e). Accordingly, provided that the TiO₂NP concentration is ≤5 mg/L, changes in the membrane potential and permeability, lipid peroxidation level, and oxidative stress necessarily mirror the

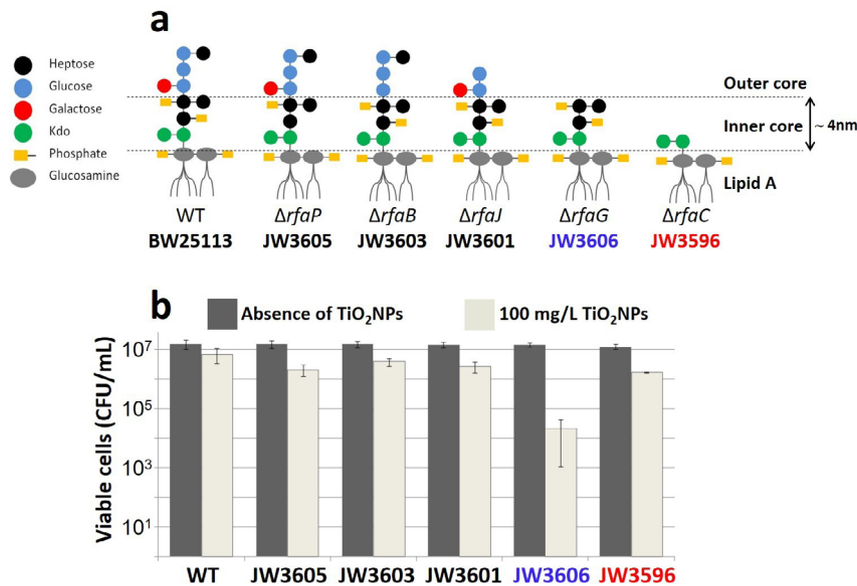


Fig. 1 Lipopolysaccharide phenotypes and preliminary assessment of TiO_2 NP toxicity towards *E. coli* K12 *rfa*-mutants at high TiO_2 NP concentration. **a *E. coli* K12 *rfa*-mutants, associated LPS phenotypes, and corresponding mutations in the *rfa* operon⁴¹. **b** Number of viable cells in CFU/mL unexposed and exposed to 100 mg/L TiO_2 NPs for 20 h in the dark. $n = 4$ for each tested condition.**

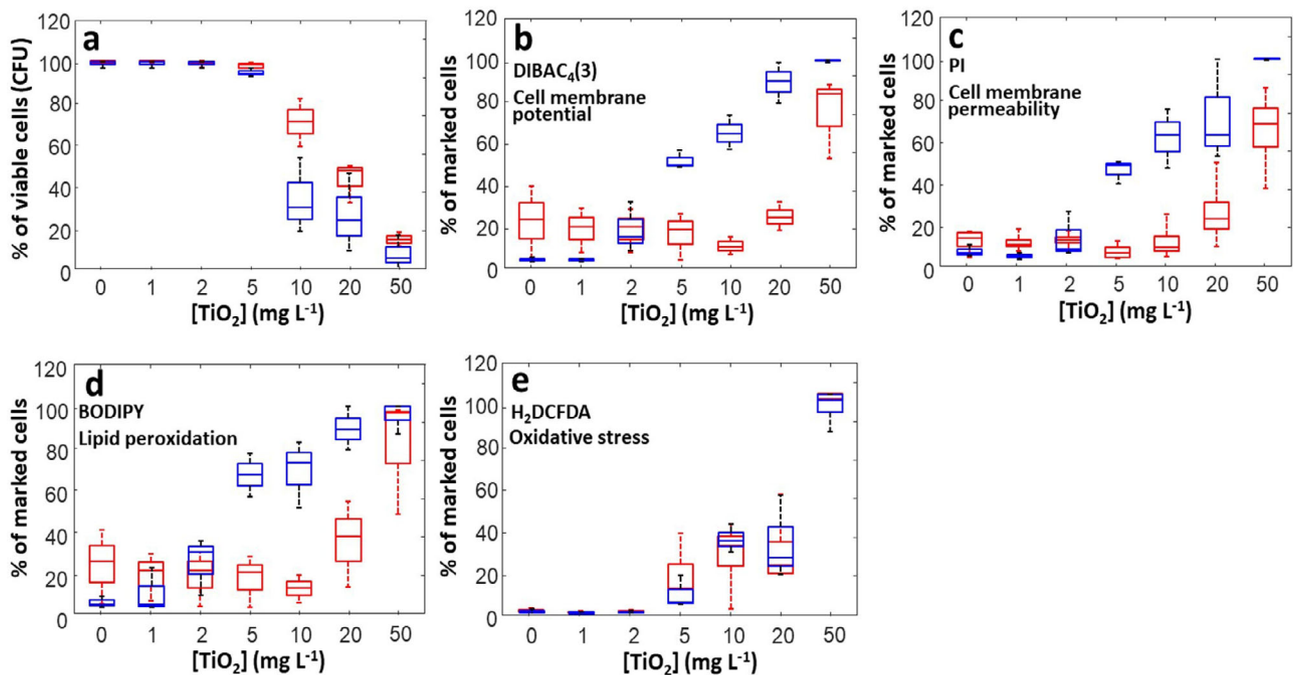


Fig. 2 JW3606 (hep+) is the deep rough mutant that is most sensitive to TiO_2 NPs. **a Dependence of JW3606 (hep+) (blue) and JW3596 (hep-) (red) viability on TiO_2 NP concentration. Normalized amounts of cells marked by membrane-selective fluorescent dyes for assessment of TiO_2 NP effects on **b** cell membrane potential, **c** cell membrane permeability, **d** lipid peroxidation, and **e** oxidative stress. As in **(a)**, results pertaining to JW3606 (hep+) and JW3596 (hep-) are represented by blue and red boxes, respectively, in the form of box plots. Data were derived from three independent experiments for each condition examined. The selected dyes and their targeted functions are specified for each panel with DIBAC₄(3) = bis-(1,3-dibutylbarbituric acid) trimethine oxonol, PI = propidium iodide, BODIPY = 4,4-difluoro-5,7-dimethyl-4-bora-3a,4a-diaza-s-indacene-3-propionic acid, and H₂DCFDA = 2',7'-dichlorodihydrofluorescein diacetate. Statistical significance testing and p -values are provided in the Supplementary Information.**

setting of cell defense mechanisms for maintaining viability (see following sections).

In agreement with Figs. 1b–2a, the situation for JW3596 (hep-) differs drastically from that for JW3606 (hep+) (Fig. 2b–e). The 10–20 mg/L TiO_2 NP concentration range marks the onset of effects on JW3596 (hep-), in contrast to the transition identified at 1–2 mg/L

for JW3606 (hep+). In addition, at TiO_2 NP concentrations ≤ 1 mg/L and in the absence of TiO_2 NPs in the exposome, membrane depolarization, membrane permeability, and the lipid peroxidation level (Fig. 2b–d) are significantly higher for JW3596 (hep-), in line with its larger LPS inner core truncation (Fig. 1)⁴³. At this stage, Fig. 2 highlights an apparent paradox: JW3596 (hep-), the mutant with a

native cell membrane that is most destabilized following inner core LPS truncation (i.e., in the absence of TiO₂NPs), is the one that exhibits greater resistance to TiO₂NPs.

The *ΔrfaG* mutant exhibits a TiO₂NP-dependent vesiculation phenotype whereas hypervesiculation of the *ΔrfaC* mutant is independent of TiO₂NP exposure conditions. Figure 3 reports the distribution profiles of electrophoretic mobilities (μ) for JW3606 (hep+) and JW3596 (hep-) in the presence of TiO₂NPs (0–50 mg/L) after 20 h exposure.

Starting with the JW3606 (hep+)–TiO₂NP system, electropherograms show the presence of two charged particle types, P1 and P2, materialized by the presence of peaks positioned at μ_{P1} and μ_{P2} in the range -4 to -5×10^{-8} and -1.6 to $-2.6 \times 10^{-8} \text{ m}^2 \text{ V}^{-1} \text{ s}^{-1}$, respectively (Fig. 3a). The apparition and extinction of the peaks depend on the TiO₂NP concentration, which underpins variation in the number of electrophoretically detected P1 and P2 entities. P1 particles refer to JW3606 (hep+) cells as evidenced by a previous electrokinetic study performed in the absence of TiO₂NPs⁴³ and by the measurements on 0.22 μm filtered TiO₂NP–bacteria suspensions (Supplementary Fig. 2a). The μ -distribution corresponding to JW3606 (hep+) is slightly shifted to negative values with increasing TiO₂NP concentrations from 0 to 2 mg/L (Fig. 3a). With a further increase in the TiO₂NP concentration up to 10 mg/L, the absolute

value of μ corresponding to the maximum of the P1 (bacteria)-related peak in Fig. 3a decreases before the bacteria-associated signal completely vanishes at TiO₂NP concentrations ≥ 20 mg/L. This signal suppression is due to the electrostatically favored formation of aggregates between the (negatively charged) cells (Fig. 3) and the micron-sized (positively charged) TiO₂NP assemblies (Supplementary Fig. 1) and their subsequent sedimentation. This sedimentation process is magnified by intracellular material and cell surface residues that are possibly released under extreme stress conditions⁴⁴. The aforementioned shift of the P1 peak to negative electrophoretic mobility values is the signature of TiO₂NP-induced modification of the cell surface structure. Indeed, any significant adsorption of positively charged TiO₂NPs onto JW3606 (hep+) is excluded as it should lead to a mobility shift in a direction opposite to that observed. Instead, the trend fits qualitatively with the following picture: the (negative) charges carried by the outer cell membrane surface increasingly contribute to the electrophoretic mobility of the cell via their enhanced exposure to the surrounding solution following TiO₂NP-mediated abrasion of peripheral cell components such as LPS. This removal of protruding components of the cell surface is further accompanied by a reduction of the hydrodynamic friction exerted by the whole-cell envelope on the electroosmotic flow developed in the vicinity of the cell surface under electrophoresis conditions: such a reduction also contributes to an increase in the

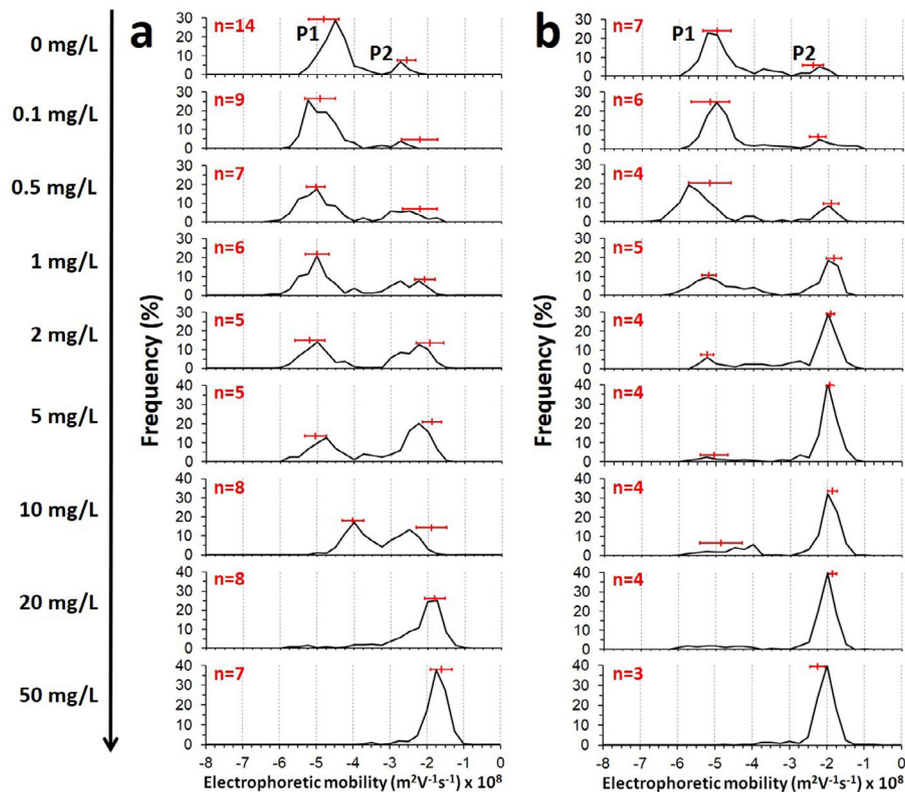


Fig. 3 Electrokinetic fingerprints of JW3606 (hep+), JW3596 (hep-), and secreted MVs as a function of TiO₂NP concentration. Illustrative electropherograms for (a) JW3606 (hep+) and (b) JW3596 (hep-) suspensions after 20 h incubation with TiO₂NPs at various concentrations (indicated). Horizontal red bars correspond to the position of the maxima (mean values \pm standard deviations) derived from the measurement of n (indicated) electrophoretic mobility distributions for the suspensions prepared from different cell cultures or colonies, with three replicates for each measurement. The reported illustrative mobility distributions are averaged over three replicates. Analysis indicates that the number of detected electrophoretic trajectories decreases by 16% and 20% in the 1–10 mg/L and 1–2 mg/L TiO₂NP concentration regimes (analysis on 1:50 diluted JW3606 (hep+)–TiO₂NP and JW3596 (hep-)–TiO₂NP samples, respectively), which features significant P1 particle sedimentation under these conditions. A 61–70% increase in the number of trajectories is further observed in 1:50 diluted JW3606 (hep+)–TiO₂NP and JW3596 (hep-)–TiO₂NP samples, respectively, on further increasing the TiO₂NP concentrations to 50 mg/L due to the released membrane material as a consequence of the mode of action of the TiO₂NPs. See text for details.

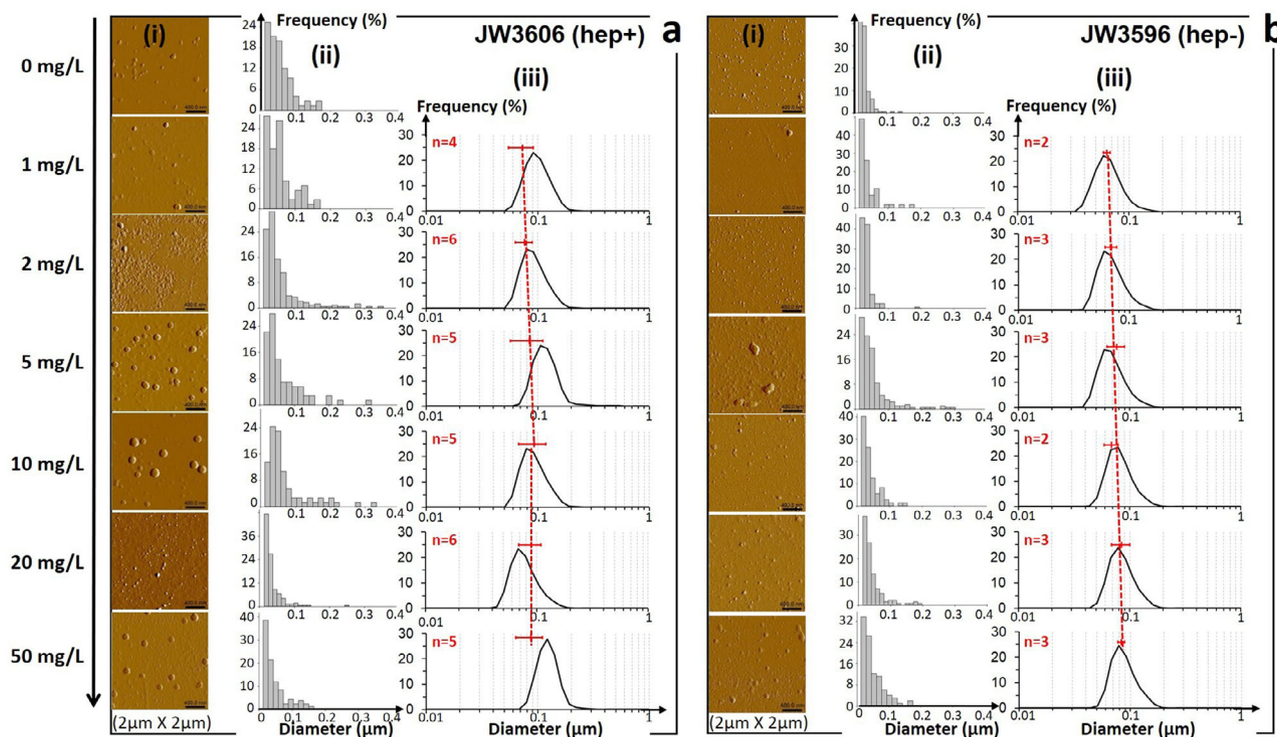


Fig. 4 MV imaging and MV size distributions as a function of TiO_2NP concentration. **i** AFM imaging of MVs and **ii** corresponding estimations of their size distribution, and **iii** DLS-derived MV size distribution as a function of TiO_2NP concentration (indicated) for **(a)** JW3606 (hep+) and **(b)** JW3596 (hep-). In **a** **(iii)** and **b** **(iii)**, red dotted lines are guides to the eye and horizontal red bars correspond to the position of the maxima (mean values \pm standard deviations) derived from DLS measurement of n (indicated) size distributions for suspensions prepared from different cell cultures or colonies, with three replicates for each measurement. The reported illustrative size distributions are averaged over three replicates. DLS measurements of MV size were performed on 0.22 μm -filtered suspensions (1:10 diluted in ultrapure water, see “Methods”) and results were confirmed by measurements on 0.45 μm -filtered suspensions. Qualitative MV size estimations from AFM imaging were based on the analysis of 0.45 μm -filtered suspensions. Detection of MVs by DLS in the absence of TiO_2NP was not possible due to an insufficient amount of particles in the solution.

absolute magnitude of the cell electrophoretic mobility. These connections between the cell electrophoretic properties and the cell surface organization are in line with predictions from the soft surface electrokinetic theory^{45,46} and with previous conclusions on the impact of surface appendages on bacteria electrohydrodynamics^{46,47}.

The presence of P2 particles is identified over the whole range of tested conditions (Fig. 3a), even in the absence of TiO_2NPs . The intensity of the P2 peak increases with increasing TiO_2NP concentrations from 0.5 to 50 mg/L. Given the positive mobility of TiO_2NPs (Supplementary Fig. 1a, b), P2 necessarily refers to particles other than TiO_2NPs , as confirmed by electrokinetic measurements on 0.22 μm -filtered cell- TiO_2NP suspensions that are free of bacteria and TiO_2NP aggregates (Supplementary Fig. 2a). Imaging of the filtrates by AFM further reveals that P2 particles are closed spheroids, that are polydisperse in size, with a diameter ranging from ca. 30–40 to 200 nm (Fig. 4a(i),(ii)), in accordance with refined measurements by dynamic light scattering (DLS) (Fig. 4a(iii)). These properties typically correspond to those of membrane vesicles (MVs for short) secreted by Gram-negative bacteria through the budding-out of their outer membrane⁴⁰. Fluorescent labeling confirms the nature of the P2 particles (Fig. 5a (i),(ii)) and the significant increase of their produced amount with increasing TiO_2NP concentration (Fig. 5b). To the best of our knowledge, these results establish for the first time the involvement of MVs in cell responses to TiO_2NP stressors. Remarkably, DLS data (Fig. 4a(iii)) indicate that the mean diameter of secreted MVs increases from ca. 70 to 95 nm in the 1–10 mg/L TiO_2NP concentration range before levelling off at higher concentrations

(recalling that the 10–20 mg/L concentration regime is that where a significant loss of cell viability is reached, Fig. 2a). Finally, an increase in the MV concentration in the exposome above a threshold value via a short-term (15 min) co-incubation procedure (see details in “Methods”) leads to a decrease in the size of TiO_2NP aggregates (Fig. 5c). This result demonstrates the existence of (electrostatically favored, Fig. 3 and Supplementary Fig. 1) MV- TiO_2NP interactions, as further detailed in the “Discussion” section. The presence of MVs in solution further leads to a reduction in the TiO_2NP -induced membrane permeabilization (Fig. 5d), thereby supporting the role of MVs in mitigating TiO_2NP toxicity.

Following the above methodology, P1- and P2-contributions to electropherograms of JW3596 (hep-)- TiO_2NPs (Fig. 3b) are attributed to JW3596 (hep-) and MVs, respectively (Figs. 4b, 5a (iii), (iv), and Supplementary Fig. 2b). The electrophoretic fingerprint of JW3596 (hep-) shifts slightly to negative values with increasing TiO_2NP concentrations from 0 to 2 mg/L and practically vanishes at concentrations >2 mg/L (Fig. 3b). This contrasts with the JW3606 (hep+)- TiO_2NP system for which the threshold TiO_2NP concentration that marks the switch from a bacteria- to an MV-dominated μ distribution is ca. 20 mg/L (Fig. 3a). This difference is possibly due to the larger propensity of JW3596 (hep-) to aggregate⁴⁸, as the absence of protruding surface LPS (Fig. 1) may reduce the magnitude of the stabilizing steric forces that are operational between neighboring cells. Like for JW3606 (hep+), MV production by JW3596 (hep-) in the absence of TiO_2NPs is revealed by electrokinetics and AFM (Figs. 3b and 4b(i),(ii)), in accordance with their known

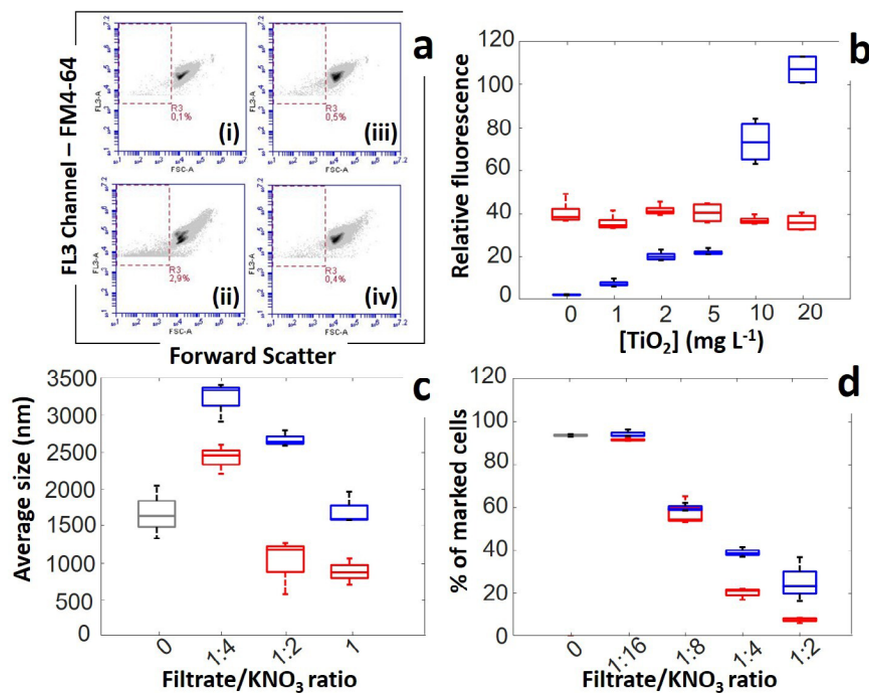


Fig. 5 The dependences of JW3606 (hep+) and JW3596 (hep-) vesiculation phenotypes on TiO₂NP concentration are strikingly different. **a** Illustrative cytograms with detection of FM4-64 labeled bacteria and MVs (red dotted window) (gate set on FL3 positive events) for JW3606 (hep+) unexposed **a(i)** and exposed to 20 mg/L TiO₂NPs **a(ii)**, and JW3596 (hep-) unexposed **a(iii)** and exposed to 20 mg/L TiO₂NPs **a(iv)**. **b** Evaluation of secreted MV amounts expressed in relative fluorescence intensity units after Syto9 labeling as a function of TiO₂NP concentration (see *Methods*). Results pertaining to JW3606 (hep+) and JW3596 (hep-) are represented by blue and red boxes, respectively. Data were derived from three independent measurements for each condition tested. **c** Average size of TiO₂NP–MV aggregates measured by DLS after 15 min co-incubation of TiO₂NPs (50 mg/L) with MVs obtained from JW3606 (hep+) (blue boxes) and JW3596 (hep-) (red boxes), for different MV/TiO₂NP concentration ratios at fixed 10 mM KNO₃ background electrolyte. MVs were produced by the mutants incubated for 20 h in 10 mM KNO₃ (in the absence of TiO₂NPs), and they were subsequently collected after 0.45 μm filtration of the suspensions. MV concentration in **(c)** was changed upon diluting the filtrate with 10 mM KNO₃ electrolyte (filtrate/KNO₃ volume ratio). The gray box corresponds to the condition without MVs (KNO₃ and TiO₂NPs are only present in the solution). **d** Normalized amounts of JW3606 cells (hep+; TiO₂NPs-sensitive strain) marked with propidium iodide (to target changes in membrane permeability) after 20 h co-incubation with TiO₂NPs (50 mg/L) and MVs previously obtained from JW3606 (hep+) (blue boxes) and JW3596 (hep-) (red boxes) at different TiO₂NP/MV concentration ratios, and fixed 10 mM KNO₃ electrolyte (MV were collected as in panel **(c)**). The gray box corresponds to the condition without added MVs (bacteria, KNO₃ and TiO₂NPs are present in the solution). In **(b, c, d)**, data are reported in the form of box plots.

vesiculation phenotypes⁴⁹. Similarly to JW3606 (hep+), the mean diameter of MVs generated by JW3596 (hep-) increases with TiO₂NP concentration with a ca. 20 nm increase over the whole range of tested conditions (Fig. 4b(iii)). Most importantly, MV production by JW3596 (hep-), unlike that by JW3606 (hep+), remains independent of TiO₂NP concentration (Fig. 5b). Additionally, at sufficiently low TiO₂NP concentrations (≤ 5 mg/L) and in the absence of TiO₂NPs in solution, vesiculation by JW3596 (hep-) is much more important than that of JW3606 (hep+) (Fig. 5b). This latter finding correlates positively with the respective magnitudes of membrane permeability/depolarization and lipid peroxidation detailed for the two mutants in Fig. 2 at low TiO₂NP doses. The larger amount of MVs secreted by JW3596 (hep-) further correlates with a more efficient reduction in the size of the TiO₂NP aggregates (Fig. 5c) and a better membrane protection against TiO₂NPs (Fig. 5d) in comparison with JW3606 (hep+).

Nanomechanical properties of the $\Delta rfaG$ mutant envelope vary non-monotonously with TiO₂NP concentration. Spatial-distributions of the cell Young modulus (E in Pa) and cell stiffness (k_{cell} in Nm^{-1}), indicative of the cell Turgor pressure^{46,50}, were evaluated for (un)exposed JW3606 (hep+) and JW3596 (hep-) (Figs. 6a, b(i),(ii) and 7a, b(i),(ii), respectively) from theoretical analysis⁵⁰ of 65,536 approach force curves collected by

atomic force spectroscopy operated in PeakForce Tapping mode on $500 \times 500 \text{ nm}^2$ single-cell surface areas (see details in “Methods”). Below, we further introduce δ defined by the value of the indentation (in nm) which marks the transition between the non-linear elastic deformation of the cell envelope and the linear compliance domain in the force versus indentation curve measured at a given location (pixel) of the cell surface⁵⁰. The spatial distribution of δ over the scanned cell surface area was obtained according to the theoretical procedure detailed elsewhere⁵⁰, and it is reported in Figs. 6b(iii) and 7b(iii) for JW3606 (hep+) and JW3596 (hep-) exposed to different TiO₂NP concentrations, respectively. The distributions of E , k_{cell} , and δ values over the probed cell surface area are further provided in Figs. 6c–7c. For the sake of comparison, E , k_{cell} , and δ derived as a function of TiO₂NP concentration were converted into the normalized quantities $R_{E,0}$, $R_{k_{\text{cell}},0}$ and $R_{\delta,0}$ defined by $R_{X,0} = (X - X_0)/X_0$ with $X \equiv E$, k_{cell} , or δ and X_0 the reference median value of the X -distribution measured at 0 mg/L TiO₂NPs. Figure 8 finally collects the median values of the E -, k_{cell} -, and δ -distributions derived as a function of TiO₂NP concentration from measurements on eight cells from distinct grown colonies.

Results for JW3606 (hep+) evidence a synchronic and non-monotonous dependence of $R_{E,0}$ and $R_{k_{\text{cell}},0}$ on TiO₂NP concentration (Fig. 8a(i),(ii)) with a $R_{E,0}$ -distribution width indicative of a spatial heterogeneity that is largest in the 1–2 mg/L range and at

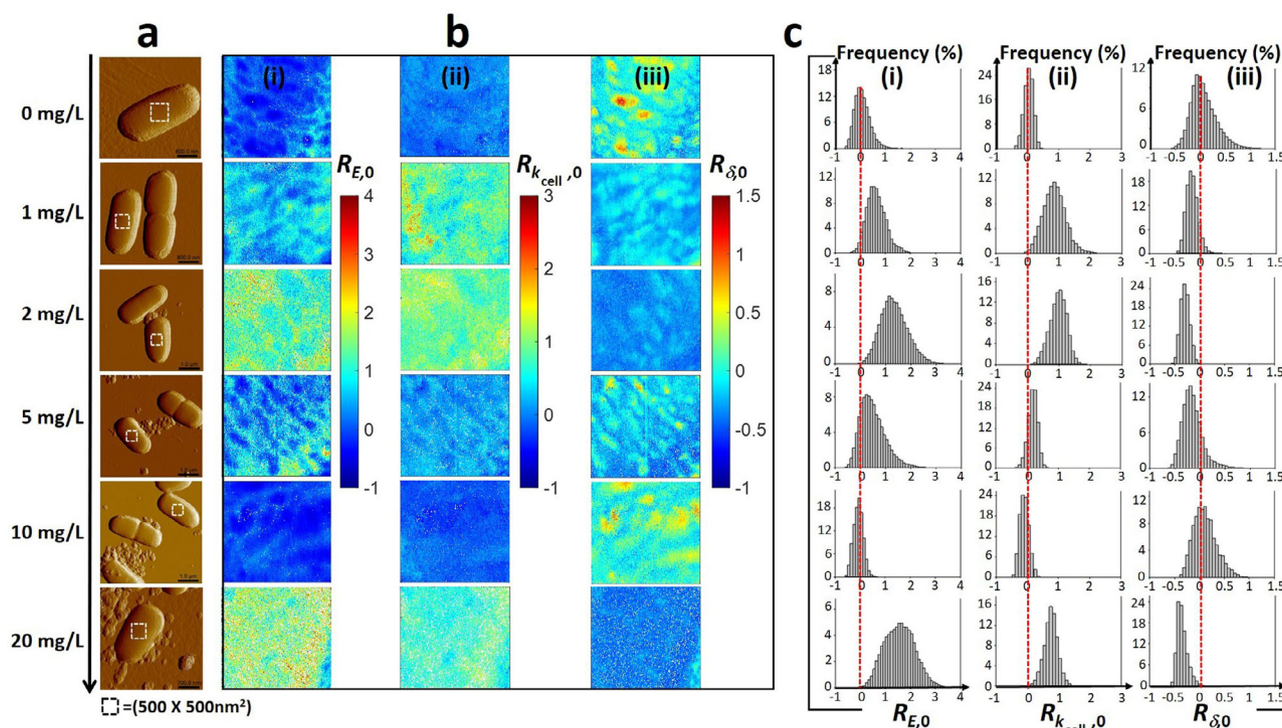


Fig. 6 Multiparametric AFM evidences TiO_2NP -mediated changes in cell surface elasticity and Turgor pressure for JW3606 (hep+) with increasing TiO_2NP concentration. **a** Illustrative AFM deflection images of JW3606 (hep+) after 20 h co-incubation with TiO_2NPs of various concentrations (indicated). The white boxes define the $500 \times 500 \text{ nm}^2$ cell surface areas where 65,536 force curve measurements were carried out. **b** Corresponding spatial maps of **(i)** Young modulus E , **(ii)** cell stiffness k_{cell} , and **(iii)** indentation δ expressed in terms of $R_{E,0}$, $R_{k_{\text{cell}},0}$ and $R_{\delta,0}$, respectively (see text for details). **c** Histogram-distributions of $R_{E,0}$, $R_{k_{\text{cell}},0}$, and $R_{\delta,0}$ corresponding to the maps given in **(b)**. Red dotted lines in **(c)** are only guides to the eye.

20 mg/L (Fig. 6b, c). In detail, $R_{E,0}$ and $R_{k_{\text{cell}},0}$ increase from 0 to 2 mg/L (regime I), decrease with a further increase in the TiO_2NP concentration up to 10 mg/L (regime II), and increase again from 10 to 20 mg/L (regime III).

The increase in $R_{E,0}$ and $R_{k_{\text{cell}},0}$ in regime I mirrors a stiffening of the cell envelope and a concomitant increase of the cell Turgor pressure. These trends compound a significant decrease of $R_{\delta,0}$ (Fig. 6c) with a median δ value decreasing from 42 to 30 nm (Fig. 8a(iii)). The findings are consistent with a TiO_2NP -mediated removal of the softer outer cell surface components, which leads to a reduction of the indentation range where non-linear deformation of the overall cell envelope is operational. The components removed by TiO_2NP action in regime I likely include LPS, which is qualitatively supported by: (i) recent nanomechanics analysis of WT, JW3601, and JW3606 (Fig. 1) unexposed to NPs⁴³, showing that the reduction in LPS length along this mutant gradient leads to increase in cell elasticity and stiffness, and (ii) TiO_2NP -induced cell surface abrasion identified from JW3606 (hep+) electrokinetics (Fig. 3a).

In regime II, the decrease in $R_{E,0}$ and $R_{k_{\text{cell}},0}$, associated with an increase in $R_{\delta,0}$, reflects a TiO_2NP -mediated softening of the cell envelope and a decrease in Turgor pressure. The increase in $R_{\delta,0}$ corresponds to an increase of the δ median from 30 to 45 nm. These different observations underpin a larger indentation into a mechanically softer biosurface (as compared to the regime I), in accordance with the loss of membrane integrity and increase in membrane permeability independently evidenced by flow cytometry for TiO_2NP concentrations $\geq 2 \text{ mg/L}$ (Fig. 2c). They further corroborate nanomechanics observations of *E. coli* exposed to SiO_2NPs ⁵¹. The drastic changes in membrane structure suggested by AFM in regime II are all the more favored as removal of the cell surface components in regime I has significantly weakened/

disorganized the outer cell membrane, thereby rendering it more prone to TiO_2NP -induced damage following lipid peroxidation (Fig. 2d) and oxidative stress (Fig. 2e). These processes lead to the leakage of intracellular ions and cell envelope components, which facilitates the formation of cohesive TiO_2NP aggregate in the cell's vicinity, as suggested by AFM imaging of cells for TiO_2NP concentrations $\geq 5 \text{ mg/L}$ (Fig. 6a).

In regime III, the increase in E and k_{cell} as the TiO_2NP concentration increases from 10 to 20 mg/L, together with the decrease in $R_{\delta,0}$ (decrease in δ from 45 to 27 nm) highlight that the AFM probe now significantly interacts with the membrane components that are more rigid and subject to reduced indentation (Fig. 8a). Accordingly, we suggest that the successive TiO_2NP -mediated LPS removal and outer membrane disruption taking place in regimes I and II lead to a significant AFM sensing of the rigid peptidoglycan layer in regime III. The heterogeneity in the bacterial surface landscape, reflected by the width of the $R_{\delta,0}$ -distributions at the single-cell level (Fig. 6c), basically decreases with ongoing cell surface scouring (regime I), then it increases upon severe action of TiO_2NPs on the outer membrane (regime II) and finally decreases significantly when the rigid peptidoglycan layer is significantly exposed and the outer membrane has significantly disintegrated (regime III). Close inspection of Fig. 6b indicates that details of $R_{E,0}$ -based surface heterogeneity do not necessarily match those inferred from $R_{k_{\text{cell},0}}$ -maps. This observation stems from the fact that E mostly reflects elastic properties of the peripheral cell surface envelope, whereas k_{cell} integrates properties of the whole membrane barrier via its connection to intracellular Turgor pressure^{43,46}. Last, the spatially resolved $R_{\delta,0}$ property unveils irregular cell surface patterns in the absence of TiO_2NPs , possibly connected to the surface distribution of protruding LPS^{43,52}.

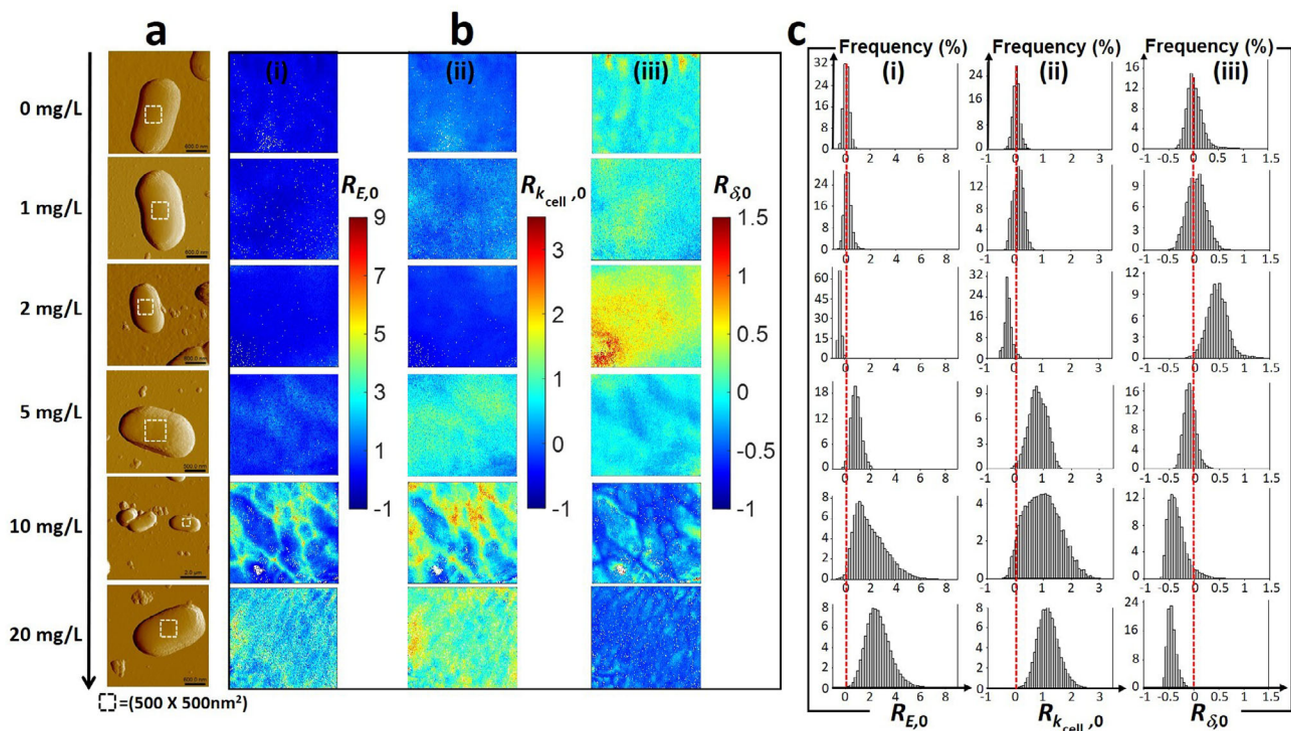


Fig. 7 Multiparametric AFM evidences TiO_2NP -mediated changes in cell surface elasticity and Turgor pressure for JW3596 (hep $-$) at larger TiO_2NP concentrations than for JW3606 (hep $+$). **a** Illustrative AFM deflection images of JW3596 (hep $-$) after 20 h co-incubation with TiO_2NPs of various concentrations (indicated). The white boxes define the $500 \times 500 \text{ nm}^2$ cell surface areas where 65,536 force curve measurements were carried out. **b** Corresponding spatial maps of **(i)** Young modulus E , **(ii)** cell stiffness k_{cell} , and **(iii)** indentation δ expressed in terms of $R_{E,0}$, $R_{k_{\text{cell}},0}$ and $R_{\delta,0}$, respectively (see text for details). **c** Histogram distributions of $R_{E,0}$, $R_{k_{\text{cell}},0}$ and $R_{\delta,0}$ corresponding to the maps given in **(b)**. Red dotted lines in **(c)** are only guides to the eye.

Concerning JW3596 (hep $-$), the dependence of $R_{E,0}$, $R_{k_{\text{cell}},0}$ and $R_{\delta,0}$ on TiO_2NP concentration and their corresponding distributions at the cell surface differ from those derived for JW3606 (hep $+$) (Figs. 7, 8). Namely, $R_{E,0}$, $R_{k_{\text{cell}},0}$ ($R_{\delta,0}$) medians slightly decrease (increase, respectively) with increasing TiO_2NP concentrations from 0 to 2 mg/L (so-called regime α), but overlap in the statistical distributions (Fig. 8b) prevents firm conclusions from being drawn. In contrast, for TiO_2NP concentrations $>5 \text{ mg/L}$ (regime β) $R_{E,0}$ and $R_{k_{\text{cell}},0}$ ($R_{\delta,0}$) notably increase (decrease, respectively). The range of TiO_2NP concentrations corresponding to regime β matches consistently the one where we measured a dramatic decrease in CFU (Fig. 2a) and the most significant changes in membrane potential (Fig. 2b), membrane permeabilization (Fig. 2c), lipid peroxidation (Fig. 2d), and oxidative stress (Fig. 2e). Regime β (5–20 mg/L) corresponds to a stiffening of the cell envelope and to an increase in cell Turgor pressure. These signatures are qualitatively similar to those described for JW3606 (hep $+$) in regimes I and III marked by cell surface abrasion and by increased contribution of the peptidoglycan layer to cell nanomechanics, respectively. The former process is detected by electrokinetics despite parasiting JW3596 (hep $-$) aggregation (Fig. 3b), and the latter requires a prior outer membrane alteration that is poorly supported by the slight decrease in $R_{E,0}$ and $R_{k_{\text{cell}},0}$ in regime α (as compared to that in regime II for JW3606 (hep $+$)). Accordingly, we hypothesize that TiO_2NPs predominantly impact on JW3596 (hep $-$) in regime β via cell envelope scouring (with resulting decrease in $R_{\delta,0}$, Fig. 8b(iii)), accompanied by significant TiO_2NP -induced increases in membrane permeability, membrane depolarization, lipid peroxidation, and oxidative stress. The heterogeneity of the so-modified cell surface is clearly identified from the maps in Fig. 7b at 10–20 mg/L TiO_2NPs . Overall,

multiparametric AFM at the single-cell level supports the results from macroscopic fluorescence-based assays: in comparison to JW3606 (hep $+$), JW3596 (hep $-$) is defined by a remarkable resistance phenotype against TiO_2NPs .

Transcriptomic analysis of deep rough mutants shows cell response to dominant osmotic stress. After 20 h exposure of JW3606 (hep $+$) and JW3596 (hep $-$) to 0–20 mg/L TiO_2NPs , the expression of selected genes involved in osmotic and oxidative stress tolerance was quantified by RT-qPCR (see “Methods”).

We first consider the osmotic stress-induced transcriptional response of JW3606 (hep $+$). Figure 9a evidences a dysregulation of the *ompF* gene that encodes OmpF protein which allows passive transport of small solutes across the membrane⁵³. This gene is downregulated at 1–2 mg/L TiO_2NPs , and its expression level increases with increasing TiO_2NP concentrations from 2 to 20 mg/L, i.e., for doses where membrane permeability significantly increases (Fig. 2c). This non-monotonous *ompF* expression with varying TiO_2NP concentration is strikingly reminiscent of that observed in regimes I–II for the Turgor pressure (Fig. 8a (ii)). In particular, reduction in *ompF* gene expression compounds the increase of the Turgor pressure in the 0–2 mg/L range, which is in accordance with the findings by Graeme-Cook⁵⁴, who reported that *ompF* expression is switched off by Turgor stress. The *osmB* gene encoding an osmotic stress-inducible lipoprotein⁵³ is further severely downregulated as the TiO_2NP concentration increases from 5 to 20 mg/L, and so is the expression of *osmC*, another osmotic stress-induced gene (Fig. 9b, c)⁵³. This marked downregulation is also observed for the *otsB* gene (Fig. 9d), which encodes a phosphatase involved in trehalose production to resist against osmotic stress⁵³, for the

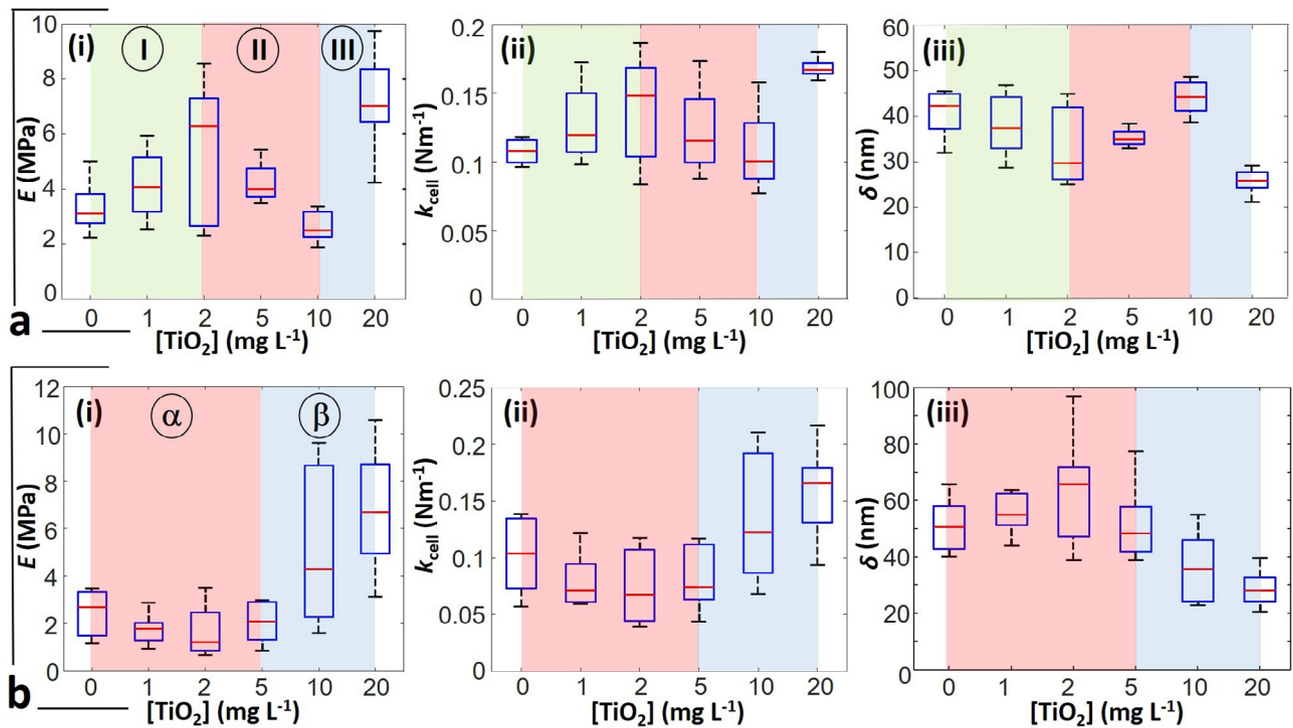


Fig. 8 Nanomechanical properties of JW3606 (hep+), unlike those of JW3596 (hep-), vary non-monotonously with increasing TiO₂NP concentration. Dependence of (i) Young modulus E , (ii) cell stiffness k_{cell} , and (iii) indentation δ as a function TiO₂NP concentration for **a** JW3606 (hep+) and **b** JW3596 (hep-) after 20 h exposure to TiO₂NPs. Data correspond to distributions of the median values of E , k_{cell} , and δ represented in the form of box plots and derived for each condition from measurements on $n = 8$ cells issued from different colonies or different cell cultures. Regimes I, II, III, α and β refer to distinct modes of action of TiO₂NPs on JW3606 (hep+) and JW3596 (hep-), see details in the text. The red lines are the median of the plotted distributions. Statistical significance testing and p -values are provided in the Supplementary Information.

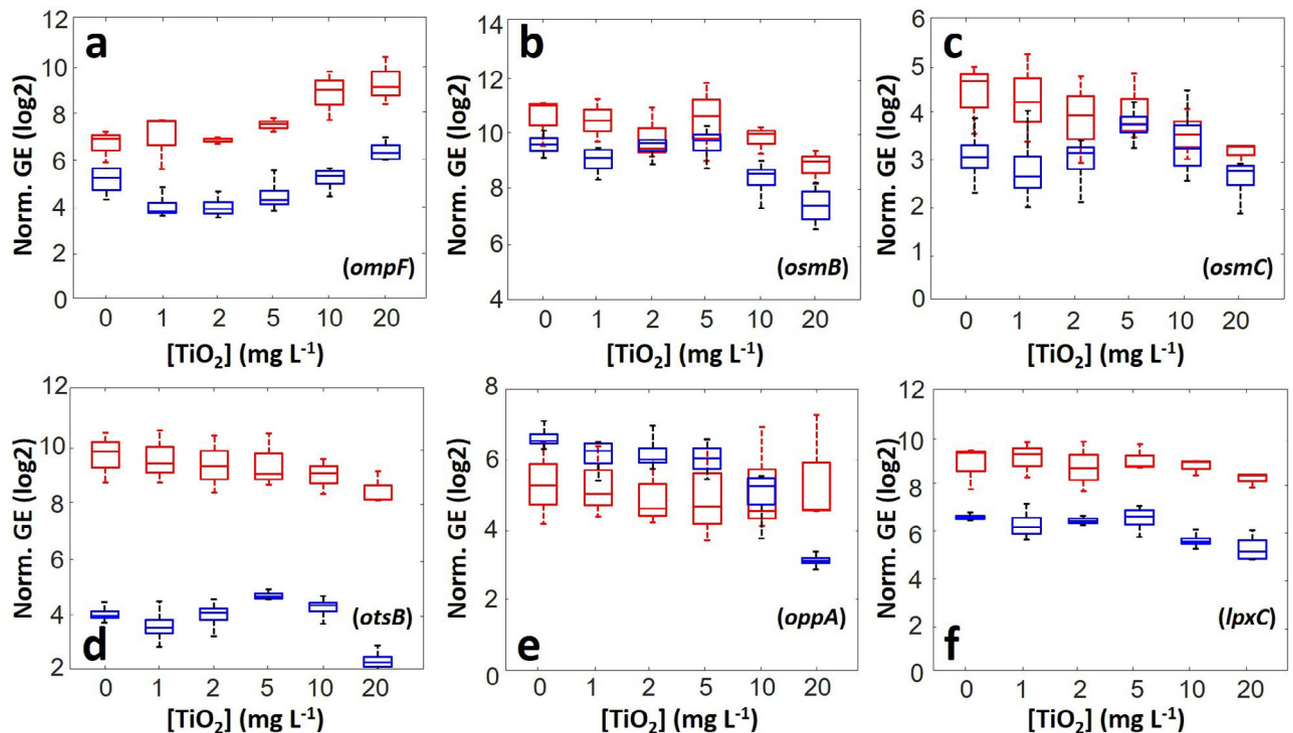


Fig. 9 Transcriptomics evidences dominant osmotic stress generated by TiO₂NPs. Expression levels of genes (**a**: *ompF*, **b**: *osmB*, **c**: *osmC*, **d**: *otsB*, **e**: *oppA*, **f**: *lpxC*) involved in the osmotic stress-response of JW3606 (hep+) (blue) and JW3596 (hep-) (red) as a function of TiO₂NP concentration, and represented in the form of box plots ($n = 4$ for each condition tested). The normalized gene expression (norm. GE) was calculated from the ratio between the intensity of the targeted gene and the geometric mean intensity of reference genes. Statistical significance testing and p -values are provided in the Supplementary Information.

oppA gene (Fig. 9e), which encodes a periplasmic binding protein of an ABC transporter that mediates high-affinity uptake of oligopeptides⁵³, and to a lesser extent for the *lpxC* gene (Fig. 9f), which is known to play a regulatory role in lipid A biosynthesis⁵³. The expressions of several genes encoding scavenger enzymes that protect cells against oxidative stress are further provided in Supplementary Fig. 3 for JW3606 (hep+). Among all tested genes, *sodB* and *ahpC* are those that are most significantly upregulated with increasing TiO₂NP concentrations from 2–5 to 20 mg/L. They encode a superoxide dismutase and an alkyl hydroperoxide reductase, respectively, which are known to participate in the antioxidant defense mechanism against O₂^{•-}- and H₂O₂-induced oxidative stress⁵³ as detected by flow cytometry for TiO₂NP concentrations ≥5 mg/L (Fig. 2e). Overall, the action of TiO₂NPs at sufficiently low TiO₂NP doses (<2–5 mg/L) mainly results in osmotic stress that couples to oxidative stress at higher concentrations. This finding is in line with literature^{55–58} suggesting that osmotic stress can lead to oxidative cell damage via disturbance of membrane components of the respiratory chain⁵⁸.

A similar conclusion is obtained for JW3596 (hep–) cells (Fig. 9 and Supplementary Fig. 3) albeit with a few remarkable differences. Namely, *ompF* expression in JW3596 (hep–) remains stable up to a TiO₂NP concentration of 2–5 mg/L and, similarly to Turgor pressure (Fig. 8b(ii)), it increases significantly at higher TiO₂NP concentrations. Whereas the downregulation of *osmB* and *osmC* at high TiO₂NP concentrations is a feature shared by JW3606 (hep+) and JW3596 (hep–), for this latter mutant the expression levels of *otsB*, *oppA*, and *lpxC* genes remain practically constant over the whole range of tested concentrations. These results imply that TiO₂NPs impact the transcriptional response of JW3596 (hep–) to osmotic stress to a lesser extent than they do for JW3606 (hep+), in comparison to their respective controls (absence of TiO₂NPs). Still, the overall magnitude of the osmotic stress, in the absence or presence of TiO₂NPs, remains larger for JW3596 (hep–) (Fig. 9) than for JW3606 (hep+) as judged by the corresponding gene expressions levels. This finding correlates with the larger vesiculation capacity of JW3596 (hep–) either in the absence of nanoparticles or at sufficiently low TiO₂NP concentrations (Fig. 5b), with their larger membrane permeability (Fig. 2c) and depolarization (Fig. 2b), and lipid peroxidation (Fig. 2d) in the 0–2 mg/L concentration range. Also, the TiO₂NP-independent expressions of *katG* and *sodB*⁵³ remain much lower than those for JW3606 (hep+) and only the transcription of *ahpC* is found to increase significantly at TiO₂NP concentrations >5 mg/L (but with lower basal level compared to JW3606 (hep+), Supplementary Fig. 3e), in line with the oxidative stress detected under such concentration conditions (Fig. 2e).

Discussion

By a combination of cell viability, fluorescence, electrokinetic, nanomechanical, and transcriptomic analyses, we provide in Fig. 10 a schematic representation of the mechanisms that govern TiO₂NP toxicity towards the most sensitive JW3606 (hep+) mutant, starting from the situation of Fig. 10a with cells featuring reduced vesiculation in the absence of TiO₂NPs (Fig. 5b). For concentrations between 0 and 2 mg/L (regime I, Fig. 10b), TiO₂NPs contribute to cell surface abrasion via removal of envelope components including LPS, and to a gradual exposure of the moderately altered outer membrane surface (Figs. 3a, 6, and 8a). Regime I is where cell osmoregulation that takes place in the absence of TiO₂NPs (Fig. 10a) is inactivated by a growing destabilization of the outer membrane and the onset of membrane permeability increase (Fig. 2b, c). As a result, cell Turgor pressure increases (Figs. 6 and 8a(ii)) as a consequence of water

entry under the selected hypotonic conditions, and the cell Young modulus increases (Fig. 8a(i)) due to the removal of the softest peripheral membrane components. The evidenced Turgor stress is further consistent with the downregulation of *ompF*, which prevents the additional entry of small hydrophilic solutes (Fig. 9a).

In regime I, MV production gently sets in with increasing TiO₂NP concentration (Fig. 5b), and membrane integrity is not yet critically compromised (Fig. 2). Secreted vesicles probably mediate membrane stress relief via their evacuation of potentially harmful products, such as proteins or LPS, that accumulate in the periplasmic space with or without modification of the outer membrane-peptidoglycan linking lipoprotein levels⁵⁹. Whereas the contribution of MVs to cell defense against e.g., antibiotics is well established⁶⁰, their existence and roles have never been documented in the context of metallic oxide NP toxicity. In regime I, hyperproduction of MVs is not required to expel the moderate amount of residues that accumulate following LPS removal and associated alteration of the outer membrane.

With further increase in the TiO₂NP concentration (regime II, 2–10 mg/L, Fig. 10c), the mechanical action of TiO₂NPs on the cell surface intensifies and leads to significant loss of membrane integrity, increase in membrane permeability, and to marked membrane depolarization, oxidative stress and membrane lipid peroxidation (Fig. 2b–e). In turn, the underlying cell surface damage generates a pronounced cell envelop softening which is materialized by a decrease in cell surface elasticity and an increase of the threshold indentation that separates the non-linear deformation and compliance regimes (Figs. 6 and 8a(i),(iii)) in the AFM force–indentation curves. Also, a decrease in cell Turgor pressure (Figs. 6 and 8a(ii)) arises due to an increase in membrane permeability (Fig. 2c) and water efflux. The subsequent release of intracellular material favors NP aggregation (Fig. 6a). Cells then attempt to cope with the enhanced production of endogenous waste molecules in the periplasm via (i) an increase in the produced amount of MVs which act as shuttles to export over-accumulated moieties (Fig. 5b) that probably include lipid peroxidation products and LPS lipid A now loosely embedded in the disrupted outer membrane (Fig. 2d), and (ii) an increased expression of *ompF* (Fig. 9) that promotes passive uptake of solutes to counteract the significant leakage of intracellular ions. In relation to (i), MV size somewhat increases (Fig. 4a(iii)), which is consistent with the increased waste content to be released towards the extracellular medium. MVs further contribute to mitigate adverse TiO₂NP effects (Fig. 5d) not only via expulsion of wastes generated by the deleterious action of TiO₂NPs (direct effect), but also via their modification of the colloidal stability of TiO₂NPs over time (depending on the concentration of secreted MVs, Fig. 5c) by electrostatically favored interactions between the MVs (negatively charged, Fig. 3 and S2) and TiO₂NPs (positively charged, Supplementary Fig. 1a, b). This results in the formation of large TiO₂NP–MV heteroaggregates and sedimentation thereof in the long term (indirect MV-mediated defense). Regime II is where significant loss of viability (Fig. 2a) is reached despite these established defense strategies.

At higher TiO₂NP concentrations (regime III, >10 mg/L, Fig. 10d), deleterious effects of TiO₂NPs in regime I and II have resulted in major outer membrane disruption, thereby exposing the thin rigid peptidoglycan layer in the periplasm to the outer aqueous environment, in agreement with nanomechanical measurements (Figs. 6 and 8a). In addition, all features delineated in regime II and derived from fluorescence-based assays (Fig. 2) are magnified in regime III, including the *ompF* over-expression (Fig. 9), consistent with a dramatic loss of membrane integrity and hyperproduction of MVs (Fig. 5b). Regime III is also where *osmB*, *otsB*, *oppA*, and *lpxC* genes (Fig. 9) are downregulated,

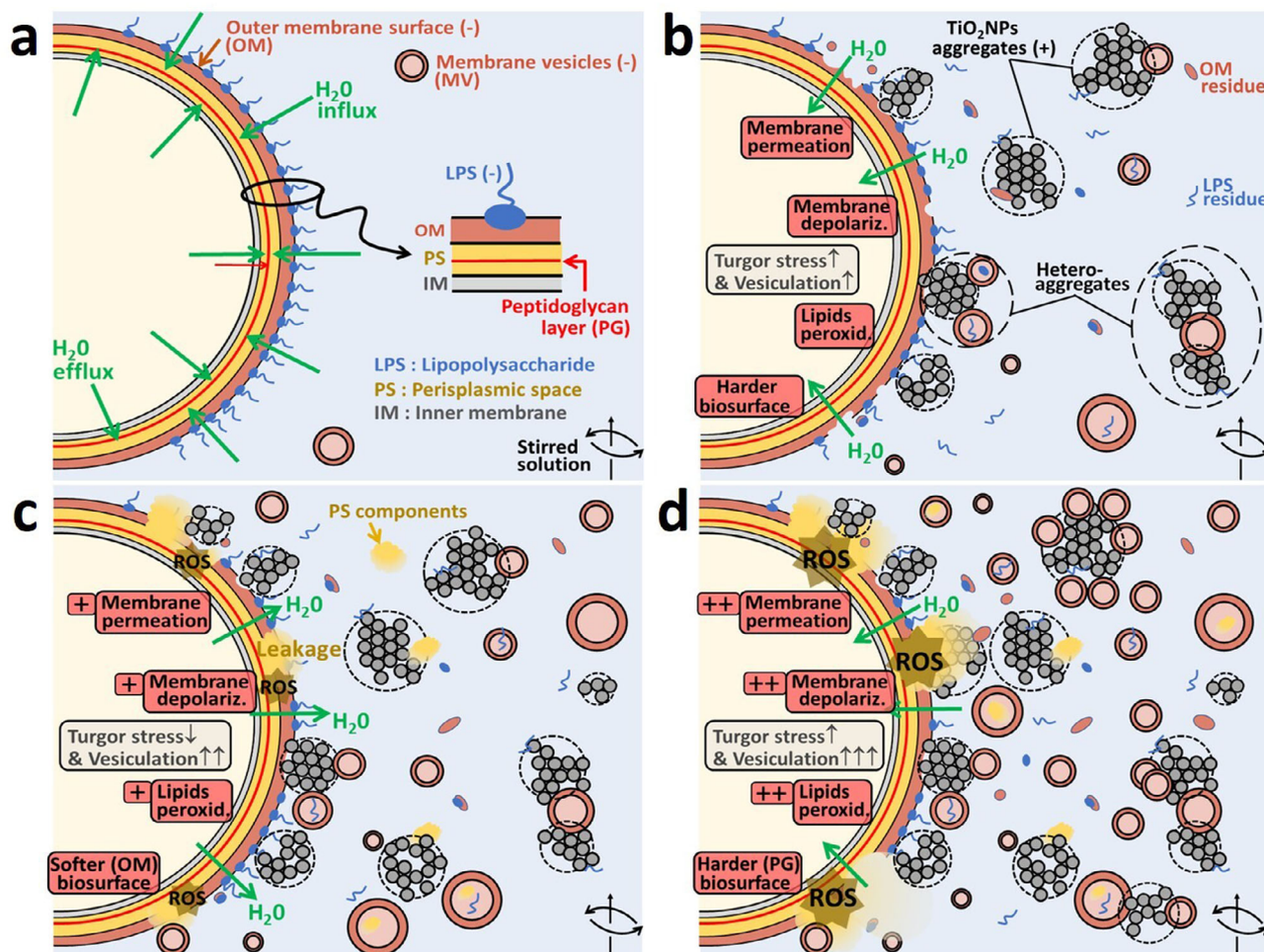


Fig. 10 Schematic representation of TiO_2NP toxicity mechanisms towards the most TiO_2NP -sensitive JW3606 (hep+) mutant with un-truncated inner core LPS. Schematic overview (not to scale) of the modes of action of TiO_2NPs on JW3606 (hep+). **a** Cells unexposed to TiO_2NPs , and osmoregulation (water flux balance in the periplasm). **b** Regime I. **c** Regime II. **d** Regime III. Nomenclature: OM outer membrane, IM inner membrane, PS periplasmic space, PG peptidoglycan layer, MV membrane vesicle, ROS reactive oxygen species. Green arrows indicate the water flux direction. (–) and (+) denote the sign of the charge carried by the OM surface, MVs and TiO_2NPs . From (a) to (d), the TiO_2NP concentration in the exposome gradually increases from 0 to 50 mg/L.

probably due to the associated energy costs required to maintain the corresponding transcription at a stage where the cell viability is minimal. Also, it is in this regime that significant oxidative stress comes into play (Fig. 2 and Supplementary Fig. 3), triggered by osmotic stress^{55–58}.

Concerning JW3596 (hep–), its nanomechanical properties remain practically unmodified in regime α (0–5 mg/L) (Figs. 7, 8b), in line with the absence of significant dependence of osmotic stress-responsive gene expression on TiO_2NP concentration (Fig. 9). This feature is further in accordance with the absence of TiO_2NP -induced lipid peroxidation and oxidative stress, with the preservation of membrane integrity and the maintenance of the membrane potential (Fig. 2). The higher native membrane permeability (i.e., at 0 mg/L TiO_2NPs) of JW3596 (hep–) following inner core LPS truncation, as compared to JW3606 (hep+) (Fig. 2c), and the associated fragilization of the membrane⁴³ offer an efficient way to circumscribe Turgor pressure perturbations caused by TiO_2NPs without the need to significantly modulate the transcription of osmotic stress-responsive genes that are expressed at similar or higher levels compared to JW3606 (hep+) (Fig. 9). In addition, the significant vesiculation of JW3596 (hep–) in the absence of TiO_2NPs (Fig. 5b) confers upon this mutant a more efficient defense, in line with the larger threshold

TiO_2NP concentration (5 mg/L) that marks the onset of significant harmful effects (Fig. 2). It is indeed only at concentrations >5 mg/L (regime β) that cell nanomechanical properties follow the trends discussed in regime I for JW3606 (hep+) (Figs. 7, 8b) and it is not until concentrations become higher than 5–10 mg/L that oxidative stress, membrane integrity loss, increase in membrane permeability, and lipid peroxidation become significant (Fig. 2). Over- and under-expressions of *ompF* and *osmB/osmC*, respectively, are also measured at TiO_2NP concentrations >5 mg/L, as for JW3606 (hep+). In contrast, *otsB*, *oppA*, and *lpxC* expression levels are maintained constant under all tested exposure conditions, which reflects a more favorable energy balance than that for JW3606 (hep+).

In summary, our multiscale approach shows that *rfaG* gene mutation (JW3606 (hep+)) results in a moderate vesiculation capacity and a preserved membrane permeability in the absence of TiO_2NPs . In turn, this dramatically reduces the efficiency of vesiculation and of osmoregulation cell strategies to circumvent the dominant osmotic stress induced by TiO_2NPs at low concentrations. Our results demonstrate that a mutant with *rfaC* gene mutation and LPS truncation (and thus a membrane that is apparently more severely altered) resists the adverse effects of TiO_2NPs in a much more efficient way due to its native

hypervesiculation capability and significant regulatory response to osmotic stress in the absence of TiO₂NPs. These results may shift current practice for fighting against harmful bacteria or preserving the viability of beneficial bacteria that are facing exogenous contaminants. Indeed, controlled stress may initiate cell acquisition of weapons, such as vesicles, and thereby increase the cell defense arsenal against toxic contaminants. On a methodological level, the study introduces multiparametric atomic force microscopy/spectroscopy as a valuable tool to diagnose spatially resolved NP effects on biosurface nanomechanics at the single-cell level. It further integrates electrophoretic cell fingerprints within the context of NP toxicity evaluation at a level that goes beyond the traditional zeta-potential concept that is unapplicable to unravel electrokinetic properties of soft (ion- and flow-permeable) bacterial surfaces^{45–47}. Finally, it succeeds to connect multiscale proxis (from the gene, single-cell to population scale) underpinning the various action modes of NPs (depending on the concentration in the exposome) and the corresponding cells response as derived by transcriptomics, cytometry, electrokinetics, and atomic force spectroscopy. Heterogeneities in cell response to TiO₂NP exposure are further revealed at the gene, single-cell, and population scales.

Methods

Bacterial culture and preparation. *E. coli* strains BW25113 (wild type WT) and the knock-out *rfa*-gene mutants⁴¹ were obtained from the Coli Genetic Stock Center, Yale University. The position of the mutations in the *rfa* operons and the structure of the LPS resulting from these mutations are reported in Fig. 1. Knock-out mutations were checked by PCR before freezing at -80°C in 50% glycerol solution. For experiments, cell cultures were first streaked from frozen stock on Luria Bertani (LB) agar (LB broth containing 1.5 w/v % agar) and incubated at 37°C . Then, 4 mL preculture of M9 medium broth (6 g/L Na₂HPO₄, 3 g/L KH₂PO₄, 1 g/L NH₄Cl, 0.5 g/L NaCl, 1 mM MgSO₄, 0.1 mM CaCl₂, 0.2% Glucose, 10 μg/mL Thiamine, 20 μg/mL Proline, and 25 μg/mL uridine) was inoculated with an isolated colony and incubated overnight at 37°C under stirring. The next day, 100 mL of M9 medium broth were seeded at 1:100 dilution with the precultured cells and further incubated at 37°C , 150 rpm, until the growth exponential phase was reached ($\text{OD}_{600\text{nm}} \sim 0.5$). Cells were subsequently harvested by centrifugation ($5000 \times g$, 5 min), washed twice with 10 mM KNO₃. $\text{OD}_{600\text{nm}}$ of the obtained bacterial suspensions was finally adjusted to 0.5 final value in 10 mM KNO₃. Except for WT, all bacterial cultures were supplemented with kanamycin (30 mg/L) as a selective pressure.

Titanium dioxide nanoparticles. Nanopowder of Aeroxide® TiO₂ P25 was purchased from Evonik Degussa GmbH (Frankfurt, Germany). TiO₂ nanoparticles (TiO₂NPs) display an 80:20 anatase:rutile composition and are defined by a pristine particle radius of 21 nm and a specific surface area of $50 \pm 15 \text{ m}^2/\text{g}$ according to the manufacturer's information. Suspensions of TiO₂NPs were prepared by dispersing 100 mg nanopowder in 10 mL sterile ultrapure water (milli-Q water, 18.2 MΩ cm) and were subsequently probe-sonicated (Sonic Vibra-cell 750 W, Sonics & Materials, frequency 20 kHz, 3 mm micro tip, amplitude 40%) in the dark for 30 min at 4°C to break apart large TiO₂NP aggregates and for homogenization purpose²⁶. The so-prepared stock suspension of TiO₂NPs (10 g/L concentration) was stable against aggregation-sedimentation for a month, and was protected from light.

Bacteria exposure to nanoparticles. TiO₂NP dispersions were prepared with concentrations in the range between 0.1 and 50 mg/L in 20 mL aliquots of bacterial suspensions ($\text{OD}_{600\text{nm}} \sim 0.5$) at 10 mM KNO₃ (pH~5.5–6). Bacteria–TiO₂NP mixtures were subsequently kept at 20°C in the dark under 150 rpm stirring conditions for 20 h. For the sake of comparison, a reference sample containing bacteria unexposed to TiO₂NPs was subjected to the same conditions. After 20 h, samples were analyzed using the procedures detailed in the sections below. In addition, bacteria–TiO₂NPs suspensions were filtered with the use of a sterile vacuum filter bottle system with 0.45 μm or 0.22 μm porosity (Corning, CA membrane) to remove bacteria and TiO₂NP aggregates. These filtrates were analyzed for evaluation of MVs charge/size properties and AFM imaging, as described below. To better identify the roles played by these MVs in mitigating TiO₂NP toxicity, co-incubation experiments combining bacteria, 50 mg/L TiO₂NPs, and filtrates at different dilution ratios in 10 mM KNO₃ were carried out (Fig. 5c, d).

Colony-forming unit (CFU). The viability of bacterial cells exposed to TiO₂NPs was assessed by CFUs per milliliter using the drop-count method²⁶. The bacteria–TiO₂NP mixtures were diluted serially at 1:100 to 1:10⁵. For each dilution condition, nine drops (20 μL per drop) were transferred onto the LB agar medium and

incubated at 37°C for 24 h. The percentage of viable cells was determined by comparing the number of CFUs obtained with exposed and unexposed samples.

Fluorescence measurements. Cells exposed and unexposed to TiO₂NPs were diluted 1:50 in 10 mM KNO₃ and labeled using different fluorescent dyes. DIBAC₄(3) (13.5 μM, 15 min at RT; Sigma Aldrich, Germany) was used to investigate TiO₂NP effects on membrane depolarization, propidium iodide (30 μM, 15 min at RT; Life Technologies, USA) for evaluation of membrane permeability, BODIPY (2.5 μM, 15 min at RT; Life Technologies, USA) for that of lipid peroxidation, H₂DCFDA (2 μM, 15 min at RT; Sigma Aldrich, Germany) to address oxidative stress, and the membrane-selective dyes FM4-64 (5 μg/ml, 15 min at RT; Invitrogen, USA) and Syto9 Green Fluorescent Nucleic Acid Stain (5 μM, 15 min at RT; Life Technologies, USA) for cell numeration. Labeled samples were then analyzed by flow cytometry on a BD Accuri™ C6 and a BD Biosciences (BD Biosciences, New Jersey, USA) equipped with a laser emitting at 488 nm. Forward scatter (FSC), side scatter (SSC), and Syto9 signal on FL1 channel (530 nm) or FM4-64 signal on FL3 channel (LP 670 nm) were used to discriminate bacteria and nanoparticle aggregates from the background, and the trigger was set at 15,000 on FSC. For detection of MVs with the use of FM4-64 labeling, the trigger was set at 1000 on FL3 and 100 on FSC (Fig. 5a). DIBAC₄(3), BODIPY, H₂DCFDA, and Syto9 fluorescence were recorded on the FL1 channel and propidium iodide on the FL2 channel (585 nm). For each sample, at least 20,000 events in the gate corresponding to the bacteria were collected in SSC versus FSC dot plot. The Accuri™ cytometer is equipped with peristaltic pumps that allow sample volume measurement and, therefore, accurate determination of cell concentration. Acquisition and further analysis were performed with BD Accuri™ software (BD Biosciences). Each set of experiments was repeated at least three times to ensure data reproducibility. In addition, due to the detection limit of the flow cytometer (i.e., >200 nm), quantitative fluorescence measurements were performed using a plate-reader fluorometer (SAFAS, Monaco) to determine the relative amount of MVs (Fig. 5b). These experiments were performed on filtrates (see “Bacteria exposition to nanoparticles” subsection above) using a DNA stain-binding assay because it was previously reported that MVs contain DNA⁴⁰. For that purpose, after 20 h exposure, cell suspensions were 0.45 μm filtrated and treated with DNase I (10 U/mL, 20 min at 37°C ; Sigma Aldrich) to remove extracellular DNA associated with MVs, and subsequently stained with Syto9 (5 μM, 30 min at RT). After excitation at 485 nm, the emission at 502 nm was measured on three replicate samples. The amount of MVs in the filtrates, expressed in relative fluorescence units, was determined after subtracting the control (i.e., Syto9 probe alone in 10 mM KNO₃).

Electrokinetics and particle size measurements. After 20 h, samples containing bacteria exposed and unexposed to TiO₂NPs were diluted at 1:10 in ultrapure water, leading to cells suspended in 1 mM KNO₃ background electrolyte. For each TiO₂NP concentration tested, the electrophoretic mobility distributions (electropherograms) of so-prepared bacteria–TiO₂NP suspensions were measured at natural pH and room temperature using a Zetaphoremeter IV (CAD Instrumentations, Les Essarts le Roi, France). Electrophoretic mobility evaluation consisted of following the displacements of particles in a quartz Suprasil® rectangular capillary upon application of a constant direct-current electric field (800 V/m) and particle tracking was monitored by the reflection of a laser beam at 90° angle with the use of a charge-coupled device camera. Trajectories were recorded in real-time and processed by CAD image analysis software to derive electrophoretic mobility distributions. For each tested condition, particle displacements generated by the applied electric field were collected from three replicates with aliquots prepared from a given bacteria–TiO₂NP preparation, and independent sets ($n = 3–14$, Fig. 3) of three measurements were further performed for each condition starting from different cell cultures and cell colonies. Electrophoretic mobility distributions of the filtrates (see “Bacteria exposure to nanoparticles” subsection) were measured following the above procedure (Supplementary Fig. 2). Distributions of the hydrodynamic diameter of the particles dispersed in the filtrates were collected with a Zetasizer NanoZS equipment (Malvern Analytical, He–Ne red laser, 633 nm) by DLS. In detail, particle diffusion coefficients were measured and converted into hydrodynamic size on the basis of the Stokes–Einstein equation. For each tested condition, three measurements were carried out in a row, and independent measurement sets ($n = 2–6$, Fig. 4) of such three measurements were also performed on samples prepared from different cell suspensions issued from distinct colonies. Distributions of hydrodynamic size and electrophoretic mobility pertaining to only the TiO₂NPs in KNO₃ electrolyte solution (pH~5.5) were further measured in 10 mM KNO₃ background electrolyte (Supplementary Fig. 1). While TiO₂NP size measurements were performed following the protocol detailed above, their electrophoretic mobility was measured with Zetasizer NanoZS device by phase analysis light scattering.

Atomic force microscopy (AFM) and force spectroscopy measurements.

Bacteria were deposited onto a cleaned borosilicate glass slide previously covered by a polyethyleneimine layer (Sigma, Mw = 750,000 g/mol) as detailed elsewhere⁴³. A few minutes after cell deposition, the glass slide was rinsed with 1 mM KNO₃ solution to remove unbound bacteria, and the remaining bacteria on the surface were kept in a 1 mM KNO₃ environment (5 ml drop) prior to AFM experiments.

Nanomechanical AFM measurements were performed with a FastScan Dimension Icon and Nanoscope V controller (Bruker) operating in PeakForce Tapping mode at room temperature in 1 mM KNO₃ electrolyte. Adopted AFM probes were NPG Silicon Nitride tips with 20–30 nm curvature radius and a nominal spring constant of 0.24 N/m (0.12–0.48 N/m range) as provided by the manufacturer. Prior to each measurement, a calibration was performed on the rigid substratum to determine the deflection sensitivity (nm/V) of the AFM probe and the cantilever spring constant by the thermal tune method⁶¹, with a resulting value of 0.40 ± 0.2 N/m. Force measurements were recorded during the approach and retraction of the AFM probe to the bacterial surface. The pixel-by-pixel force curves were recorded at the apex of the cell with 500 nm scan size (256×256 local force curve measurements at 1 Hz scan rate and 1 $\mu\text{m/s}$ probe velocity). The setpoint adopted for all force measurements was 5 nN. As previously shown⁴⁶, the liquid environment adopted for the AFM measurements (1 mM KNO₃) allows a proper detection of changes in cell surface structure from modulations of the cell Young modulus (elasticity) and cell Turgor pressure (related to the cell stiffness) derived here by analysis of the approach force curves along the lines detailed elsewhere⁵⁰. Briefly, nanomechanical cell properties (E , k_{cell}) and indentation δ , which marks the transition between the non-linear part of the force-indentation curve and the linear compliance regime, were evaluated on the basis of the Sneddon model corrected for finite cell thickness and Hook's law using a home-made MATLAB program able to handle rapidly the analysis of a large number of force curves (65,536 here per cell examined, with $n = 8$ probed cells issued from similar and different colonies, Fig. 8)⁵⁰. All reported spatial maps are based on only the force curves that were successfully fitted with $R^2 > 0.95$ ⁵⁰, and curves that did not comply with this condition were systematically rejected (white dots in Figs. 6, 7). Cell elasticity and stiffness were derived for TiO₂NP concentrations in the range 0–20 mg/L as measurements at higher concentrations were significantly impaired by AFM probe contamination by TiO₂NPs. AFM imaging of bacteria and filtrates (see “Bacteria exposure to nanoparticles” subsection above) was also performed by PeakForce Tapping mode that best preserves the integrity of fragile biosurfaces upon probe scanning. The first-order estimation of the size distribution of MVs obtained by 0.45 μm -filtration of bacteria–TiO₂NP suspensions and subsequently deposited (50 μl) on a cleaned borosilicate glass slide, was derived (after sample drying) from AFM images collected in air and analyzed with WSXM free software⁶². For that purpose, based on literature results⁴⁰ a minimum cutoff MV diameter of 20 nm was selected and a minimal value was further imposed for the height of the particles to be included in the analysis. As the size evaluation of soft MV particles deposited on a rigid surface and imaged after drying in the air is necessarily approximate due to e.g., capillarity-driven particle deformation, DLS measurements were further conducted to refine MV size estimation in aqueous solution (Fig. 4).

Transcriptomics. Bacteria–TiO₂NP suspensions were centrifuged ($7000 \times g$, 10 min) and pellets were stored at -80°C . RNA extractions were performed using an UltraClean Microbial RNA isolation kit (MOBIO, CA, USA). After extraction, contaminating DNA was digested with DNase I (Sigma Aldrich), and total RNA was purified by phenol/chloroform extraction and ethanol precipitation. RNA quantity and purity were assessed by OD measurements ($\text{OD}_{260\text{nm}}$ and ODs ratio $260\text{nm}/280\text{nm}$ and $260\text{nm}/230\text{nm}$) and RNA integrity was checked using Bioanalyzer 2100 (Agilent, CA, USA). The cDNA was synthesized in a final volume of 20 μl using 550 ng of RNA, 2.5 μM of random hexamer primers, and SuperScript[®] IV reverse transcriptase according to the manufacturer's instructions (Invitrogen). RT-qPCR was performed on 12 selected genes that encode enzymes involved in ROS scavenging and osmotic stress regulatory pathways. Genes and primers are listed in the Supplementary Information, Supplementary Table 1. Primers were designed using Primer3Plus⁶³. The qPCR reaction was conducted with 2 μl of cDNA (30 ng/reaction) as a template, 150 or 250 nM primers and Fast SYBR[®] green master mix (Applied Biosystem[®], CA, USA) in a reaction mixture with a final volume of 20 μl . The cycling conditions were 20 s at 95°C , followed by 40 cycles of 3 s at 95°C and 30 s at 60°C . Amplification efficiencies (between 90 and 110%) of all primers were verified and amplicon sizes were also verified on agarose gel. All PCR amplifications were performed in four biological replicates using the StepOnePlus RT-PCR system (Applied Biosystems[®]). Gene expression levels (Fig. 9—Supplementary Fig. 3) were analyzed using the relative quantification method ($\Delta\Delta\text{Ct}$)⁶⁴. In order to select a suitable reference gene, the stability of five genes was tested on 12 cDNA produced from 12 cell cultures exposed to different concentrations of TiO₂NPs and analyzed with Genorm⁶⁵. *IhfB* and *idt1* were assigned as the most stable genes and these genes have already been used several times as references^{13,66–68}. The ΔCt and the pooled standard deviation were calculated by normalizing the gene of interest Ct value by the geometric mean of the two Ct from reference genes.

Statistics and reproducibility. Data reported in Figs. 2, 8, 9, and Supplementary Fig. 3 were statistically analyzed with R software, version 4.0.3. Data were first tested using the Shapiro–Wilk test for normality and the Bartlett test for homogeneity of variances. Based on the outcome of these tests, we used either a parametric one-way ANOVA followed by Tukey post hoc test or a non-parametric Kruskal–Wallis ANOVA with Dunn post hoc test. Post hoc tests were only performed when the overall ANOVA or Kruskal–Wallis ANOVA revealed overall significance. Statistical results are provided in Supplementary Tables 2–5. Statistical testing of the data was performed by excluding the few outliers of the box plots.

Note that the statistical analysis (i) should be cautiously considered for appreciation of the significance of cytometry and transcriptomic measurements carried out with $n = 3–4$ independent replicates, as commonly done in literature (Figs. 2, 9 and Supplementary Fig. 3) due to the dependence of the test outcome on n , and (ii) does not inform on the significance of the overall data dependence on TiO₂NP concentration (including that of their distribution widths, which reflects multiscale heterogeneities in cell response to TiO₂NP stressors). The numbers of replicates adopted in this work for the various experiments conform to what is classically reported in the literature.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The authors declare that the data supporting the findings of this study are available within the paper and its supplementary information file. All source data underlying the graphs presented in Figs. 1–9 are made available as Supplementary Data with accompanying captions. All other data are available from the corresponding author on reasonable request.

Code availability

All codes used for the analysis of the data within this paper are available upon request to the authors.

Received: 27 November 2020; Accepted: 11 May 2021;

Published online: 03 June 2021

References

- Chen, X. B. & Mao, S. S. Titanium dioxide nanomaterials: synthesis, properties, modifications, and applications. *Chem. Rev.* **107**, 2891–2959 (2007).
- Fu, G., Vary, P. S. & Lin, C.-T. Anatase TiO₂ nanocomposites for antimicrobial coatings. *J. Phys. Chem. B* **109**, 8889–8898 (2005).
- Hou, J. et al. Toxicity and mechanisms of action of titanium dioxide nanoparticles in living organisms. *J. Environ. Sci.* **75**, 40–53 (2019).
- Dasari, T. P., Pathakoti, K. & Hwang, H. Determination of the mechanism of photoinduced toxicity of selected metal oxide nanoparticles (ZnO, CuO, Co₃O₄, and TiO₂) to *E. coli* bacteria. *J. Environ. Sci.* **25**, 882–888 (2013).
- Erdem, A., Metzler, D., Cha, D. K. & Huang, C. P. The short-term toxic effects of TiO₂ nanoparticles toward bacteria through viability, cellular respiration, and lipid peroxidation. *Environ. Sci. Pollut. Res.* **22**, 17917–17924 (2015).
- Lin, X. et al. Toxicity of TiO₂ nanoparticles to *Escherichia coli*: effects of particle size, crystal phase and water chemistry. *PLoS ONE* **9**, e110247 (2014).
- Tong, T., Chu, T. T. B., Kelly, J. J., Gaillard, J. & Gray, K. A. Cytotoxicity of commercial nano-TiO₂ to *Escherichia coli* assessed by high-throughput screening: effects of environmental factors. *Water Res.* **47**, 2352–2362 (2013).
- Nel, A., Xia, T., Mädler, L. & Li, N. Toxic potential of materials at the nanolevel. *Science* **311**, 622–627 (2006).
- Warheit, D. B. Debunking some misconceptions about nanotoxicology. *Nano Lett.* **10**, 4777–4782 (2010).
- Martyniuk, C. J. & Simmons, D. B. Spotlight on environmental omics and toxicology: a long way in a short time. *Comp. Biochem. Physiol. Part D. Genomics Proteom.* **19**, 97–101 (2016).
- Léonard, L., Chibane, L. B., Bouhedda, B. O., Degraeve, P. & Oulahal, N. Recent advances on multi-parameter flow cytometry to characterize antimicrobial treatments. *Front. Microbiol.* **7**, 1225 (2016).
- Nosaka, Y. & Nosaka, A. Y. Generation and detection of reactive oxygen species in photocatalysis. *Chem. Rev.* **117**, 11302–11336 (2017).
- Sohm, B., Immel, F., Bauda, P. & Pagnout, C. Insight into the primary mode of action of TiO₂ nanoparticles on *Escherichia coli* in the dark. *Proteomics* **15**, 98–113 (2015).
- Carré, G. et al. On the use of capillary cytometry for assessing the bactericidal effect of TiO₂. Identification and involvement of reactive oxygen species. *Photochem. Photobiol. Sci.* **12**, 610–620 (2013).
- Ranjan, S. & Ramalingam, C. Titanium dioxide nanoparticles induce bacterial membrane rupture by reactive oxygen species generation. *Environ. Chem. Lett.* **14**, 487–494 (2016).
- Dalai, S., Pakrashi, S., Kumar, R. S. S., Chandrasekaran, N. & Mukherjee, A. A comparative cytotoxicity study of TiO₂ nanoparticles under light and dark conditions at low exposure concentrations. *Toxicol. Res.* **1**, 116–130 (2012).
- Pathakoti, K., Manubolu, M. & Hwang, H.-m. Mechanistic insights into TiO₂ and ZnO nanoparticle-induced metabolic changes in *Escherichia coli* under solar simulated light irradiation. *Water Air Soil Poll. Res.* **231**, 16 (2020).

18. Cho, M., Chung, H., Choi, W. & Yoon, J. Linear correlation between inactivation of *E. coli* and OH radical concentration in TiO₂ photocatalytic disinfection. *Water Res.* **38**, 1069–1077 (2004).
19. Kiwi, J. & Nadochenko, V. Evidence for the mechanism of photocatalytic degradation of the bacterial wall membrane at the TiO₂ interface by ATR-FTIR and laser kinetic spectroscopy. *Langmuir* **21**, 4631–4641 (2005).
20. Nadochenko, V. A., Rincon, A. G., Stanca, S. E. & Kiwi, J. Dynamics of *E. coli* membrane cell peroxidation during TiO₂ photocatalysis studied by ATR-FTIR spectroscopy and AFM microscopy. *J. Photochem. Photobiol. A* **169**, 131–137 (2005).
21. Li, Y., Zhang, W., Niu, J. & Chen, Y. Mechanism of photogenerated reactive oxygen species and correlation with the antibacterial properties of engineered metal-oxide nanoparticles. *ACS Nano* **6**, 5164–5173 (2012).
22. Erdem, A., Metzler, D., Cha, D. & Huang, C. P. Inhibition of bacteria by photocatalytic nano-TiO₂ particles in the absence of light. *Int. J. Environ. Sci. Technol.* **12**, 2987–2996 (2015).
23. Kumar, A., Pandey, A. K., Singh, S. S., Shanker, R. & Dhawan, A. Engineered ZnO and TiO₂ nanoparticles induce oxidative stress and DNA damage leading to reduced viability of *Escherichia coli*. *Free Radic. Biol. Med.* **51**, 1872–1881 (2011).
24. Djurišić, A. B. et al. Toxicity of metal oxide nanoparticles: mechanisms, characterization, and avoiding experimental artefacts. *Small* **11**, 26–44 (2015).
25. Carré, G. et al. TiO₂ photocatalysis damages lipids and proteins in *Escherichia coli*. *Appl. Environ. Microbiol.* **80**, 2573–2581 (2014).
26. Pagnout, C. et al. Role of electrostatic interactions in the toxicity of titanium dioxide nanoparticles toward *Escherichia coli*. *Colloids Surf. B* **92**, 315–321 (2012).
27. Zhukova, L. V., Kiwi, J. & Nikandrov, V. V. TiO₂ nanoparticles suppress *Escherichia coli* cell division in the absence of UV irradiation in acidic conditions. *Colloids Surf. B* **97**, 240–247 (2012).
28. Adams, L. K., Lyon, D. Y. & Alvarez, P. J. J. Comparative eco-toxicity of nanoscale TiO₂, SiO₂, and ZnO water suspensions. *Water Res.* **40**, 3527–3532 (2006).
29. Leung, Y. H. et al. Toxicity of ZnO and TiO₂ to *Escherichia coli* cells. *Sci. Rep.* **6**, 35243 (2016).
30. Applerot, G. et al. Enhanced antibacterial activity of nanocrystalline ZnO due to increased ROS-mediated cell injury. *Adv. Funct. Mater.* **19**, 842–852 (2009).
31. Leung, Y. H. et al. Mechanisms of antibacterial activity of MgO: non-ROS mediated toxicity of MgO nanoparticles towards *Escherichia coli*. *Small* **10**, 1171–1183 (2014).
32. Leung, Y. H. et al. Antibacterial activity of ZnO nanoparticles with a modified surface under ambient illumination. *Nanotechnology* **23**, 475703 (2012).
33. Angel, B. M., Vallotton, P. & Apte, S. C. On the mechanism of nanoparticulate CeO₂ toxicity to freshwater algae. *Aquat. Toxicol.* **168**, 90–97 (2015).
34. Lyon, D. Y., Brunet, L., Hinkal, G. W., Wiesner, M. R. & Alvarez, P. J. J. Antibacterial activity of fullerene water suspensions (nC60) is not due to ROS-mediated damage. *Nano Lett.* **8**, 1539–1543 (2008).
35. Burello, E. & Worth, A. P. A theoretical framework for predicting the oxidative stress potential of oxide nanoparticles. *Nanotoxicology* **5**, 228–235 (2011).
36. Gali, N. K., Ning, Z., Daoud, W. & Brimblecombe, P. Investigation on the mechanism of non-photocatalytically TiO₂-induced reactive oxygen species and its significance on cell cycle and morphology. *J. Appl. Toxicol.* **36**, 1355–1363 (2016).
37. Fenoglio, I., Greco, G., Livraghi, S. & Fubini, B. Non-UV-induced radical reactions at the surface of TiO₂ nanoparticles that may trigger toxic responses. *Chemistry-Eur. J.* **15**, 4614–4621 (2009).
38. Buchman, J. T. et al. Using an environmentally-relevant panel of Gram-negative bacteria to assess the toxicity of polyallylamine hydrochloride-wrapped gold nanoparticles. *Environ. Sci.: Nano* **5**, 279–288 (2018).
39. Macdonald, I. A. & Kuehn, M. J. Stress-induced outer membrane vesicle production by *Pseudomonas aeruginosa*. *J. Bacteriol.* **195**, 2971–2981 (2013).
40. Schwachheimer, C. & Kuehn, M. J. Outer-membrane vesicles from Gram-negative bacteria: biogenesis and functions. *Nat. Rev. Microbiol.* **13**, 605–619 (2015).
41. Yethon, J. A., Vinogradov, E., Perry, M. B. & Whitfield, C. Mutation of the lipopolysaccharide core glycosyltransferase encoded by waaG destabilizes the outer membrane of *Escherichia coli* by interfering with core phosphorylation. *J. Bacteriol.* **182**, 5620–5623 (2000).
42. Nikaido, H. Molecular basis of bacterial outer membrane permeability revisited. *Microbiol. Mol. Biol. Rev.* **67**, 593–656 (2003).
43. Pagnout, C. et al. Pleiotropic effects of rfa-gene mutations on *Escherichia coli* envelope properties. *Sci. Rep.* **9**, 9696 (2019).
44. Krapf, M.-E. et al. Polyethyleneimine-mediated flocculation of *Shewanella oneidensis* MR-1: impacts of cell surface appendage and polymer concentration. *Water Res.* **46**, 1838–1846 (2012).
45. Duval, J. F. L. & Gaboriaud, F. Progress in electrohydrodynamics of soft microbial particle interphases. *Curr. Opin. Colloid Interface Sci.* **15**, 184–195 (2010).
46. Francius, G. et al. Bacterial surface appendages strongly impact nanomechanical and electrokinetic properties of *Escherichia coli* cells subjected to osmotic stress. *PLoS ONE* **6**, e20066 (2011).
47. Gaboriaud, F., Gee, M. L., Strungnell, R. & Duval, J. F. L. Coupled electrostatic, hydrodynamic and mechanical properties of bacterial interfaces in aqueous media. *Langmuir* **24**, 10988–10995 (2008).
48. Nakao, R., Ramstedt, M., Nyunt Wai, S. & Eric Uhlén, B. Enhanced biofilm formation by *Escherichia coli* LPS mutants defective in Hep biosynthesis. *PLoS ONE* **7**, e51241 (2012).
49. Kulp, A. J. et al. Genome-wide assessment of outer membrane vesicle production in *Escherichia coli*. *PLoS ONE* **10**, e0139200 (2015).
50. Offroy, M., Razafitianamaharavo, A., Beaussart, A., Pagnout, C. & Duval, J. F. L. Fast automated processing of AFM PeakForce curves to evaluate spatially resolved Young modulus and stiffness of turgescient cells. *RSC Adv.* **10**, 19258–19275 (2020).
51. Mathelié-Guinlet, M. et al. Detrimental impact of silica nanoparticles on the nanomechanical properties of *Escherichia coli*, studied by AFM. *J. Colloid Interface Sci.* **529**, 53–64 (2018).
52. Lins, R. D. & Straatsma, T. P. Computer simulation of the rough lipopolysaccharide membrane of *Pseudomonas aeruginosa*. *Biophys. J.* **81**, 1037–1046 (2001).
53. EcoCyc E. coli Database <https://ecocyc.org/>
54. Graeme-Cook, K. A. The regulation of porin expression in *Escherichia coli*: effect of Turgor stress. *FEMS Microbiol. Lett.* **79**, 219–224 (1991).
55. Bojanović, K., D'Arrigo, I. & Long, K. S. Global transcriptional responses to osmotic, oxidative, and imipenem stress conditions in *Pseudomonas putida*. *Appl. Environ. Microbiol.* **83**, e03236–16 (2017).
56. Mager, W. H., de Boer, A. H., Siderius, M. H. & Voss, H. P. Cellular responses to oxidative and osmotic stress. *Cell Stress Chaperones* **5**, 73–75 (2000).
57. Gunasekera, T. S., Csonka, L. N. & Paliy, O. Genome-wide transcriptional responses of *Escherichia coli* K-12 to continuous osmotic and heat stresses. *J. Bacteriol.* **190**, 3712–3720 (2008).
58. Smirnova, G. V., Muzyka, N. G. & Oktyabrsky, O. N. The role of antioxidant enzymes in response of *Escherichia coli* to osmotic upshift. *FEMS Microbiol. Lett.* **186**, 209–213 (2000).
59. Schwachheimer, C., Kulp, A. & Kuehn, M. J. Modulation of bacterial outer membrane vesicle production by envelope structure and content. *BMC Microbiol.* **14**, 324 (2014).
60. Kulkarni, H. M., Nagaraj, R. & Jagannadham, M. V. Protective role of *E. coli* outer membrane vesicles against antibiotics. *Microbiol. Res.* **181**, 1–7 (2015).
61. Butt, H.-J. & Jascke, M. Calculation of thermal noise in atomic force microscopy. *Nanotechnology* **6**, 1–7 (1995).
62. Horcas, I. et al. WSXM: a software for scanning probe microscopy and a tool for nanotechnology. *Rev. Sci. Instrum.* **78**, 013705 (2007).
63. Untergasser, A. et al. Primer3Plus, an enhanced web interface to Primer3. *Nucleic Acids Res.* **35**, W71–W74 (2007).
64. Livak, K. J. & Schmittgen, T. D. Analysis of relative gene expression data using real-time quantitative PCR and the 2^{-(delta delta C(T))} method. *Methods* **25**, 402–408 (2001).
65. Vandesompele, J. et al. Accurate normalization of real-time quantitative RT-PCR data by geometric averaging of multiple internal control genes. *Genome Biol.* **3**, research0034.1–0034.11 (2002).
66. Caspeta, L., Flores, N., Perez, N. O., Bolivar, F. & Ramirez, O. T. The effect of heating rate on *Escherichia coli* metabolism, physiological stress, transcriptional response, and production of temperature-induced recombinant protein: a scale-down study. *Biotechnol. Bioeng.* **102**, 468–482 (2009).
67. Baez, A., Flores, N., Bolivar, F. & Ramirez, O. T. Metabolic and transcriptional response of recombinant *Escherichia coli* to elevated dissolved carbon dioxide concentrations. *Biotechnol. Bioeng.* **104**, 102–110 (2009).
68. Zhou, K. et al. Novel reference genes for quantifying transcriptional responses of *Escherichia coli* to protein overexpression by quantitative PCR. *BMC Mol. Biol.* **12**, 18 (2011).

Acknowledgements

J.F.L.D. and C.P. acknowledge funding of this work by la Fondation Rovaltain (<https://fcsrovaltain.org/>), France (EVAMINTOX project). J.F.L.D. acknowledges Héloïse Gendre for preliminary electrokinetic and AFM measurements, and J.F.L.D. thanks Prof. Raewyn M. Town (University of Antwerp, Antwerp, Belgium) for some final language corrections.

Author contributions

J.F.L.D. and C.P. designed the study and interpreted data. J.F.L.D. wrote the manuscript. C.P., A.R., B.S., C.C., A.B., E.D., I.B., M.O., and J.F.L.D. contributed to the acquisition, analysis, and interpretation of the data. C.P., A.R., B.S., C.C., A.B., E.D., I.B., and M.O. revised the paper and all authors approved the final version.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s42003-021-02213-y>.

Correspondence and requests for materials should be addressed to J.F.L.D.

Reprints and permission information is available at <http://www.nature.com/reprints>

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2021