



Published in final edited form as:

Nat Microbiol. ; 1(11): 16186. doi:10.1038/nmicrobiol.2016.186.

Optical and force nanoscopy in microbiology

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Abstract

Microbial cells have developed sophisticated multicomponent structures and machineries to govern basic cellular processes, such as chromosome segregation, gene expression, cell division, mechanosensing, cell adhesion and biofilm formation. Because of the small cell sizes, subcellular structures have long been difficult to visualize using diffraction-limited light microscopy. During the last three decades, optical and force nanoscopy techniques have been developed to probe intracellular and extracellular structures with unprecedented resolutions, enabling researchers to study their organization, dynamics and interactions in individual cells, at the single-molecule level, from the inside out, and all the way up to cell–cell interactions in microbial communities. In this Review, we discuss the principles, advantages and limitations of the main optical and force nanoscopy techniques available in microbiology, and we highlight some outstanding questions that these new tools may help to answer.

The functioning of a microbial cell relies on a complex set of multicomponent cellular structures. These structures often adopt specific shapes, reside in specific subcellular spaces, and carry out functions specific to these properties. Examples can be found both inside and outside the cell. Inside the cell, large assemblies formed by bacterial cytoskeletal proteins play important roles in cell division, chromosome segregation and cell shape maintenance¹. On the outside of the cell, the macromolecular cell wall structure forms a barrier to separate the inner cell from its environment and fulfils essential functions such as defining cell shape, cell motility and cell adhesion processes^{2,3}.

A current challenge in cellular and molecular microbiology is to understand how these cellular structures and machineries are assembled at the correct time and space to achieve their functions. Clearly, microscopy is one powerful method to investigate these cellular structures. However, the small sizes of bacterial cells, often in the range of $\sim 1 \mu\text{m}$, are not much larger than that allowed by diffraction in conventional light microscopy ($\sim 200\text{--}300$

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Competing interests

The authors declare no competing financial interest.

nm), making it impossible to visualize these structures, or to assess their assembly dynamics and interactions with other cellular partners, in nanoscopic details. Using much shorter wavelengths as in electron microscopy (EM) methods, such as electron cryo-tomography (ECT), provides views of microbial structures with much higher resolution, but requires substantial instrumentation and cannot be carried out in live cells.

Since the late 1980s, scientists have attempted to break the diffraction barrier in light microscopy. One branch of efforts concentrated on the 'near-field', in which a tiny probe is placed near the sample and scanned, thus providing images of surfaces with a resolution that is no longer limited by diffraction. Among near-field microscopy methods, atomic force microscopy (AFM), also referred to as force nanoscopy, is particularly well suited for microbiology as it can typically reach resolutions of a few nanometres for surface components of living cells. Another branch of efforts focused on the diffraction-limited 'far-field', in which super-resolution microscopy, also called optical nanoscopy, can probe the localization and motion of single molecules in live cells with a resolution in the range of a few tens of nanometres. These nanoscopy techniques, when further combined with biochemical and genetic interrogations, have profoundly impacted our perception of microbial structures, enabling us for the first time to analyse the organization, dynamics and interactions of cellular structures and machineries at the single molecule level, throughout the cell, and further up to intercellular interactions in microbial communities. In this Review, we describe the main principles of force and optical nanoscopy modalities currently available and discuss distinct advantages and limitations of each. We then survey recent breakthroughs that have been made possible using these new tools, going from the interior of the cell to the cell surface.

Principles of nanoscopy methods

Currently there is a large number of different nanoscopy methods available for imaging and probing small microbial cells. While each comes with different acronyms and varied detection schemes, there are common operational principles as detailed below.

Force nanoscopy

Invented in 1986, AFM was initially designed for imaging surfaces. The key breakthrough was the realization that samples can be imaged at atomic or molecular resolution, without using an incident beam of photons or electrons, but by measuring the near-field interaction between a sharp tip and the surface⁴. The tip is brought in contact with the specimen and raster scans the surface while sensing the minute forces (in the piconewton range) acting on the tip. A piezoelectric scanner moves the tip (or sample) in 3D with high accuracy. The tip is mounted on a soft cantilever made of silicon or silicon nitride. The deformation of the cantilever is measured with a laser beam, enabling the determination of the force on the tip. AFM can work in various environments, including aqueous buffers, thus making it appropriate to analyse biological systems under physiological conditions without the need for labelling or fixation.

AFM is commonly used for imaging the topography of biological structures, with a resolution hundreds of times better than that allowed by the optical diffraction limit. As the

technique shows exquisite sensitivity for surfaces, it is ideally suited for imaging membranes and cell walls^{5,6} (Fig. 1, top). Several imaging modes may be used, which essentially differ in the way the tip is scanned on the surface. In the contact mode, the tip always touches the sample surface and is scanned at a constant applied force. In the dynamic (or intermittent) mode, the tip is oscillated to lower the forces during imaging, thereby minimizing deformation and alteration of the biological specimen. An important issue for reliable imaging of microbiological structures is sample preparation. While isolated structures such as membranes and cell walls can be readily imaged following adsorption on mica, whole cells need to be firmly immobilized on a solid substrate, for instance by attachment onto glass slides coated with positively charged macromolecules like polylysine or by mechanical trapping into porous polymer membranes⁵. Advances in instrumentation, data recording and interpretation, and sample preparation, have enabled researchers to routinely capture molecular details of microbial structures^{5,7}. As images are recorded in buffer and in real-time, it is possible to track dynamic changes during processes like cell growth or interaction with antibiotics⁵. While conventional AFMs have a rather slow time resolution (one image per min), high-speed AFMs (HS-AFMs) have recently been developed, in which the use of small cantilevers and improved electronics enables imaging biosystems with millisecond resolution⁸. This has opened new vistas for studying physiologically relevant molecular and cellular dynamics⁸.

Besides imaging, AFM may also be used as an ultrasensitive force probe, yielding direct information on the localization, adhesion, elasticity and interactions of individual molecules^{9,10} (Fig. 1, top). In this mode, called force spectroscopy, interaction forces are measured by recording the deflection of the cantilever while the tip is moved up and down, thus yielding a force–distance curve. Furthermore, acquiring spatially resolved force–distance curves makes it possible to map interactions and properties of surfaces⁹. Mapping and functionally analysing single molecules using so-called single-molecule force spectroscopy (SMFS) requires labelling the tip with specific ligands, like antibodies or lectins, and then measuring the characteristic adhesion force between the ligand and its receptor^{9,11}. Biofunctionalization of the tip is achieved using crosslinker molecules that anchor ligands firmly at low density, while maintaining their mobility and functionality. In microbiology, SMFS can help us understand how cell surface proteins such as adhesins assemble into nanodomains on the surface of living cells^{12,13}. SMFS may also be exploited to pull on single molecules in order to learn about their elasticity, a property that plays an important role in cell behaviour^{13,14}. These single-molecule manipulations have shown that microbial molecules and structures feature unanticipated mechanical responses when subjected to force (for example, protein unfolding and unzipping, pilus extension and spring properties), which contribute significantly to cellular functions such as mechanosensing and adhesion.

Single-cell force spectroscopy (SCFS), a variation of SMFS, measures forces between a single cell and a target surface (Fig. 1, top)¹⁵. This modality is highly useful to understand the extent to which single-molecule properties and interactions contribute to the behaviour of whole cells. Here, a key issue is to attach a single cell to the cantilever while at the same time ensuring that cell viability and functionality are maintained. An assay was recently proposed for preparing single microbial probes by using colloidal cantilevers and a bio-

inspired polydopamine wet adhesive^{16,17}. The method is versatile, nondestructive and provides precise control of cell positioning, therefore enabling reliable single-cell analysis. Today, the combination of SMFS and SCFS is increasingly used in microbiology to decipher forces driving microorganism–microorganism, microorganism–host, and microorganism–substrate interactions. Importantly, dissecting the molecular nature of measured forces generally requires integrating AFM with modern tools of molecular genetics (mutant strains deficient for cell wall constituents, expression of specific proteins and their mutants in cell display models).

There is much interest in correlating the nanostructure of microbial systems with their chemical and biophysical properties. Multiparametric imaging is a new force-spectroscopy-based AFM modality that can simultaneously image the structure of a sample and quantify its properties and interactions, at the speed of conventional imaging (~30 sec per image)¹⁸. This technology has been applied to the native purple membrane from *Halobacterium salinarum*, which is composed of the light-driven proton-pump bacteriorhodopsin^{19,20}. The adhesion and elasticity of native bacteriorhodopsins were observed at a resolution of a few nanometres¹⁹, and the flexibility of individual proteins was mapped and correlated with the protein crystal structure²⁰. Applied to living microbial cells, multiparametric imaging has enabled observation of single filamentous bacteriophages extruding from living bacteria²¹, mapping adhesive nanodomains on fungal pathogens²², and unravelling structural and physical dynamics of the *Staphylococcus aureus* cell wall²³ and extracellular matrix substances²⁴. As such, this novel modality provides a means of relating surface structure with biophysical properties, which is essential to understand cell surface functions. Despite the numerous advantages of force nanoscopy highlighted above, there are also a number of limitations that are important to keep in mind (Box 1).

Optical nanoscopy

To probe the inside of a cell without being invasive, fluorescence light microscopy is ideal. The ultimate goal of resolving a fluorescent cellular structure with nanoscopic details is to obtain the spatial coordinates of all molecules making up the structure with nanometre precision.

Stimulated emission depletion (STED) microscopy overcomes the diffraction barrier by optically confining the excitation beam in a confocal microscope to a spot smaller than a diffraction-limited area²⁵. The confinement can be achieved by overlaying two beams: an excitation beam to induce fluorescence; and a donut-shaped STED beam to inhibit fluorescence at the outer rim of the excitation beam (Fig. 1, bottom). As such, only molecules at the centre of the excitation beam, that is, where the inhibition STED beam intensity is zero, are allowed to fluoresce, hence effectively shrinking the point spread function (PSF) of the fluorescent spot. The two overlaid beams then scan point-to-point across a sample the same way as that in a confocal laser scanning microscope to generate a super-resolution image²⁶. Thus, the spatial resolution is determined by the size of the intensity-zero centre of the STED beam, and the temporal resolution is determined by the scanning speed. Currently, STED can routinely reach a spatial resolution of ~40–80 nm with a frame rate of ~1 s⁻¹, but a recent development using an ultrafast electro-optical scanning

technique achieved an impressive 5–10 ms per frame with ~70 nm resolution²⁷. STED is also naturally capable of 3D imaging with an axial resolution of up to 40 nm because of its confocal-like setup²⁸.

In wide-field microscopy, one commonly used optical nanoscopy is single-molecule localization-based microscopy (SMLM) such as PALM (photoactivated localization microscopy), STORM (stochastic optical reconstruction microscopy) and other derivatives^{29–32}. The basic concept is that if a particular feature of a molecule can be isolated from surrounding molecules in the same focal volume, and used to determine the molecule's position with greater spatial precision than the focal volume, each individual molecule's coordinates can then be sequentially determined and finally superimposed to reconstruct a super-resolved image (Fig. 1, bottom). To realize this concept, two key components, single-molecule detection and photoswitchable fluorophores, are required. Single-molecule detection allows the localization of the centroid position of a single molecule's PSF with a few nanometre's precision, effectively 'breaking' the diffraction barrier^{33,34}. Photoswitchable fluorophores allow the stochastic activation of individual fluorophores one at a time within a diffraction-limited area by a low dosage of the activation light, effectively isolating single fluorophores from the others in the same area³⁵. Along the axial direction, sub-diffraction-limit resolution can also be achieved by a variety of different methods^{36–38}. For example, a simple setup using cylindrical-lens-induced astigmatism can achieve an axial resolution of ~80 nm (ref.³⁹), and the two-objective, interference-based iPALM (interferometric PALM) can reach ~15 nm axial resolution⁴⁰.

The spatial resolution of SMLM is limited by the following three factors: (1) the precision in localizing single fluorophores, which is mainly influenced by the number of photons a fluorophore can emit³⁴; (2) labelling density (ρ) of the cellular structure, or sampling frequency, dictated by the Nyquist criterion, $d = 2/\rho$ for 2D imaging⁴¹; and (3) the spread of repeat localizations of same molecules^{42,43}. When the first two factors are not limiting, the spread of repeat localizations of same molecules determines the actual spatial resolution in SMLM imaging. Currently SMLM can routinely reach a spatial resolution of ~30 nm in 2D, but the temporal resolution is low because often tens of thousands of images are required to reconstruct one super-resolution image. However, in a recent study, by using a fast CMOS (complementary metal-oxide-semiconductor) camera and a combination of noise-minimizing strategies, an impressive frame rate of up to 32 super-resolution images per second has been achieved in both fixed and live mammalian cells⁴⁴, raising the promise of SMLM's application to microbiology.

A different concept of wide-field super-resolution imaging, structured illumination microscopy^{45,46} (SIM, Fig. 1, bottom), utilizes a predetermined, sinusoidal illumination pattern with alternating maxima and minima to excite a fluorescently labelled cellular structure. The emitted fluorescence from the structure produces the so-called moiré fringes, which are the product of both the structure pattern and the illumination pattern. Moiré fringes contain fine details of the underlying cellular structure, and are coarse enough to be easily visualized by conventional fluorescence microscopy. Thus, by applying multiple (>3, often 10–15) illumination patterns with different orientation and phases to the same sample, information of the underlying cellular structure can be mathematically extracted⁴⁵. The

spatial resolution of SIM is determined by the spatial frequency of the spacing of the illumination pattern. In the linear excitation regime the spatial resolution in 2D can only be improved by a factor of two to ~120 nm. In 3D, SIM can reach an axial resolution of ~300 nm by applying a spatial illumination pattern where out-of-focus light is not detected due to interference⁴⁶. However, because only ~10–15 illumination patterns are needed to reconstruct a super-resolution image, SIM can achieve a fast frame rate of less than 1 s⁻¹. Thus, it is possible to probe the dynamics of cellular structures in super-resolution even in fast growing bacterial cells. In addition, SIM does not require special photoswitchable fluorophores and is fully compatible with regular fluorescent proteins. Because of these properties, when temporal resolution is the primary concern, SIM is often the best choice. Recent employment of nonlinearity to generate high frequency harmonics such as that in saturated SIM (SSIM)^{47,48} can further push the spatial resolution to ~50 nm, hence holding promise for achieving both high temporal and spatial resolution.

While different in formats, all optical nanoscopy methods such as STED, SMLM, and SIM rely on one central principle to achieve sub-diffraction-limit resolution. That is, by switching on and off the fluorescence of a subpopulation of molecules within a diffraction-limited area, molecules that are spatially indistinguishable from each other can be separated in time. This key principle of temporal separation by switching on and off fluorescence unifies all optical nanoscopy methods⁴⁹. Variations of the key concept led to the development of a diverse set of optical nanoscopy methods with overly abundant acronyms. For example, turning molecules off by ground state depletion (GSD) instead of stimulated emission led to GSD microscopy⁵⁰; using reversibly photoswitchable fluorophores to reduce the depletion beam intensity substantially for less photo-damage of the sample led to RESOLFT (ref.⁵¹). Using fluorescence change of fluorophores upon binding and unbinding to DNA or membrane led to PAINT (points accumulation for imaging in nanoscale topography^{52,53}) or BALM (binding activated localization microscopy)⁵⁴. Readers are referred to excellent articles for in-depth discussions of operational principles of these methods⁵⁵, selection of fluorophores^{56,57}, practical concerns and applications^{58,59}, and quantitative analyses^{60,61}. In Box 1 we summarize the major limitations of optical nanoscopy.

Applications in microbiology

By employing new physical principles and imaging technologies, force and optical nanoscopy techniques have provided novel insights into the organization, dynamics and functions of the microbial cell. Here, we survey some of the new biological insights learned from various nanoscopy methods in microbiology and discuss potential new directions, going through cellular structures from the inside to the outside of the cell.

Spatial organization of the nucleoid

Genetic and biochemical studies have shown that the bacterial chromosome is spatially organized into segregated, activity-insulated domains⁶². Optical nanoscopy has complemented population-based studies by directly visualizing the overall morphology and cellular positions of the nucleoid at the single-cell level (Fig. 2).

In fixed cells, using the reversible binding and unbinding of PicoGreen, a dye that increases its fluorescence ~1000-fold upon its binding to the major groove of double-stranded DNA (dsDNA), fine fibres and small voids inside the nucleoid were observed that were not discernible in conventional light microscopy⁵⁴ (Fig. 2a). In other studies (Fig. 2b), chromosomal DNA was directly labelled by incorporating a thymidine analogue EdU (5-ethynyl-2'-deoxyuridine) during DNA replication, which is then conjugated with a photoswitchable dye⁶³⁻⁶⁵ by performing the 'click' reaction⁶⁶. The use of a bright organic dye such as Alexa647 led to much-improved resolution, revealing a cycle of decondensation and recondensation of the nucleoid upon replication initiation and completion, and previously unseen hetero-structures of the nucleoid with regions connected by thin fibres^{63,65} (Fig. 2b). In live cells, DAPI (4',6-diamidino-2-phenylindole), another dsDNA intercalating dye that can diffuse through bacterial membrane, has been used in SIM to visualize the nucleoid structure in both *Bacillus subtilis*⁶⁷ (Fig. 2c) and *Escherichia coli* cells⁶⁸. The ~2-fold increase of resolution in SIM is enough to visualize high-density chromosomal regions (HDRs) heterogeneously distributed along the long axis of *B. subtilis* cells (Fig. 2c) and also *E. coli* cells grown in rich medium^{67,68}.

Alternatively, researchers have fused non-specific DNA-binding proteins (commonly nucleoid-associated proteins, NAPs) to photoactivatable fluorescent proteins (PA-FPs) and visualized their spatial distributions by SMLM as a proxy for the underlying nucleoid organization. However, in both *E. coli* and *Caulobacter crescentus*, all NAPs investigated so far showed rather dispersed, uniform distribution with no distinct organization patterns (Fig. 2d,e)^{69,70}. This discrepancy is likely due to differences of the nucleoid organization and dynamics between live and fixed cells, growth conditions, or bacterial species. Further investigations, combined with mutational and functional analyses, will likely provide new insight into the roles of these NAPs in organizing the nucleoid.

Labelling of the nucleoid as described permits the imaging of the global nucleoid structure in super-resolution, but is not sequence-specific. FROS (Fluorescent repressor operator system⁷¹) uses the binding of a repressor protein such as LacI or TetR fused with a FP to tandem arrays of operator sites to label chromosomal DNA segment specifically. A recent study modified the FROS method and achieved a precision of ~15 nm in determining the cellular position of labelled DNA sites⁷², allowing looped and unlooped conformations of a short (~2 Kbps) chromosomal DNA segment to be distinguished in live *E. coli* cells⁷². If further scaled up to allow systematic, multiplexed sequence-specific imaging, this method may make it possible to investigate how the spatial organization of the nucleoid is coupled to specific gene expression activities.

Distribution and dynamics of DNA-associated machineries

Many DNA-associated molecular machineries exhibit complex spatial distributions and dynamics on the nucleoid. Here optical nanoscopy, in particular SMLM imaging, is often complemented by single-molecule tracking (SMT) to probe where these proteins localize to, how fast they are moving, and how they change in spatial distribution and mobility in response to different cellular signals or genetic mutations (Fig. 3). From these measurements

it is possible to deduce reaction kinetics and dissect molecular mechanisms of these machineries at the single-molecule level in live cells.

Using two-colour PALM imaging and SMT, *B. subtilis* MutS, an enzyme involved in DNA mismatch repair⁷³, was observed to sample rapidly the entire nucleoid, but dwell at the replisome region (Fig. 3a,f). Importantly, MutS was found only to identify errors in newly replicated DNA through its transient association with the replisome. Using a similar strategy, the spatial search and repair pattern of DNA polymerase I and ligase upon DNA methylation damage have been mapped with high resolution, providing quantitative measurement of the search time, binding time, and enzymatic rates of these two enzymes in live cells⁷⁴. Additionally, PALM imaging helped discover that in *E. coli*, Pol V, the translesion DNA synthesis enzyme⁷⁵, was initially sequestered on the inner membrane upon the induction of SOS response; only at a late stage was it released in a RecA-dependent manner into the cytosol to perform translesion DNA synthesis⁷⁶. Consistent with this observation, it was recently shown by a combination of SIM imaging and biochemical assays that in response to dsDNA break and genetic mutations, RecA forms bundled filaments along the inner membrane through its interaction with anionic phospholipids^{77,78} (Fig. 3b). With much enhanced resolution, these studies revealed that RecA bundles are likely composed of multiple laterally associated RecA filaments, with thinner RecA-ssDNA filaments extending from both ends (Fig. 3b,f), which dynamically sample the inner cell compartment likely in search for homology^{77,78}.

Similar strategies also revealed interesting spatial distribution patterns and diffusive behaviours of RNA polymerase (RNAP)^{64,68,79–81}. RNAP forms dense clusters of ~150 nm in size preferably at the edge of nucleoid in fast growing cells, but distributes more homogeneously and becomes more mobile in slow-growing or transcription-inhibited cells^{64,68,81} (Fig. 3d). Furthermore, two-colour SIM imaging found that dense RNAP clusters colocalize with transcription anti-termination factors NusA and NusB (ref.⁸²), likely suggesting that multiple RNAP molecules cluster on ribosomal RNA (*rnm*) operons⁸³, although it remains unknown whether these RNAP clusters are actively engaged in transcription.

Finally, quantitative PALM imaging and two-colour super-resolution colocalization studies have helped elucidate molecular mechanisms of DNA-associated machineries by providing precise cellular positioning and molecular counting measurements. SpoIIE, a DNA translocase responsible for exporting the replicated chromosome from the mother cell to the forespore compartment during sporulation in *B. subtilis*⁸⁴, was shown to form paired DNA channels on each side of the sporulation septum, the separation of which is too small (<50 nm) to be visible by conventional fluorescence microscopy^{85,86} (Fig. 3c,f). Another example is that ParB and ParA, two proteins involved in chromosome segregation in *C. crescentus*, were found to localize with ~50 nm difference with respect to the pole organizing protein PopZ (Fig. 3e,f), revealing the differential roles of PopZ in modulating these two proteins' spatial organization and dynamics⁸⁷.

Cellular localization and morphology of cytoskeletal structures

It is now well documented that a large number of bacterial proteins, in particular polymeric proteins such as actin and tubulin homologues, form subcellular structures that have specific shapes and cellular locations. Optical nanoscopy can probe spatial features and organization of these structures at unprecedented resolution, providing new information regarding their possible molecular compositions, assembly mechanisms and functional roles (Fig. 4).

The most studied subcellular structure in microbial optical nanoscopy so far is the bacterial cytokinesis ring formed by the essential division protein FtsZ and its partners. The FtsZ protein is a tubulin homologue and a GTPase^{88,89}; it polymerizes at the midcell to form a ring-like structure, called the Z-ring, to recruit all other division proteins and initiate cytokinesis. While appearing as a smooth ring under conventional light microscopy, optical nanoscopy revealed that the Z-ring is a discontinuous, patchy structure in many bacterial species^{90–96}, with FtsZ clusters loosely and heterogeneously organized in a toroid zone of ~70–100 nm by 40–60 nm (refs^{90,92,97–100}; Fig. 4a). Most remarkably, due to much improved resolution, other proteins originally believed to form structures that are dependent on and colocalize with the Z-ring, were found to localize to places adjacent to, but distinct from, that of the Z-ring, providing new insight into their roles in positioning and stabilizing the Z-ring^{101–103} (Fig. 4b,c).

Beside FtsZ, actin homologues MreB and ParA, and intermediate-filament-like protein CreS also forms polymeric structures. Under super-resolution, the patchy, irregular foci formed by MreB, the protein involved in maintaining the rod-cell shape in many bacterial cells, resolved into thin, sometimes discontinuous filaments of variable lengths^{104–107}. Importantly, live cell SIM–STED imaging observed length-dependent moving velocity of individual filaments (Fig. 4d), leading to a novel mechanistic model in which multiple cell wall synthesis motors are coupled on the same MreB filament and hence exert concerted peptidoglycan (PG) synthesis¹⁰⁴. The other actin homologue ParA, a protein involved in chromosome segregation, was found in one study to form a narrow (~40 nm thickness) linear structure spanning the long axis of *C. crescentus* cells, possibly providing a track for the directional movement of the segregating chromosome guided by the ParB-*parS* complex¹⁰⁸. Finally, CreS, or crescentin, a protein required for maintaining the crescent shape of *C. crescentus*¹⁰⁹, has recently been imaged together with the surrounding cell membrane in live cells, both with 20–40 nm resolution in 3D (Fig. 4e)¹¹⁰.

Assembly and dynamics of membrane proteins

Microorganisms are surrounded by cell envelopes consisting of an inner membrane, a cell wall made of PG layers, and for Gram-negative bacteria, an outer membrane. Cell envelopes play essential roles, such as defining cell shape and division, resisting turgor pressure, providing receptor sites for viruses and antibiotics, and controlling cell adhesion and biofilm formation. At this level, optical nanoscopy continues to enable the mapping of unique membrane distribution patterns and dynamics of a variety of membrane proteins, including chemotaxis receptors¹¹¹, sporulation complexes¹¹², toxins¹¹³, respiratory systems¹¹⁴ and surface sugar metabolizing complexes¹¹⁵, and helped elucidate possible mechanisms of how

a membrane protein's localization and dynamics are coupled to its biological function. Yet here, force nanoscopy offers unique advantages over optical nanoscopy in probing structural and biophysical properties of cell surfaces with unmatched spatiotemporal resolution.

Force nanoscopy has enabled probing of cell surface structures directly in live cells¹¹⁶. AFM captured the surface ultrastructure of spores from the fungus *Phanerochaete chrysosporium*, showing that the surface was uniformly covered with crystalline protein layers (rodlets) with a periodicity of 10 nm (ref.¹¹⁶). However, the highest resolution was achieved on isolated membranes. Since the first molecular images of native *E. coli* OmpF porin surfaces were reported in 1995¹¹⁷, AFM has become an increasingly popular technique for observing protein assemblies and their structural changes in bacterial membranes, including surface layers, outer membranes from Gram-negative bacteria, and intracytoplasmic membranes of purple photosynthetic bacteria^{4–7}. AFM may also be combined with cryo-EM to resolve the architecture of surface layers like the exosporium in *B. cereus* and *B. thuringiensis* spores¹¹⁸.

Currently, there is much effort in applying HS-AFM in microbiology. A prominent example is the fast imaging of the purple membrane, which contains 2D arrays of the light-driven proton pump bacteriorhodopsin. The structure of individual bacteriorhodopsin trimers could be observed at a speed of 100 ms per image¹¹⁹, and the motion of monomers and trimers was visualized, suggesting that the protein arrays were in dynamic association–dissociation equilibrium. A pioneering dynamic imaging study revealed structural changes of bacteriorhodopsin in response to light within 1 s (Fig. 5a)¹²⁰. Under illumination with green light, cytoplasmic parts of monomers were brought into contact with neighbouring trimers and a cooperative interplay between excited proteins was observed. Additional observations brought novel insights into the trimer–trimer interaction energy and the amino acids involved in trimer interactions¹²¹.

The high-speed technology has also provided key information about the diffusion of membrane proteins, which are often coupled to their functional states. An elegant example is the characterization of the diffusion and interaction of the OmpF porin¹²² (Fig. 5b). The motion of about 70 OmpF trimers was tracked and individual subunits were resolved, revealing wide distribution in the membrane due to diffusion-limited aggregation. Protein motion correlated with the local density of proteins, but single protein molecules were also found to diffuse freely.

Three-dimensional architecture and synthesis mode of cell wall

PG is a major component of bacterial cell walls, and has received considerable attention in view of its role in controlling cell shape, growth and division. Force and optical nanoscopy methods are often used in conjunction with each other to probe not only the overall architecture of PG with nanometre precision, but also its remodelling and synthesis dynamics with specificity. Together with EM, they have disclosed a diversity of PG supramolecular structures in different bacterial species^{123,124}.

For example, while all Gram-positive microorganisms have thick cell walls, their PG architectures are drastically different from each other. Analysis of purified sacculi from *B.*

subtilis revealed PG nanocables running parallel to the short cell axis and led to a model in which glycan strands polymerize to form PG ropes¹²⁵. The coccal bacterium *S. aureus* showed a complex surface pattern with rings and knobble architectures associated with nascent and old PG synthesis respectively, and a large belt of PG forming a piecrust structure in the cell division plane¹²⁶. Ovococcal bacteria such as *Streptococcus pneumoniae*, *Enterococcus faecalis* and *Lactococcus lactis*, however, showed relatively smooth surfaces, with thin PG bands parallel to the short axis of the cell and no visible twists associated with cable-like structures¹²⁷. In the Gram-negative bacterium *E. coli*, high-resolution AFM revealed features running parallel to the plane of the purified sacculi and bands of porosity. Further combined with STORM imaging, it was suggested that only porous regions of PG are permissive for synthesis¹²³. Of note, cell wall architectures have also been explored in living cells: AFM imaging revealed networks of PG fibres and nanocables on *Bacillus atrophaeus* spores¹²⁸ and *L. lactis* cells¹²⁹, respectively.

In optical nanoscopy, labelling of PG using dye-conjugated wheat germ agglutinin (WGA, binds to GlcNAc residues¹³⁰), fluorescent d-amino acids (FDAAs, can be incorporated into PG when included in growth media^{131,132}) and fluorescent vancomycin (binds to terminal D-Ala–D-Ala residues¹³³) have allowed novel PG synthesis modes to be identified in a variety of different bacterial species^{93,134–136}. Alternatively, labelling of proteins involved in PG synthesis and remodelling using PA-FP fusions^{96,137}, immunofluorescence^{136,138}, or fluorescent antibiotics that specifically bind to PBPs (penicillin binding proteins¹³⁹) has revealed distinct cellular localization patterns of these enzymes with respect to other cell division and cell wall components, providing new insight into their roles at different stages of cell cycle.

Molecular mechanisms of biofilm formation

A remarkable feature of microorganisms is their ability to switch between two lifestyles, namely free-living planktonic cells and biofilm-associated cells³. Two major players in biofilm formation are cell surface adhesins, which mediate adhesion to biomaterial and host surfaces, and extracellular matrix polymers that hold the cells together. AFM-based force spectroscopy has been used to study the binding mechanism of single molecules (SMFS), and to quantify the forces driving the adhesion of whole cells (SCFS). Much of our recent knowledge comes from the analysis of *Staphylococcus epidermidis* and *S. aureus*, which form biofilms on indwelling medical devices. Using SMFS, weak interactions (~100 pN) were measured between single fibronectin (Fn) molecules and fibronectin-binding proteins (FnBPs) from *S. epidermidis*¹⁴⁰. FnBP binding forces were also measured on isolates from patients with an infected device, revealing a distinct binding force signature and specific single-amino-acid polymorphisms¹⁴¹. Notably, the *S. epidermidis* SdrG protein was found to bind to the blood plasma protein fibrinogen with a force of 2 nN, which is much stronger than the force measured so far for all other adhesins¹⁴². This strong binding originates from the high affinity ‘dock, lock, and latch’ (DLL) binding mechanism, which involves dynamic conformational changes leading to highly stable complexes. Unlike SdrG, the collagen-binding protein SdrF featured a dual-ligand-binding activity¹⁴³. Adhesion to collagen-coated substrates was mediated by weak and strong bonds involving two distinct regions of the

protein. The high dissociation rates of the bonds suggested they were less stable than DLL bonds.

The extracellular matrix mediates intercellular adhesion during the development of biofilms, but the underlying mechanisms are poorly understood. A widely investigated biofilm matrix component is the polycationic polysaccharide intercellular adhesin (PIA). Applying force nanoscopy to a clinically-relevant methicillin-resistant *S. aureus* (MRSA) strain, revealed the binding mechanism of PIA²⁴ (Fig. 6a). Whereas multiparametric imaging showed that bacteria expressing PIA were surrounded by a soft and adhesive matrix, force measurements demonstrated that PIA mediates multivalent electrostatic interactions with polyanionic teichoic acids on the surface of neighbouring cells.

There is evidence that the development of microbial biofilms also involves protein-based matrices. FnBPA proteins from *S. aureus* were shown to promote intercellular adhesion *via* low-affinity, zinc-dependent homophilic bonds between FnBPA domains on neighbouring cells¹⁴⁴. Unlike the very strong and stable DLL bonds, homophilic bonds showed moderate strength and fast dissociation, which could be important for biofilm dissemination. For another *S. aureus* matrix protein SasG, AFM demonstrated zinc-dependent homophilic interactions between proteins on opposing bacteria, and unravelled the molecular elasticity of individual SasG molecules²³ (Fig. 6b). While the protein mechanics measured on live cells was consistent with those measured on purified proteins¹⁴⁵, absorption of zinc to the bacterial cell surface favoured SasG exposure and promoted homophilic binding of SasG proteins on opposing cells (Fig. 6b).

Cell–cell adhesion interactions have also been studied in fungal cells. For example, aggregation of the yeast *Saccharomyces cerevisiae* is mediated by lectin-like flocculin (Flo) proteins through molecular interactions that are not fully understood. Weak lectin–sugar interactions as well as strong unfolding forces associated with the force-induced extension of the hydrophobic tandem repeats of Flo proteins, have been shown to be involved in yeast flocculation¹⁴⁶. The same behaviour was demonstrated for agglutinin-like sequence (Als) adhesins from *C. albicans*¹³, suggesting that lectin-like and Ig-like fungal adhesins have evolved similar domains with high mechanical strength to achieve adhesion. In *C. albicans*, single-molecule manipulations demonstrated that mechanical force triggers the formation of amyloid-like nanodomains of Als adhesins on the yeast cell surface, thus showing that microorganisms can use functional amyloids to activate adhesion, both through the lateral clustering of adhesins, and through amyloid bonds between cells.

Recently, SCFS has been implemented to study forces guiding microorganism–host interactions. The nanoscale adhesion forces between *P. aeruginosa* and host epithelial cells were shown to involve the extension of bacterial type IV pili and the formation of membrane tethers from host cells¹⁷. These mechanical responses may play a role in host colonization by increasing the adhesion lifetime of bacteria. In another study, the adhesion forces between *C. albicans* and macrophages were found to involve multiple specific molecular bonds between lectin receptors on the macrophage membrane and mannan carbohydrates on the fungal cell surface¹⁴⁷. Furthermore, an innovative method combining multiparametric

imaging with single bacterial probes was developed to map simultaneously the topography and adhesion properties of human skin at high spatiotemporal resolution¹⁴⁸.

Optical nanoscopy has also provided insight into biofilm formation. Using an *in vivo* labelling strategy, the extracellular matrix of developing biofilms of *Vibrio cholerae* was imaged with single-molecule precision and three distinct levels of spatial organization of cells in the biofilm revealed, suggesting complementary architectural roles of the main matrix constituents¹⁴⁹. In another study, biogenesis of bacterial membrane vesicles in *Pseudomonas aeruginosa* biofilms was investigated, and explosive cell lysis was found to play a crucial role in forming membrane vesicles that contribute to the structural integrity of the biofilm matrix¹⁵⁰.

Combining the unique advantages of optical and force nanoscopy is particularly powerful in biofilm research. In recent work, AFM force measurement was combined with 3D-SIM imaging to show that cell surface localization of the LapA protein from *Pseudomonas fluorescens* correlated with adhesion forces, thus providing strong evidence that LapA function as a cell-surface biofilm adhesin¹⁵¹.

Towards novel applications in diagnosis and therapy

Force nanoscopy has also established itself as a valuable approach for diagnosis and therapy. An AFM cantilever-based technology¹⁵² was exploited to understand how cell wall PG interacts with the clinically important antibiotics vancomycin and oritavancin¹⁵³, which suggested that mechanical stress alters the membrane and cell wall, thus leading to cell death. The device could also be used to better understand the influence of dosing and competing ligands on the functionality of the drugs. Another example involves monitoring the fluctuations of cantilevers in order to quickly assess the sensitivity and resistance of bacteria to antibiotics¹⁵⁴. The technology could also determine the presence of viable microorganisms in complex, uncontrolled environments, such as soil and river samples¹⁵⁵. These studies illustrate the power of AFM for studying the mechanism of action of antibiotics, and for helping select the most efficient treatments against pathogens.

Lastly, SCFS revealed the mechanism by which carbon nanoparticles (fullerenes) functionalized by multiple mannose residues are capable of blocking the adhesion of uropathogenic bacteria to their carbohydrate receptors via high-affinity multivalent bonds¹⁵⁶. The authors suggested that this direct, label-free method could lead to novel applications in anti-adhesion therapy, that is, for the design of peptides or antibodies capable of treating microbial infections.

Conclusions

Characterization of multi-molecular structures and machineries of microbial cells is an essential step towards understanding cellular processes and functions, and could potentially lead to novel applications in medicine and biotechnology. Because of their small dimensions, microbial subcellular structures have long been difficult to study. Examples discussed here demonstrate that optical and force nanoscopy methods can tackle this problem with unprecedented resolution and sensitivity. Super-resolution and AFM

techniques represent a powerful toolkit for probing the organization, dynamics, interactions and functionality of single molecules, from the inside out, and up to intercellular interactions, thereby allowing microbiologists to answer outstanding questions that were impossible to address before and to develop ultrasensitive assays for diagnosis and therapy.

Ultimately, the full potential of nanoscopy will be achieved when combining optical and force modalities. Establishing these correlated platforms in microbiology should allow the identification and tracking of specific cellular components, while probing their biophysical properties (adhesion, elasticity) simultaneously on the same single cell, thus contributing to the important connection between their structures and functions. Toward this goal, correlated AFM-fluorescence imaging has been exploited to track cell surface dynamics during cellular morphogenesis^{157,158}. A recent study established a correlated SMLM and AFM imaging platform for localizing specific proteins within high-resolution AFM images¹⁵⁹. Although correlative nanoscopy is still in its infancy, this approach offers promising prospects for the comprehensive analysis of the structure, dynamics and interactions of single molecules in microbial cells.

Acknowledgments

Work in the Dufrêne team was supported by the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme (grant agreement no. 693630), the National Fund for Scientific Research (FNRS), the FNRS-WELBIO (grant no. WELBIO-CR-2015A-05), the Federal Office for Scientific, Technical and Cultural Affairs (Interuniversity Poles of Attraction Programme), and the Research Department of the Communauté française de Belgique (Concerted research action). Y.F.D. is Research Director at the FNRS. Work in the Xiao lab is supported by National Institute of Health General Medicines 1R01GM086447-06, 1R01GM112008-01 (Multi-PI), National Science Foundation grant EAGER MCB1019000. Y.F.D. and J.X. thank Carla Coltharp and Xinxing Yang for their critical reading and suggestions of the manuscript.

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Box 1**Limitations of nanoscopy approaches****Force nanoscopy**

- Surface-sensitive technique, so cannot probe intracellular structures.
- AFM imaging and force spectroscopy (SMFS, SCFS) are not routine and require several months of advanced training to obtain reliable results.
- Success will depend on the quality of sample and tip preparation protocols, the accuracy of recording conditions, data collection and interpretation, the type of specimen being analysed (live cells versus purified structures), and the availability of proper controls (blocking experiments, mutant strains).
- Not all microbiological specimens will be well suited for AFM. Softer cells, like Gram-negative bacteria, will generally feature poor data because the cell is deformed or damaged due to the local pressure applied by the tip, or because the tip is contaminated by loosely bound molecules.
- Imaging speed has always been a limiting factor and recent HS-AFM instruments may not be appropriate for imaging live cells.
- Conventional SMFS and SCFS assays are low-throughput, preparation of single-molecule and single-cell probes is time-consuming, meaning performing statistically significant numbers of measurements is difficult.

Optical nanoscopy

- High spatial resolution in optical nanoscopy comes at the expense of temporal resolution, and requires the optimal combination of both imaging and labelling conditions.
- Fluorescent labelling of molecules of interest may alter the biological function/localization of labelled molecules, thus requiring careful validation of the functionality of the labelled molecules prior to imaging.
- Phototoxicity of cells caused by long time illumination of activation and excitation light prevents time-lapse live cell studies.
- Reconstructed images can be easily affected by sample drifting during imaging, mis-registration of multi-colour channels, and/or image reconstruction algorithms. Calibrations and control samples using fiducial markers should be used to avoid artefacts.

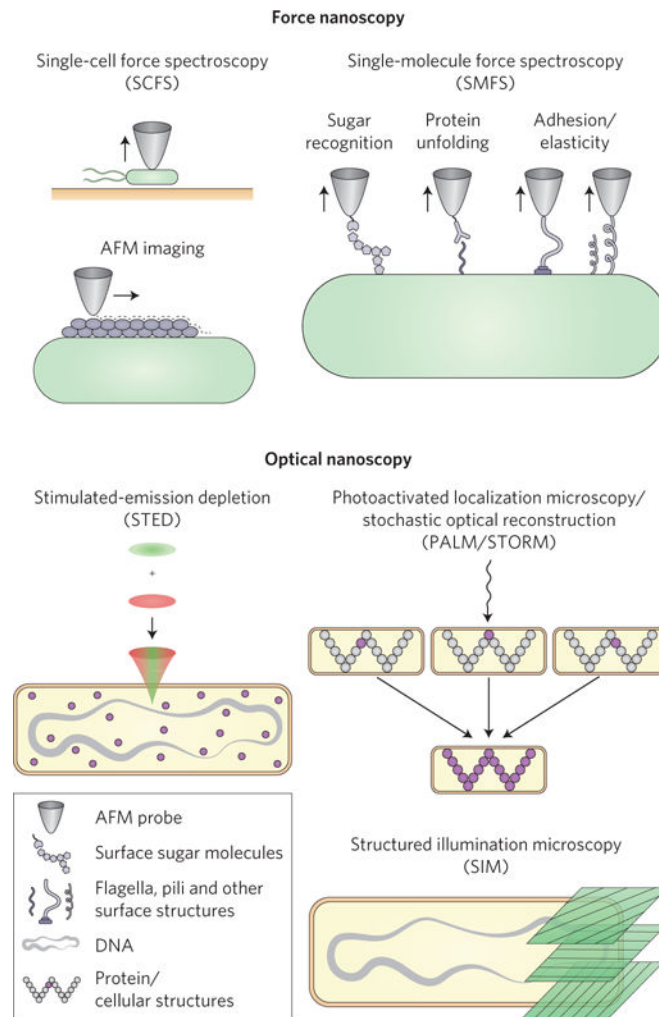


Figure 1. Probing the microbial cell from the inside out using nanoscopy

Top, force nanoscopy probes the outside of the cell. AFM imaging unravels the architecture of membranes and cell walls with nanoscale resolution. Whereas SMFS measures the localization, adhesion, elasticity and interactions of individual surface molecules and appendages, SCFS quantifies the forces driving the adhesion of whole cells. Bottom, optical nanoscopy probes the inside of the cell. The three commonly used imaging modes, STED, PALM/STORM, and SIM are shown. STED combines an excitation beam (green) and a depletion beam (red) to generate an excitation volume smaller than a diffraction-limited spot, which then raster scans the sample to generate a super-resolved image. PALM/STORM stochastically activates single molecules (purple) to acquire their positions subsequently, and reconstructs a super-resolution imaging by adding all the molecules' positions together. SIM uses wide-field illumination of different excitation patterns and takes advantage of the generated moiré fringes (convolutions between green illumination patterns and cellular structures) to extract the information about the underlying structure.

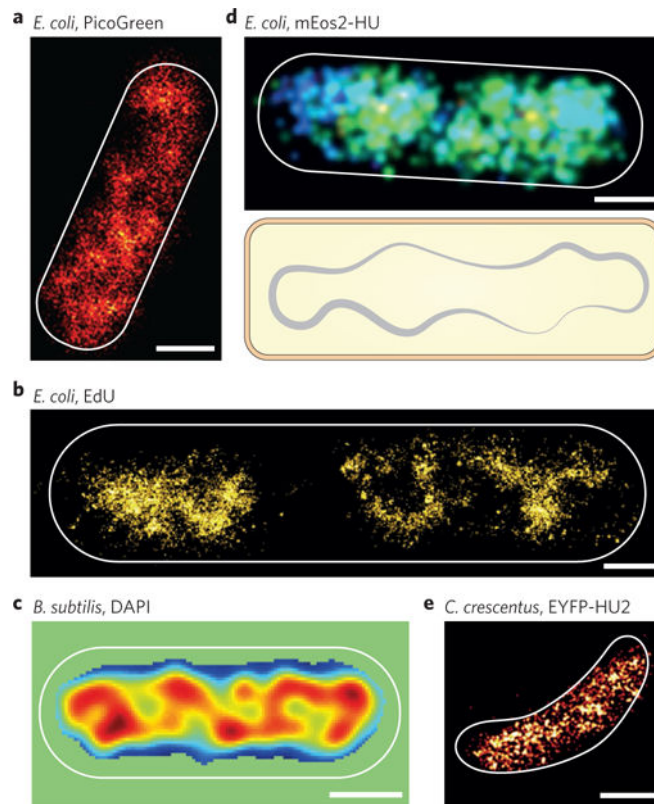


Figure 2. Spatial organization of bacterial nucleoid

a, BALM image of a fixed *E. coli* cell showing voids and inhomogeneous nucleoid organization when labelled with PicoGreen⁵⁴. **b**, STORM image of a fixed *E. coli* cell showing asymmetric and heterogeneous nucleoid structures⁶³. The cell was incubated with EdU in LB medium, subsequently fixed and labelled with Alexa647 by click chemistry. **c**, 3D-SIM image of a DAPI-stained live *B. subtilis* cell showing high-density chromosomal regions (red peaks)⁶⁷. **d**, Top, a live *E. coli* cell grown in minimal M9 medium showed dispersed, rather uniform distribution of mEos2-labelled HU protein in 3D super-resolution imaging⁷⁰. Bottom, schematic of a bacterial nucleoid. **e**, Super-resolution image of EYFP-HU2 in a fixed *C. crescentus* cell grown in minimal M2G medium⁶⁹. All cell contours are indicated by white outlines. Scale bars, 500 nm. Figure reproduced from: ref.⁵⁴, ACS (**a**); ref.⁶³, Elsevier (**b**); ref.⁶⁷, Elsevier (**c**); ref.⁷⁰, AAAS (**d**); ref.⁶⁹, Biophys. Soc. (**e**).

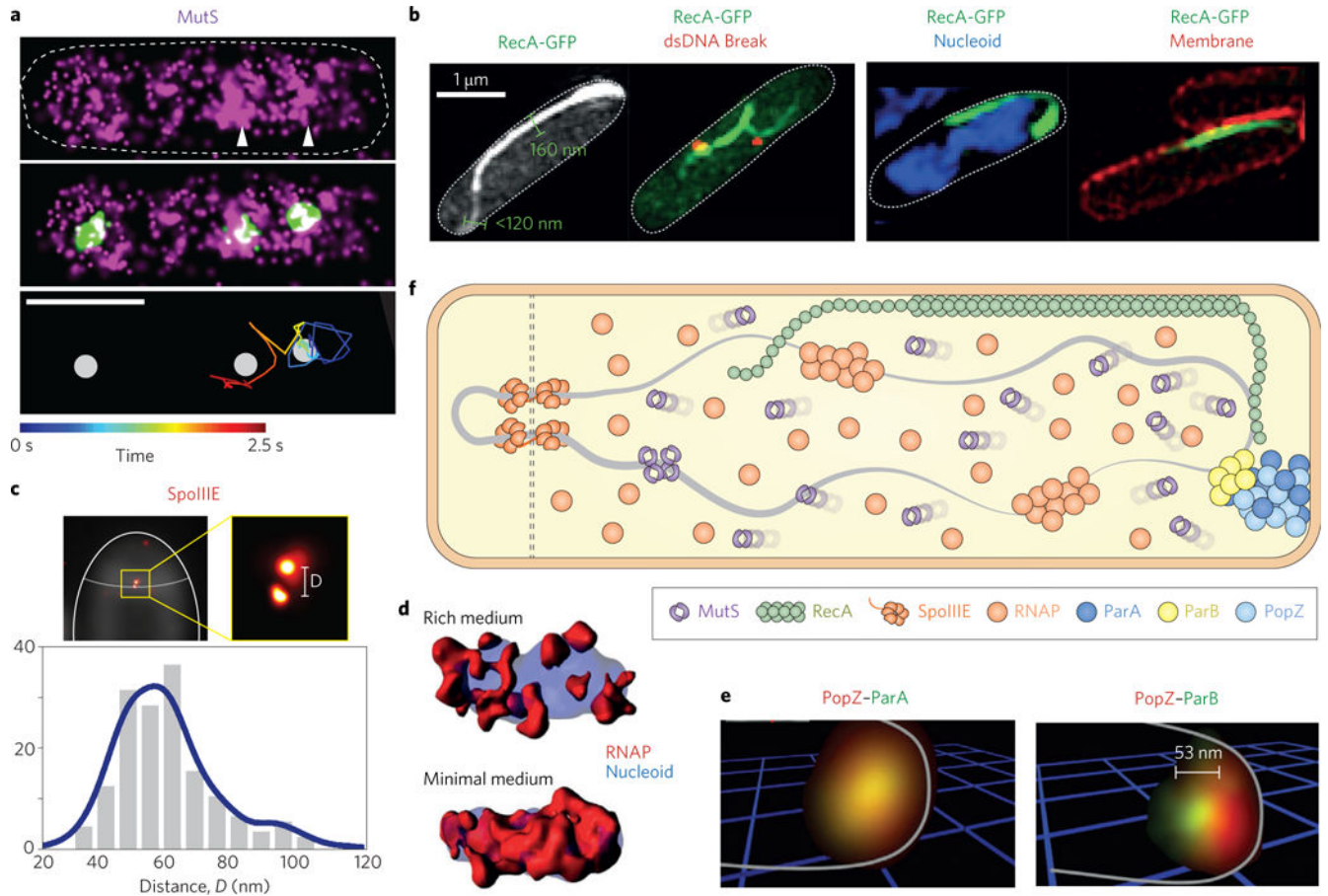


Figure 3. Spatial distribution and dynamics of DNA-associated machineries

a, Spatial distribution of MutS-PAMCherry in a live *E. coli* cell (top), superimposed with DnaX-Citrine-labelled replisome (middle, green spots), and a representative time-coloured SMT trajectory of MutS-PAMCherry showing its slowed diffusion in the replisome regions (bottom, white spots)⁷³. Scale bar, 1 μm. **b**, 3D-SIM images of *E. coli* cells showing RecA-GFP alone, with ParB-mCherry labelled dsDNA break ends (red spots), with DAPI-stained DNA (blue), or with FM4-64 stained membrane (red)⁷⁷. The thickness of RecA bundles was measured at 160 nm on average, and the thinner filaments at the ends of the bundle were measured <120 nm, the resolution limit of SIM. **c**, PALM imaging to identify the spatial arrangement of SpoIIIE motor complexes at the septum between the forespore and mother cell in *B. subtilis*⁸⁵. Top left, a diffraction-limited, FM5-95 stained membrane image of a σ^E *B. subtilis* cell overlaid with the corresponding PALM image of SpoIIIE-tdEos. Cell and septum are outlined in white. Top right, zoomed PALM image (red foci) of SpoIIIE-tdEos showed clear separation between two complexes. Bottom, the histogram of the separation distance *D* measured from individual dual foci showed a mean separation value at 55 nm. **d**, Surface rendering of 3D-SIM images of the nucleoid (DAPI, blue) and RNAP-GFP (red) in live *E. coli* cells showed highly clustered RNAP distribution in rich medium, but more uniform distribution in minimal medium⁶⁸. **e**, Two-colour PALM imaging of PAMCherry-PopZ (red) with ParA_{G16V}-EYFP (green, left) or EYFP-ParB (green, right) showing that inactive ParA_{G16V} colocalizes with PopZ, but ParB is displaced ~53 nm away

from PopZ (ref.⁸⁷). **f**, A cartoon summarizing cellular localizations and spatial organizations of DNA-associated machineries as depicted in panels **a–e**. Figure reproduced from: ref.⁷³, PNAS (**a**); ref.⁷⁷, NPG (**b**); ref.⁸⁵, eLife (**c**); ref.⁶⁸, PNAS (**d**); ref.⁸⁷, PNAS (**e**).

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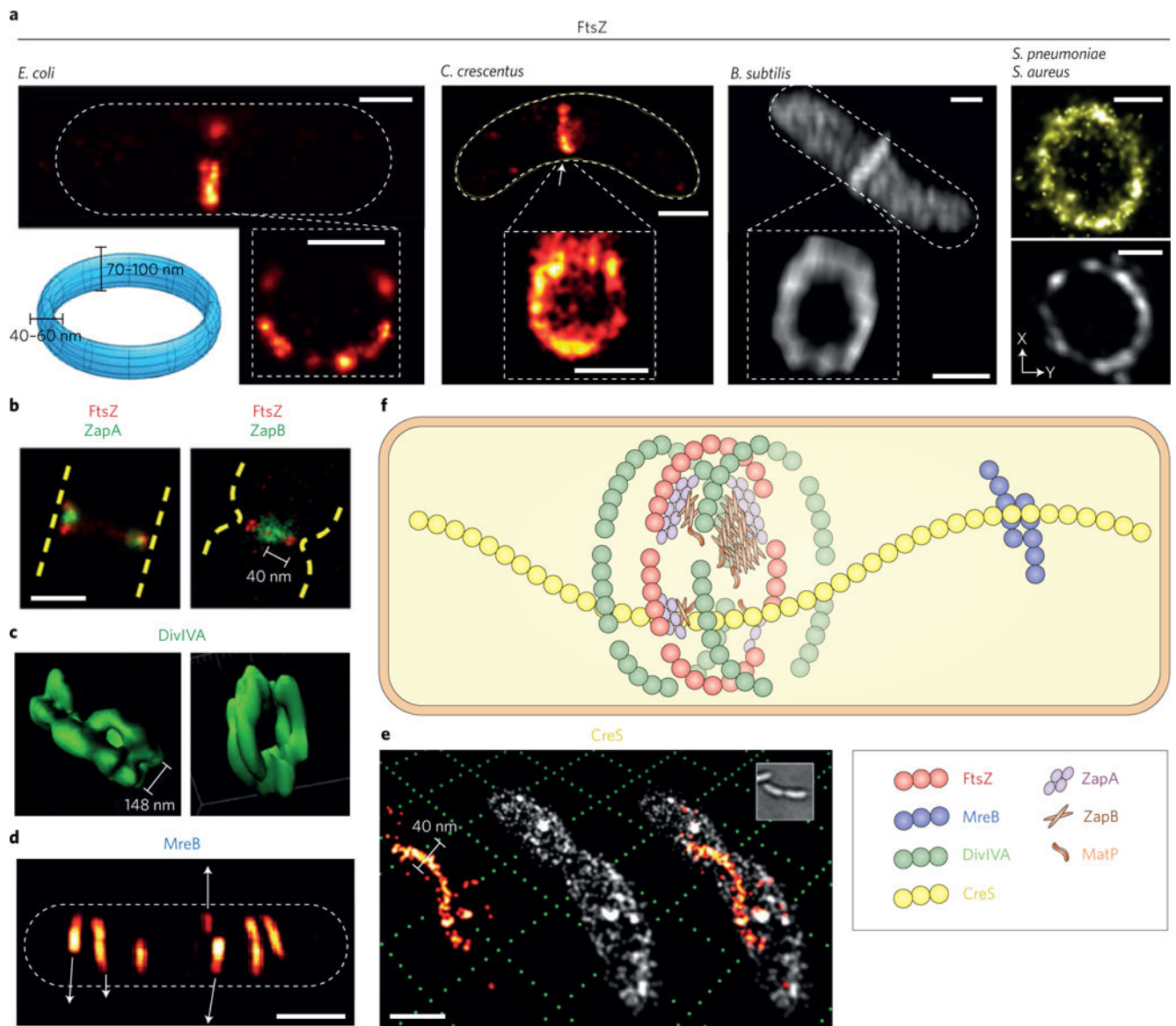


Figure 4. Intracellular location and morphology of cytoskeletal structures

a, 3D-PALM or SIM imaging showed patchy, discontinuous FtsZ-ring structures in different bacterial species. From left to right: FtsZ-mEos3.2 in *E. coli*¹⁰⁰, FtsZ-Dendra2 in *C. crescentus*⁹², FtsZ-GFP in *B. subtilis*⁹¹, FtsZ-spDendra2 in *S. pneumoniae*⁹⁴ and in *S. aureus*⁹¹. A blue toroid at the bottom left shows the dimensions of the midcell zone the FtsZ ring occupies in most bacterial species measured. **b**, Two-colour PALM imaging in live *E. coli* cells shows that ZapA (Z-ring associated protein A) largely colocalizes with the Z-ring while ZapB is displaced inward ~40 nm (ref.¹⁰³). **c**, SIM imaging shows that DivIVA, a peripheral membrane protein targeting FtsZ antagonist MinC protein to membrane in *B. subtilis*, forms a double ring flanking the septum, and hence likely regulating the midcell positioning of the FtsZ-ring¹⁰¹. **d**, SIM imaging of GFP-MreB in *B. subtilis* showed MreB filaments moving independently along both directions of the membrane¹⁰⁴. **e**, 3D, two-colour super-resolution imaging of Nile Red-stained membrane (white) and EYFP-labelled

CreS, an intermediate-filament like protein that determines the crescent shape of *C. crescentus*¹¹⁰. The thickness of CreS filament was measured at ~40 nm. Scale bars, 500 nm except **d** (1 μ m). **f**, Cartoon of cytoskeletal structures observed by optical nanoscopy as depicted in panels **a–e**. Figure reproduced from: ref.¹⁰⁰, Wiley (**a**, *E. coli*); ref.⁹², PNAS (**a**, *C. crescentus*); ref.⁹¹, PLoS (**a**, *B. subtilis*, *S. aureus*); ref.⁹⁴, ASM (**a**, *S. pneumoniae*); ref.¹⁰³, PLoS (**b**); ref.¹⁰¹, ASM (**c**); ref.¹⁰⁴, Biophys. Soc. (**d**); ref.¹¹⁰, PNAS (**e**).

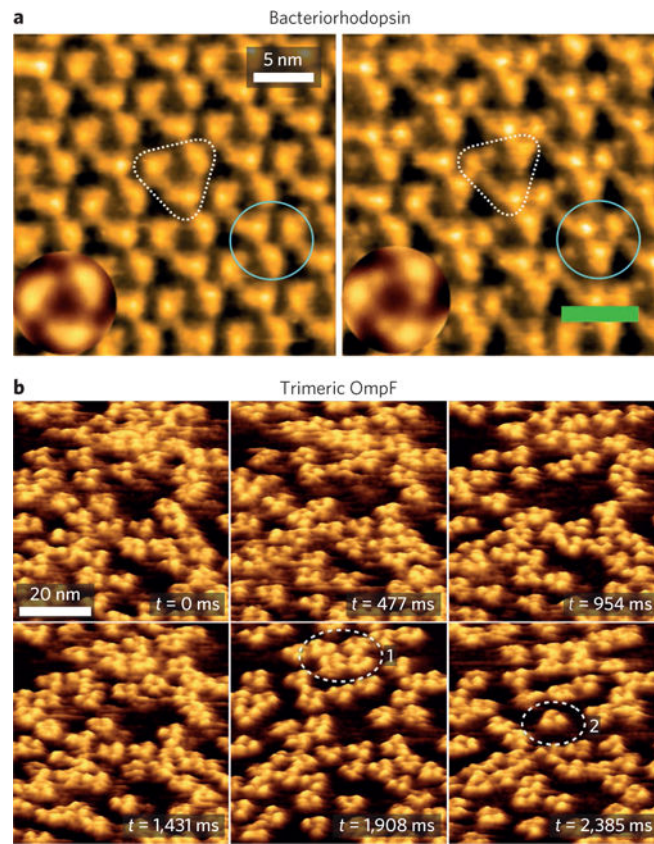


Figure 5. Imaging the structure and dynamics of single proteins in bacterial membranes
a, High-speed AFM images of bacteriorhodopsin documenting structural differences under dark (left) and green light illumination (right)⁸. The circles and triangles highlight trefoils and trimers (insets: averaged images of trimers). **b**, High-speed AFM images of the diffusion and interaction dynamics of trimeric OmpF (ref.⁸). Unlike protein trimers outlined in (1), the protein outlined in (2) is highly mobile. Figure reproduced from ref.⁸, ACS.

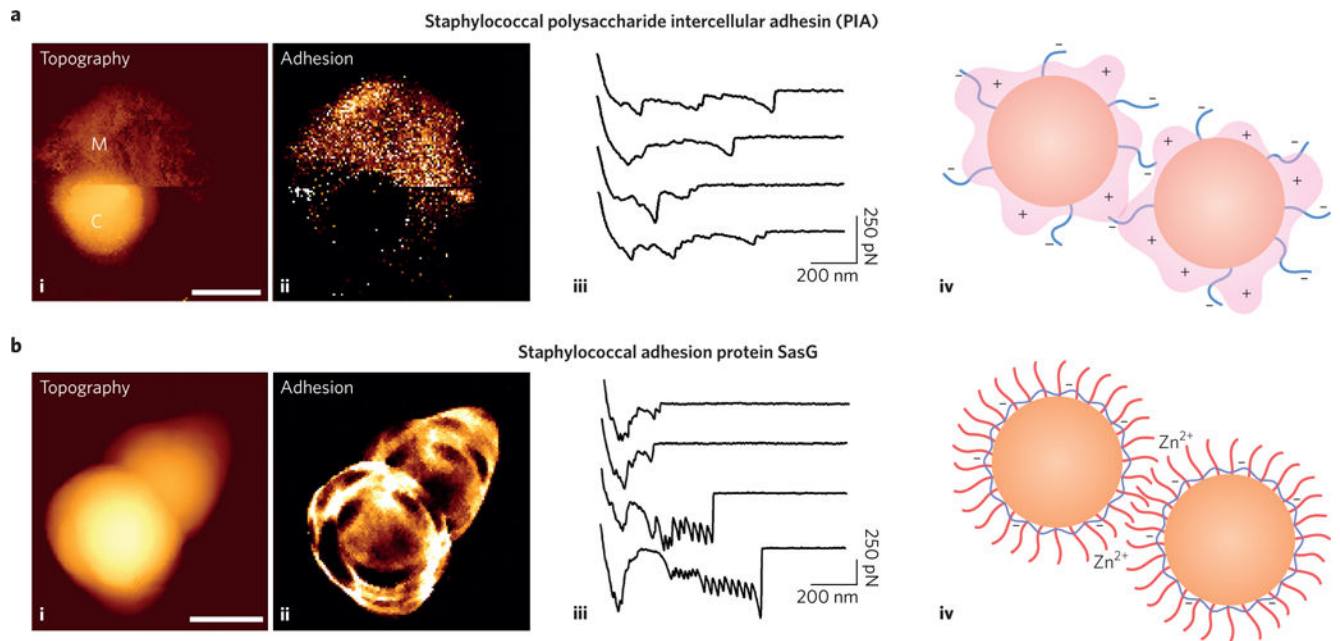


Figure 6. The matrix revolutions, dissecting the binding mechanisms of biofilm matrices
a. AFM unravels the molecular forces of the staphylococcal polysaccharide intercellular adhesin (PIA)²⁴: (i) topographic and (ii) adhesion images of a living *S. aureus* cell expressing PIA (labels M and C highlight the PIA matrix and the cell surface, respectively); (iii) adhesion force signatures recorded between two cells; and (iv) the binding mechanism involving the multivalent electrostatic interaction of cationic PIA (pink) with polyanionic teichoic acids (blue) on the cell surface. **b.** AFM deciphers the zinc-dependent mechanical properties of the staphylococcal adhesion protein SasG (ref.²³): (i) topographic and (ii) adhesion images of *S. aureus* cells expressing SasG recorded in the presence of zinc; (iii) adhesion force profiles recorded between two bacteria in the presence of zinc; and (iv) a proposed model for the zinc-dependent activation of intercellular adhesion (SasG proteins are in red, teichoic acids in blue). Scale bars, 1 μ m. Figure reproduced from: ref.²⁴, ACS (**a**); ref.²³, PNAS (**b**).