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**DOI**

[10.1038/nnano.2017.45](https://doi.org/10.1038/nnano.2017.45)

**Publication date**

2017

**Document Version**

Accepted author manuscript

**Published in**

Nature Nanotechnology

**Citation (APA)**

Dufrêne, Y. F., Ando, T., Garcia, R., Alsteens, D., Martinez-Martin, D., Engel, A., Gerber, C., & Müller, D. J. (2017). Imaging modes of atomic force microscopy for application in molecular and cell biology. *Nature Nanotechnology*, 12(4), 295-307. <https://doi.org/10.1038/nnano.2017.45>

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*Review article for Nature Nanotechnology*

## **Atomic Force Microscopy Imaging Modalities in Molecular and Cell Biology**

Yves F. Dufrêne<sup>1\*</sup>, Toshio Ando<sup>2</sup>, Ricardo Garcia<sup>3</sup>, David Alsteens<sup>1</sup>, David Martinez-Martin<sup>4</sup>,  
Andreas Engel<sup>5</sup>, Christoph Gerber<sup>6</sup> and Daniel J. Müller<sup>4\*</sup>

<sup>1</sup>Institute of Life Sciences and Walloon Excellence in Life sciences and Biotechnology (WELBIO), Université catholique de Louvain, Croix du Sud 4-5, bte L7.07.06., B-1348 Louvain-la-Neuve, Belgium.

<sup>2</sup>Department of Physics, Kanazawa University, Kanazawa 920-1192, Japan.

<sup>3</sup>Instituto de Ciencia de Materiales de Madrid, CSIC, Sor Juana Inés de la Cruz 3, 28049 Madrid, Spain.

<sup>4</sup>Department of Biosystems Science and Engineering, Eidgenössische Technische Hochschule (ETH) Zürich, Mattenstrasse 28, 4056 Basel, Switzerland.

<sup>5</sup>Department of BioNanoscience, Delft University of Technology, Van der Waalsweg 8, 2628 CH Delft, The Netherlands.

<sup>6</sup>Swiss Nanoscience Institute, University of Basel, Klingelbergstrasse 80, 4057 Basel, Switzerland.

\*e-mail: yves.dufrene@uclouvain.be; daniel.mueller@bsse.ethz.ch

## **Abstract**

Within three decades, atomic force microscopy (AFM) has become a powerful multifunctional imaging platform, enabling the visualization and manipulation of biological samples, from single molecules to living cells. Soon after its invention, it was recognized that the full potential of AFM imaging in biology requires key technological developments in order to solve a number of limitations and drawbacks. This need has been a key driving force towards developing new imaging modalities, pushing continuously the field ahead. Here, we survey the basic principles, advantages and limitations of the most common AFM bioimaging modalities available today, starting from the popular contact and dynamic modes, to newly developed modes, including multiparametric, molecular recognition, multifrequency and high-speed imaging. We discuss recent examples that highlight the unique capabilities of these emerging new modalities. We anticipate that, in the next decade, these nanotools will have a profound influence on the way researchers look at biological systems, thereby helping them to solve fundamental questions that could not have been addressed with traditional techniques.

## Introduction

The emergence of atomic force microscopy (AFM) 30 years ago<sup>1</sup> in the then fledgling field of nanotechnology<sup>2</sup> has opened new avenues in physics, chemistry, biology, and medicine, and since then has continuously inspired researchers all over the world, as testified by more than 340,000 scientific articles in peer reviewed journals (web of science). The key invention was to contour nonconductive surfaces much below the diffraction limit of light by controlling a conglomerate of forces acting between a tiny probe and the object. Whereas first results on the atomic scale were obtained within a year<sup>3</sup>, it took another few years to accomplish atomic imaging of nonconductive surfaces in vacuum. Meanwhile, the technique started to be adapted to work over a vast temperature scale and basically in every environment<sup>2,4,5,6</sup>. The ability to investigate surfaces with exceptional signal-to-noise ratio at sub-nanometer resolution triggered a wealth of AFM-related techniques using a variety of probes to locally sense interactions and manipulate matter from the atomic to microscopic scale<sup>2,7</sup>. This unique flexibility of AFM to image, probe and manipulate materials made it the most versatile toolkit in nanoscience and -technology, changed our perception of hard and soft matter and stimulated revolutionary discoveries and technologies<sup>2</sup>. The possibility to operate in fluidic environments and at ambient temperature moved AFM towards biology, opening the door to image and probe molecules and cells at (sub-)nanometer resolution<sup>4,5,6,8,9</sup>. To address the wide complexity of biological systems, ranging from lipids, nucleic acids, proteins, assemblies thereof, to cells and tissues, a wealth of AFM modalities have been developed over the years (**Fig. 1**). Major advances in high-resolution imaging have also been achieved in complementary methods including super resolution microscopy and cryo-electron microscopy, which significantly enrich the imaging toolbox now available to molecular and cell biology (**Table 1**).

Many reviews have been published in the past two decades that describe the use of certain AFM imaging modalities to characterize biological systems. However, as it is difficult for newcomers and often even for advanced users to overview the principles of these quickly developing and diverse imaging modalities and to evaluate their applicability, advantages and limitations, we here survey the most significant steps that have led to establish AFM as a powerful toolbox in molecular and cell biology. We outline for each AFM imaging modality to which kind of biological systems it can be preferably applied, their current limitations and future perspectives.

## A journey into AFM imaging techniques

### Imaging native biological systems in liquid

The key breakthrough that led to biological AFM was the development of an optical detection system, followed by the design of a fluid chamber, enabling imaging in buffer solution and thus maintaining the native state of the biological system<sup>4,5</sup>. The first AFM imaging mode invented, *contact mode*, raster scans a tip over the sample and adjusts pixel-by-pixel the height of the tip so that the force applied to the sample is kept constant (**Fig. 2a**). The resulting height image resembles the sample topography with the resolution depending on the radius of the tip, the sample corrugation, the physical properties of the sample, and how precisely the feedback system contours the tip over the soft biological

sample.

Shortly after introducing the first commercially available AFM, biological specimens imaged included animal cells<sup>10,11</sup>, cell membrane patches and membrane proteins<sup>12,13,14</sup>, DNA and RNA<sup>15</sup>, as well as lipid films<sup>16,17</sup>. For flat, smoothly corrugated surfaces such as proteins protruding  $\approx 1$  nm from membranes contact mode AFM can provide topographs of single membrane proteins at lateral and vertical resolution of  $<1$  nm and  $<0.1$  nm, respectively (**Fig. 2b**)<sup>14,18,19</sup>. This exceptionally high resolution and signal-to-noise ratio of AFM allowed for example to unravel the functionally relevant oligomeric state of various water-soluble and membrane proteins<sup>20,21,22,23,24</sup>. Operated in the time-lapse contact mode AFM visualized the morphological dynamics of cells<sup>10,11</sup>, the growth of pathological amyloid fibrils<sup>25</sup>, the enzymatic degradation of DNA<sup>26</sup> or lipid membranes<sup>27</sup>, and provided insight into the working principles of bacterial outer membrane pores<sup>28</sup>, gap junctions enabling intercellular connections between certain animal cells<sup>29</sup> and nuclear pore complexes<sup>30</sup>. Other exciting examples monitored the insertion of pathological toxins into membranes<sup>31</sup> and the supramolecular architecture of photosynthetic membranes changing in response to light<sup>32</sup>. Such insight allowed static structural models to be complemented with functional dynamics<sup>33</sup>.

Although contact mode AFM is widely used to characterize solid substrates, its application to soft biological systems requires expert skills to adjust the force applied to the tip. As a rule of thumb, forces  $>100$  pN should be avoided as they can cause reversible or even irreversible deformations<sup>33</sup>. *Dynamic mode imaging* (originally termed tapping or oscillation mode) was invented to minimize the friction and the force applied between tip and sample (**Fig. 2a,c**). In its simplest application, the cantilever is oscillated close to resonance while scanning across a sample<sup>1</sup>. Ideally the tip only touches the sample at the very end of its downward movement thus considerably minimizing friction. In close proximity to the sample surface, the interactions between tip and sample change both the cantilever amplitude and resonance frequency allowing them to be used as feedback parameters for contouring fragile biological samples<sup>34,35,36</sup>. Using the amplitude as feedback is technically simpler because it requires only one feedback loop compared to using frequency as feedback requiring three such loops. Thus, amplitude modulation AFM is currently more often applied than frequency modulation AFM. Besides these two well-known AFM imaging modes, other dynamic modes have been developed which employ different signals as feedback parameters or excite the cantilever at different frequencies simultaneously (see section Multifrequency Imaging)<sup>37</sup>. Importantly, as dynamic modes considerably reduce force and friction between tip and sample, they can be applied to image biological objects, which are only weakly adsorbed to supports, like DNA, single proteins, and filaments<sup>38,39,40,41</sup>. Dynamic modes also allow highly corrugated objects, like living cells, to be depicted in their unperturbed state<sup>42</sup>. However, the topographic contrast relies on rather complex interaction mechanisms between the AFM tip and sample. Stiffness, roughness, surface charge and chemistry, or friction of the sample can change the oscillation of the tip and thus alter or even invert the contrast<sup>37</sup>. To record faithful high-resolution images it can therefore be helpful to image unknown biological systems in the presence of structurally well-characterized reference samples<sup>37,43,44</sup>.

Applied to cellular systems contact and dynamic mode AFM reveal topographs below the resolution limit of conventional light microscopy. The ease of use and the exceptional signal-

to-noise ratio quickly raised the hope that AFM would revolutionize live-cell imaging<sup>4,5,10</sup>. Yet, only part of the dream came true. For example, the resolution of animal cell surfaces remained generally limited to  $\approx 50\text{--}100$  nm due to their soft and corrugated nature<sup>9</sup>. As opposed to animal cells, surfaces of microbes, which are mechanically much more rigid and generally smoother, have been routinely imaged approaching a resolution of  $\approx 10$  nm<sup>[45,46]</sup>. However, polysaccharides of the plasma membrane can contaminate the scanning tip thus changing the image contrast. An elegant approach for imaging living cells and circumventing tip contamination problems is scanning ion conductance microscopy (SICM), which scans a nanopipette over the sample while measuring the ion current<sup>47,48,49</sup>. The ion current is then used to control the vertical position of the nanopipette and thus to contour the sample. If adjusted properly this feedback parameter can be adjusted to avoid physical contact between pipette and cell. As a result SICM contours living cellular systems including hair cells or hippocampal neurons at superior resolution ( $\approx 50$  nm) and in the unperturbed state<sup>48</sup>. Excitingly SICM can be combined with single-channel patch clamp recordings. However, to apply SICM more widely requires overcoming bottlenecks including the intrinsically slow imaging process and the convolution of the rather large SICM probe with corrugated cell surfaces.

Last but not least, AFM cannot only be used to image but also to manipulate biological samples. The force applied to the AFM tip can simply be adjusted for mechanical manipulation, and the tip can be functionalized with chemical groups to manipulate specific sample regions. Thus AFM has been used to manipulate and dissect cells, chromosomes, viruses, membranes or single nucleic acids and proteins early on<sup>2,8,50</sup>. The possibility to mechanically manipulate biological systems guided the development of the AFM tip as nanotool to cut, pick up, release or to sculpt biomolecules at nanometer precision and very recently even to control the division of animal cells<sup>51,52,53,54</sup>.

### From force-distance curves to multiparametric imaging

The question came up whether AFM can do more than just contouring a surface. A milestone was the realization that, simultaneously with structural imaging, AFM is capable to probe biophysical properties. Initially such properties were measured by approaching the AFM tip to and retracting it from the biological sample while recording single force-distance (FD) curves<sup>55</sup>. Approach FD curves can quantify the height, surface forces, mechanical deformation of the sample, or derive its elastic modulus and energy dissipation. Retraction FD curves allow adhesion forces to be measured (**Fig. 3a**). To reliably characterize the properties of the sample implies precisely controlling the interaction between tip and sample, thus requiring AFM tips with well-defined geometry and surface chemistry. Sophisticated commercial micro- and nanomachined cantilevers and tips are now available, which are customized in terms of shape, tip radius and physical and chemical properties. As further discussed below, several imaging modes have been developed to extract the sample properties while imaging the sample<sup>56,57,58,59,60</sup>. A versatile and widely distributed approach among these is the FD curve-based imaging mode<sup>61,62,63</sup>, which, pixel-by-pixel, approaches and retracts the AFM tip to locally measure forces (**Fig. 3b**).

Modern FD curve-based AFMs (FD-based AFMs) acquire several hundreds of thousands of FD curves while imaging the biological sample<sup>63</sup>. As each FD curve locally quantifies physical

properties and interactions, this information can be directly mapped to the sample topography (**Fig. 3c**). FD-based AFM thus opens the door to image complex biological systems and to simultaneously quantify and map their intrinsic physical properties, including elasticity and adhesion (**Fig. 3d-e**). Although AFM provides an absolute measurement of the tip position ( $x,y,z$ ), it is often a challenge to determine the exact contact point between tip and sample (zero separation), particularly when long-range surface forces, surface roughness and deformation of the soft biological sample play roles. Knowledge of the contact point is needed to differentiate surface forces from the mechanical deformation of the soft cell. However, for most applications linearly extrapolating the contact region to zero force is sufficiently accurate (**Fig. 3b**).

Currently, the most widely used application of FD-based AFM is the mapping of the mechanical properties of biological systems. This is important because pertinent cellular functions rely on mechanical properties. Pioneering contributions applied the method to image and mechanically map drug-induced changes of the cytoskeleton of fibroblasts<sup>64</sup> and to spatially map the stiffness of the actomyosin cortex of adherent cultured cells during cell division<sup>65</sup> (**Fig. 3d**). Mapping the viscoelasticity of non-tumorigenic cells and breast tissues showed that they are less deformable compared to cancerous cells and malignant breast tissues, respectively<sup>66,67</sup>. This led to the conclusion that diseased cellular systems expose considerably altered mechanical properties. Imaging and mechanically mapping yeast cells (*Saccharomyces cerevisiae*) revealed a substantial stiffening of the chitin accumulating bud scar compared to the surrounding cell wall<sup>68</sup>.

Two interconnected issues in FD-based AFM are the lateral and temporal resolutions. In modern AFMs, the lateral resolution is mainly related to the tip radius, the tip-sample drift, the distance dependence of the tip-sample interaction, imaging force and the properties of the biological sample. Long-range surface forces interacting over several tens of nm reduce the resolution at which these interactions can be localized. Technically, when recording an AFM image at a certain frame size the number of pixels recorded determines the theoretically approachable resolution. However, the amount of pixels and thus the amount of force curves collected per FD-based AFM image is limited by the data acquisition time. In the early days of FD-based AFM<sup>61,62</sup>, the time required for recording a single force curve was between  $\approx 0.1$  and 10 s, and the time needed to acquire a FD-based AFM image of  $32 \times 32$  pixels ranged from  $\approx 2$  min to  $\approx 3$  h. Until recently, this slow imaging speed strongly limited the use of FD-based AFM imaging in biology, but the introduction of faster piezo elements, feedback loops, data acquisition systems, oscillation modes changing the tip-sample distance<sup>69</sup>, and of tailored cantilevers reducing hydrodynamic drag<sup>70,71,72,73</sup> largely solved this problem.

As a consequence, nowadays FD-based AFM can record  $512 \times 512$  pixels multiparametric images of native biosystems with a resolution approaching 1 nm, within time ranges of 15–30 min<sup>[63]</sup>. For instance, the method can image even individual membrane proteins in their native state at  $\approx 1$  nm resolution and simultaneously map the mechanical properties of their secondary structures<sup>74,75</sup> and of interfacing lipids<sup>74</sup>. FD-based AFM also mapped the mechanical properties of heterogeneous lipid membranes<sup>76</sup> and correlated mechanical properties of human keratinocytes<sup>77</sup> and bacteria<sup>78,79</sup> to their morphology and state. Applied to viruses FD-based AFM has shed new light into the relationship of structural, functional and mechanical properties of herpes simplex viruses<sup>50</sup>, bacteriophages<sup>79,80</sup>, southern bean

mosaic viruses<sup>81</sup> and parvovirus minute viruses<sup>82</sup>. Excitingly, FD-based AFM can map various molecular and surface forces from the micro- to nanometer scale, including complex and heterogeneous biological systems<sup>83,84</sup>. We are now beginning to understand the time-dependence of mechanical interactions, and we can measure for example the strength of chemical bonds<sup>85</sup>, as well as the mechanical response of biological materials under different loading rates<sup>59,86</sup>. Although technological improvements have considerably reduced the acquisition time of FD-based AFM images, it remains an important challenge to further increase the imaging speed so that the multiparametric complexity of dynamic molecular and cellular processes can be fully addressed.

### Molecular recognition imaging

Soon after introducing FD-based imaging, the idea to map specific chemical and biological properties was born<sup>83,84,87,88,89,90</sup>. This approach requires tip-sample interactions to be known, which is facilitated by functionalizing AFM tips with specific chemical groups or ligands<sup>88,89</sup>. FD curves then allow adhesion and mechanical strength of specific bonds formed between tip and sample to be measured<sup>91,92</sup>. Accordingly, FD-based AFM can map such specific forces while imaging the biological system<sup>63,93</sup>. Chemical tips can be obtained by functionalizing gold-coated tips with self-assembled alkanethiol monolayers terminated by specific functional groups<sup>87</sup>. Alkanethiols functionalized with nitrilotriacetate (NTA)-terminated groups that attach histidine-tagged biomolecules of interest have been used<sup>94</sup>. Silicon tips can be amino-silanized and reacted with PEG linkers, which carry benzaldehyde functions to attach peptides or proteins through lysine residues<sup>88</sup>.

Using functionalized probes, FD-based AFM could detect and localize specific interactions of biological systems ranging from antibodies to living human cells<sup>8,63,88,89,90,93,95</sup>. Biospecific FD-based AFM has proven useful to map receptor sites on animal cells. In an early work, AFM tips bearing the Helix pomatia lectin were used to map N-acetylgalactosamine-terminated glycolipids on group A red blood cells<sup>90</sup>. Since then, receptors mapped on animal cells include vitronectin receptors on osteoblasts<sup>96</sup>, prostaglandin receptors on CHO cells<sup>97</sup>, and glycosylphosphatidylinositol (GPI)-anchored proteins in neuronal membranes<sup>98</sup>. In another example human G-protein coupled receptors were imaged in membranes while measuring and mapping their single binding events of native and synthetic ligands<sup>99</sup>. By moving the AFM tip in a non-linear manner the unbinding forces of the ligands were measured over a very wide loading rate, which allowed the free-energy landscape of receptors binding to ligands to be reconstructed (**Fig. 3f**). Applied to live bacteria and yeast, the main components of microbial cell walls have been localized and force probed, including peptidoglycans<sup>46</sup>, teichoic acids<sup>100</sup>, and cell adhesion proteins<sup>83,101</sup>. These studies revealed the heterogeneous distribution of microbial cell surface molecules, which is related to the cell state. Additionally, the assembly machinery of bacteriophages was imaged on live bacteria and localized near the septum in soft nanodomains surrounded by the stiffer cell wall<sup>79</sup>. Whereas these applications functionalized the AFM tip with one type of biomolecule, a recent approach functionalized the AFM tip with two different ligands to map two binding sites of human G-protein coupled receptors<sup>102</sup>. Such application opens the door to AFM-based multifunctional recognition imaging.

A critical issue when analyzing adhesion forces detected by FD-based AFM is to proof their



specificity and to separate them from unspecific ones. Controls include blocking the specific interactions with antibodies or chemical compounds, as well as using mutant cells lacking the specific recognition sites. For direct comparison fluorescently labeled target and mutant cells may be co-cultured, identified by fluorescence microscopy and simultaneously imaged with the functionalized tip. Tip contamination is another problem that needs to be addressed. With complex samples like living cells, adsorption of loosely bound molecules may quickly change the functionalized tip, making the tip to record unknown interactions with the sample. Therefore, before engaging functionalized tips, it is useful to characterize the sample with unmodified tips. Also, one should always keep the applied force below 100 pN.

An alternative to FD-based AFM is TREC imaging, which records topography and specific recognition images at a similar speed as contact mode AFM<sup>103,104</sup>. This method was used to map the binding sites of cadherins on vascular endothelial cells<sup>105</sup>. TREC oscillates functionalized tips at very small (5–10 nm) amplitudes while scanning the sample. A specific binding event is then detected *via* an amplitude change. However, as FD curves are not recorded quantitative information of the molecular binding events is lacking.

### **Multifrequency imaging**

Besides topographic imaging AFM can map mechanical and functional properties of the biological sample. However, applying modes such as FD-based AFM considerably increases the data acquisition time<sup>63</sup>. Advanced dynamic mode AFM, including frequency or amplitude modulation, or multifrequency mode AFM offer higher frame rates. Recently developed multifrequency AFM modes<sup>37,106</sup>, which promise exciting possibilities to study biological systems are therefore discussed. Multifrequency AFM involves the simultaneous excitation and/or detection of several frequencies of the cantilever motion. These frequencies are usually associated with multiple integers (harmonics) of the fundamental frequency or intrinsic resonance frequencies (eigenmodes) of the cantilever<sup>37</sup>. There are several multifrequency AFM approaches<sup>37</sup>, however, their physical foundations can be quite complex and mostly their theoretical description is still under development. One key issue is to develop analytical expressions that relate the observables (amplitude, phase or frequency shifts) to material properties such as topography, flexibility, adhesion, stiffness, magnetic or electrostatic<sup>107</sup>. A straightforward explanation of how these methods operate is provided by bimodal AFM, which excites two eigenmodes of the cantilever and measures their observables (**Fig. 4a-b**). This combination of 1<sup>st</sup> and 2<sup>nd</sup> eigenmodes multiplies the number of observables to characterize the sample properties by a factor of two and requires just four data points per topographic pixel.

Bimodal AFM has been applied to measure different properties of DNA and proteins in liquid. The observation of the major and minor grooves of DNA in buffer is an example of the high-resolution provided by bimodal AFM (**Fig. 4c**). Topography and flexibility maps of a single IgM antibody have been acquired at a spatial resolution of  $\approx 2$  nm and showing that the uppermost part of the protein complex has an effective Young's modulus of 18 MPa while the antibody domains are much softer (8 Pa)<sup>37,59</sup>. Bimodal AFM has also been used to image ferritin while separating short-range mechanical ( $\approx 0.5$  nm) from long-range magnetic ( $\approx 5$ –1,000 nm) forces. The separation of mechanical forces provided by the stabilizing

protein shell and of magnetic forces of ferritin is possible because the 1<sup>st</sup> eigenmode is more sensitive to short-range repulsive forces while the 2<sup>nd</sup> eigenmode measured long-range interactions (**Fig. 4d**)<sup>108</sup>. Imaging water layers covering the chaperone GroEL at forces < 20 pN exemplifies the potential of bimodal AFM to provide novel insight about sample properties (**Fig. 4e**)<sup>109</sup>. Complementary to this frequency modulation AFM has also been applied to image hydration layers at the water-lipid interface of lipid membranes<sup>110</sup>.

Multiharmonics AFM excites the cantilever with a single frequency while recording multiple harmonics of the flexural or torsional cantilever motion. Initially, this AFM imaging mode has been applied to measure topography and viscoelastic properties of relatively large biological objects including viruses and cells (**Fig. 4e**)<sup>111,112</sup>. Torsional harmonics allow the topograph of the sample and the time-varying forces to be recorded by integrating the higher harmonics of the torsional movement. These forces quantify the mechanical properties of the sample, including Young's modulus or adhesion. Torsional harmonics also detect interactions in the  $\mu\text{s}$  range<sup>73</sup> and measure recognition forces of chemical groups or protein complexes (**Fig. 4g,h**)<sup>113</sup>. However, torsional harmonics AFM require the use of specially designed T-shaped cantilevers, which are not yet commercially available. This necessity together with the need to use complex algorithms to analyze the harmonics data is currently limiting wider application of the technique.

Accessing the sub-surface morphology of complex biological systems has been a longstanding challenge for AFM. Recently, ultrasonic microscopy and dynamic AFM have been combined to mechanically excite sample and cantilever, which generates mechanical waves that propagate through the biological sample. Waves mechanically interacting with the inside of the sample change amplitude and phase<sup>114,115</sup>. Thus, by using the AFM tip to probe these changes pixel-by-pixel can provide topography and structures beneath. This method shows potential for the imaging of embedded or buried sub-surface structures of animal and plant cells. However, currently sub-surface imaging requires the application of relatively large forces ( $\approx 100$  nN), which questions to which extent the structures imaged are representative of a native unperturbed cell. Additionally, the use of delocalized ultrasonic waves to generate images of sub-surface structures leaves interpretative challenges and limits the spatial resolution<sup>116</sup>. There is thus progress to be made before this AFM imaging mode will be applicable by a broad audience to address pertinent biological problems.

### High-speed imaging: imaging biological processes in real time

Compared to fluorescence microscopy, AFM imaging is limited by its rather slow time resolution. In the past years however tremendous technological advances allowed increasing the imaging speed, thus offering a means to study dynamic molecular processes by high-speed AFM (HS-AFM). Among AFM components, the slowest is the cantilever. Therefore, to achieve high-speed using amplitude modulation AFM, the cantilever's response time  $\tau = Q/(\pi f_0)$  has to be shortened, with  $Q$  being the quality factor and  $f_0$  the first resonance frequency of the cantilever in water (**Fig. 5a**). To increase  $f_0$ , while keeping the spring constant  $k$  small, small cantilevers (100–140 nm thick, 2–5  $\mu\text{m}$  wide and 9–14  $\mu\text{m}$  long) were developed, thereby approaching  $f_0 = 100$ –650 kHz and  $k = 0.1$ –0.3 N m<sup>-1</sup><sup>[70,71]</sup>. Because the  $Q$  value of these small cantilevers approaches  $\approx 2$  in water, their response time of  $\approx 1$ –6  $\mu\text{s}$  is 40–240-fold shorter than conventional cantilevers. Presently, small cantilevers with  $f_0 = 400$ –

800 kHz and  $k = 0.1\text{--}0.2 \text{ N m}^{-1}$  are commercially available. To achieve HS-AFM, it is also important to suppress mechanical vibrations of the Z-scanner that is moved at much higher frequencies than X- and Y-scanners (**Fig. 5b**). For this, three approaches were taken; counterbalancing the impulse generated by quick Z-scanner displacements<sup>71</sup>, designing robust scanner structures<sup>117,118,119</sup> and actively damping vibrations based on a Q-control technique (**Fig. 5c**)<sup>120</sup>. The last component to be noted is a controller that can dynamically tune the feedback gains during imaging to minimize the tip-sample force (**Fig. 5c,d**)<sup>121</sup>. The highest possible imaging rate of HS-AFM as a function of various parameters is quantitatively described elsewhere<sup>122</sup>.

In the early days of HS-AFM, DNA<sup>70</sup>, GroEL-GroES<sup>123</sup> and myosin V<sup>71,124</sup> were observed to evaluate the performance of newly developed devices. Recently, HS-AFM provided unique mechanistic insight into the function of bacteriorhodopsin<sup>125</sup>, myosin V<sup>126</sup>, F<sub>1</sub>-ATPase<sup>127</sup>, ESCRT-III<sup>128</sup> and nuclear pore complexes<sup>129</sup>. HS-AFM images of the light-driven proton pump bacteriorhodopsin showed that upon light illumination the cytoplasmic E-F helix portion of each bacteriorhodopsin displaces outwards by  $\approx 0.7 \text{ nm}$  and contacts bacteriorhodopsins from adjacent trimers (**Fig. 5e**)<sup>125</sup>. Myosin V processively walks along actin filaments in a handover-hand manner, resulting in a  $\approx 36 \text{ nm}$  step for every ATP hydrolyzed. HS-AFM observations of myosin V interacting with actin provided a direct observation of the process, and visualized the lever-arm swing, which had been hypothesized for a long time (**Fig. 5f**)<sup>126</sup>. The results suggested that myosin V steps forward without transitioning through an ADP-Pi bound state, and hence, that the actin-myosin binding energy is harnessed to generate the lever-arm swing.

In the rotary motor F<sub>1</sub>-ATPase, the  $\gamma$  subunit rotates in the stator  $(\alpha\beta)_3$  ring upon ATP hydrolysis in the catalytic sites mainly located in the  $\beta$  subunits. This rotation is made possible by rotary propagation of three chemical states (empty, ATP-bound and ADP-bound states) and hence corresponding structural states over the  $\beta$  subunits. HS-AFM visualization of  $\gamma$ -less  $(\alpha\beta)_3$  rings revealed that the three states can propagate without the  $\gamma$  subunit (**Fig. 5g**)<sup>127</sup>. So, the  $\beta$ - $\beta$  interplay through the  $\alpha$  subunits engenders this cooperativity, ruling out a previous  $\gamma$ -dictator model that the cooperativity would be caused by different  $\gamma$ - $\beta$  interactions for the three  $\beta$  subunits because of an asymmetric structure of the  $\gamma$  subunit.

Snf7, an endosomal sorting complex required for transport (ESCRT-III), plays a key role in lipid membrane budding and abscission. HS-AFM of Snf7 placed on supported planar lipid bilayers showed concentric spiral filaments (**Fig. 5h**)<sup>128</sup>. Upon disrupting large spirals with the cantilever tip, the broken polymers spontaneously formed smaller rings, suggesting a preferred diameter of 25 nm for Snf7 as well as “unbending” of the spiral filaments from their natural curvature. Thus, it was proposed that in cellular conditions energy would be accumulated during the growth of the spiral spring and eventually released through shrinking of the spiral diameter and buckling of the inner spirals, which would cause the membrane to buckle, bud and abscise.

Nuclear pore complexes (NPCs) facilitate the molecular exchange between cytoplasm and nucleus in eukaryotic cells. However, how nucleoporins form a selective barrier facilitating this transport has been unclear. Applying HS-AFM it became possible to visualize the spatiotemporal dynamics of nucleoporins inside NPCs of *Xenopus laevis* oocytes at timescales of 100 ms<sup>129</sup>. It was observed that the cytoplasmic orifice is circumscribed by

highly flexible, dynamically fluctuating nucleoporins that rapidly elongate and retract. This transient entanglement in the NPC channel manifests as a central plug when averaged in space and time.

Beside these molecular studies, HS-AFM has also been successfully used to observe dynamic processes of live bacteria<sup>130,131</sup> and eukaryotic cells<sup>132</sup>. However, HS-AFM has long relied on scanning the sample-stage, which excludes the use of large heavy sample stages and makes it difficult to combine with optical microscopy. The tip-scan HS-AFM developed very recently will thus significantly expand the applicability to study biological processes by AFM<sup>133</sup>. Observations, for example, living cells cultured in Petri dishes, membrane proteins in suspended membranes, or proteins responding to external forces applied by optical tweezers, will become possible. Cell biological applications most of which require the combination of AFM and sophisticated optical techniques (next chapter) will be made easier. It is also possible to transfer this knowledge to high-speed SICM for studying dynamics of live cells and isolated intracellular organelles.

### Correlative imaging

Living cells present a high level of structural and functional complexity. Cell surfaces consisting of thousands of different macromolecules represent a small heterogeneous and dynamic portion of the cellular complexity<sup>134</sup>. It is thus challenging to identify even simple cell surface structures such as receptors, channels, transporters or assemblies thereof in topographs recorded by AFM. In such cases the full potential of AFM is achieved in combination with complementary microscopy techniques that can identify and correlate complex cellular structures of interest<sup>9</sup>. These complementary techniques include optical microscopy, fluorescence microscopy, confocal microscopy, FRET, TIRF, or super-resolution microscopy. In most cases AFM has been adapted to fit to optical microscopes. Environmental chambers allowing cellular systems to be kept in their close-to-native state had to be engineered (**Fig. 6a**). Nowadays, such multimicroscopic combinations allow the unique characterization of a wide range of complex biological systems ranging from membranes, cells to tissues.

A popular combination of AFM is either with epifluorescence or confocal microscopy. Exciting applications range from single animal cells, to tissues microbial cells, and to their assemblies. In such studies, structures of interests were fluorescently labeled, optically imaged at  $\mu\text{m}$  resolution and correlated to AFM topographs contoured at nm precision. These approaches identified hitherto unknown supramolecular assemblies of cell surface structures and contributed to the understanding of their function. For example, various steps of the interaction between fungal pathogens and macrophages were captured, including initial cellular contact, fungal cell internalization, and hyphal elongation resulting in membrane piercing and escape from the macrophage. While fluorescence imaging distinguished fungal cells from macrophages, AFM revealed biological relevant nanostructures on both cell types (**Fig. 6b-c**)<sup>135</sup>. The AFM has also been used to image cell surface structures including microvilli, actin ridges and nanodomains of cellular membranes and to characterize their dynamic mechanical properties (**Fig. 6d**)<sup>98,136,137</sup>. Optical microscopy is frequently applied to characterize cell morphology and state while employing AFM to characterize the mechanical properties (e.g., stiffness, elasticity, pressure) of the cell or its

mechanical interaction (e.g., adhesion, migration) with the environment<sup>9</sup>. Such experiments allowed the furrow stiffening during cell division<sup>65</sup> to be observed, the adhesion of *Dictyostelium discoideum* to their substrate to be measured to molecular scale<sup>138</sup>, or to unravel whether cell adhesion or cortex tension determine cell sorting in the developing embryo<sup>139</sup>. Importantly, some of the experiments contributed answers to controversial debate lasting for more than three decades. Combined AFM and confocal microscopy was used to monitor angiotensin-induced contractile response and cytoskeleton remodeling in human embryonic kidney cells<sup>140</sup>. Other examples used confocal microscopy to monitor eukaryotic cells transiently expressing GFP-actin, tubulin, vimentin and LaminA and imaged the mechanical properties of cytoskeleton and nucleus during early apoptosis<sup>141</sup>. AFM was also applied to measure the cell pressure and cortex tension while quantifying the actin and myosin accumulating at the cell cortex by confocal microscopy (**Fig. 6e-g**)<sup>142</sup>. The latter approach contributed to the understanding of how adherent animal cells facilitate and regulate their rather drastic cell shape changes required to progress through mitosis<sup>143</sup>.

As discussed above, cantilevers functionalized with biological molecules, chemical groups or even with living cells can reveal specific sites and their interactions on live cells<sup>9</sup>. Applying molecular recognition AFM in conjunction with optical microscopy can reveal a comprehensive picture of the distribution of cell surface receptors and of cell morphology and state. Recent examples include the localization of receptors on CHO cells and endothelial cells<sup>144</sup>, and the visualization of the peptidoglycan insertion into the cell wall of *L. lactis*<sup>46</sup> while mapping the distribution of single peptidoglycan molecules on the outermost cell surface using the AFM. Molecular recognition AFM and fluorescence microscopy also linked the spatial localization and functional role of cell wall teichoic acids in *Lactobacillus plantarum*<sup>100</sup>. Polarized cell-wall organization was found to play a key role in controlling cell morphogenesis. In yeast, both AFM recognition imaging and confocal microscopy demonstrated that Als adhesion proteins form nanodomains on live cells through amyloid interactions<sup>145</sup>. Very recently, AFM tips functionalized with single rabies viruses were used to correlate fluorescence images of cell surface receptors to viral binding events to the animal cell<sup>146</sup>. Analysis of the initial binding events revealed that the viral glycoproteins bind cell surface receptors in an allosteric mode until all three binding sites of the trimeric cell surface receptor are occupied and viral fusion can be initiated.

## Conclusions

This year we are celebrating the 30<sup>th</sup> birthday of AFM, which undoubtedly has revolutionized nanotechnology and now shows a considerable impact in life sciences. Here we highlighted the wealth of AFM-based modalities that have been implemented over the years, opening the door towards the multiparametric and multifunctional characterization of biological systems. These methods range from the high-resolution imaging of native biostructures and the simultaneous mapping of mechanical, kinetic and thermodynamic properties, of functional groups and binding sites, of free energy landscapes of ligand-receptor bonds, or of electrostatic properties ranging from charge distributions to ion currents. In the past years many new AFM-imaging modalities have been introduced, which in principle can be readily applied to biological systems and thus will further extend the variety of information that can be quantified and structurally mapped while imaging complex biological systems.

Currently, force sensitivity and thermal stability (drift) of AFM limit the precision at which biological systems can be imaged and manipulated. It may be thus expected, that recently introduced ultrastable AFMs providing sub-pN force precision and high positional stability ( $< 0.03 \text{ \AA}$ ) at extremely low lateral drift ( $\approx 5 \text{ pm min}^{-1}$ )<sup>147,148</sup>, will guide the development of AFMs for new applications of biological significance. Today most bio-AFM users apply single AFM-imaging modalities in their specific field of interest. However, biological systems are rather complex and require the acquisition of a wealth of information to be understood. Therefore, we foresee that in the near future many of the AFM-modalities discussed here will be combined into one instrument and thus into one set of correlated measurements. Such multimodal, multiparametric, multifrequency, and high-speed AFM imaging platforms will guide us towards a more comprehensive understanding of the dynamic, structural, mechanical, chemical and functional heterogeneity of complex biological systems. Together with advances in complementary techniques (**Table 1**), this will thrive the use of AFM to address outstanding questions in biology in the next decades.

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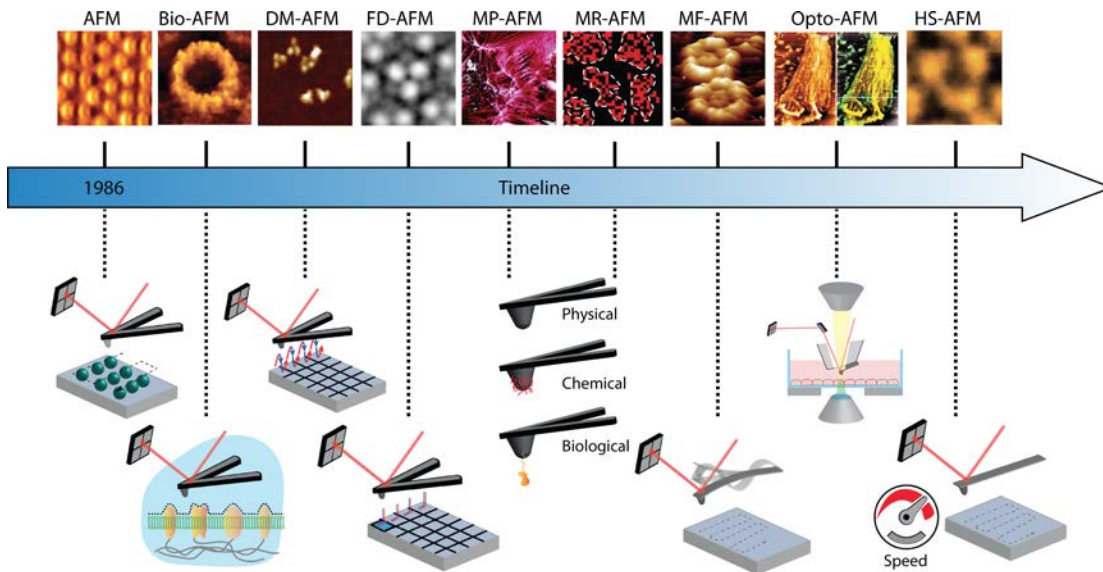
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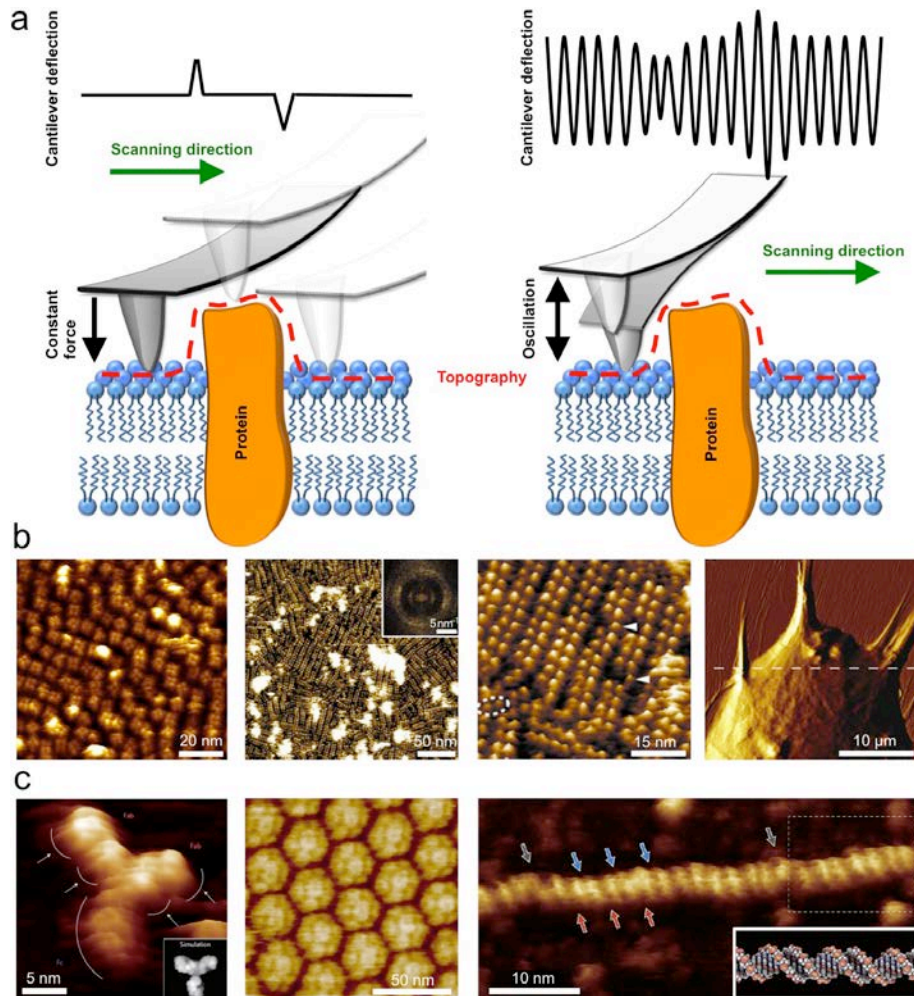
## Acknowledgements

Y.F.D. was supported by the Université catholique de Louvain, the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme (grant agreement No [693630]), the WELBIO (Grant n°WELBIO-CR-2015A-05), the National Fund for Scientific Research (FNRS), 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). D.A. and D.M.M. were supported by the European Molecular Biology Organization (EMBO; ALTF 265-2013 and ALTF 506-2012). D.A. and Y.F.D. are Research Associate and Research Director at the FNRS, respectively. D.J.M. was supported by the Swiss National Science Foundation (SNF; Grant 310030B\_160225 to D.J.M.), the NCCR Molecular Systems Engineering and the Swiss Commission for Technology and Innovation (CTI, Grant 17970.1). R.G. acknowledges financial support from the European Research Council AdG no. 340177 and the Ministerio de Economía y Competitividad MAT2013-44858-R. T.A. was supported by the Japan Society for the Promotion of Science (JSPS; Grants 24227005 and 26119003) and by the Japan Science and Technology Agency (JST; CREST program on Structural Life Science and Advanced Core Technology for Innovative Life Science Research).

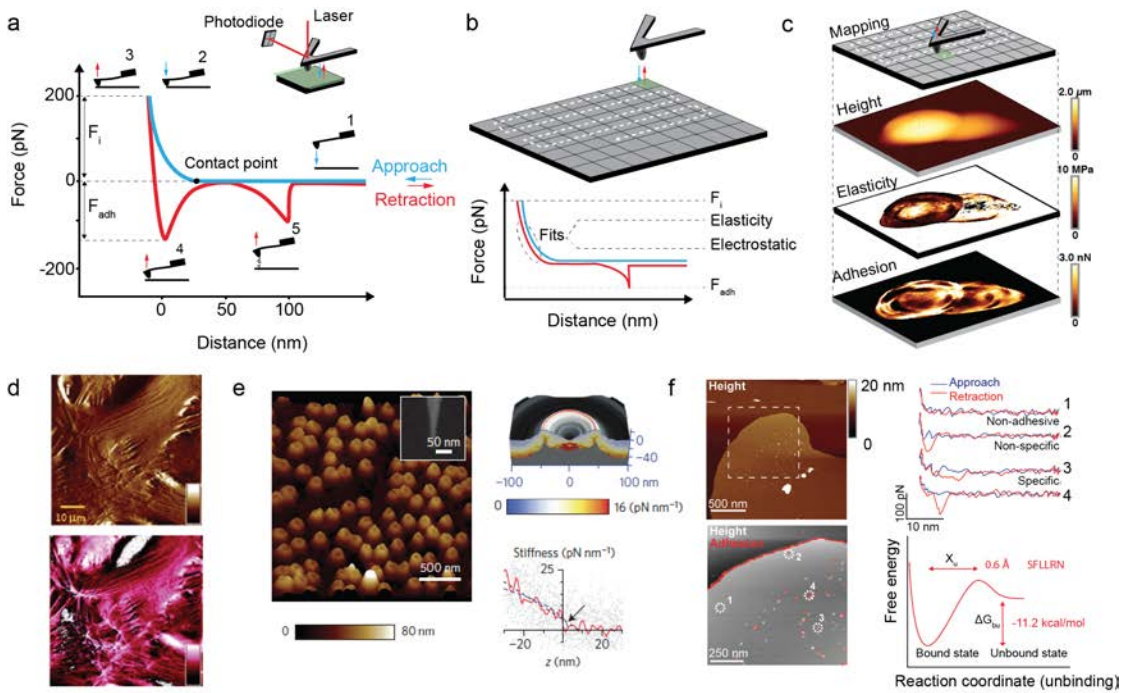
## Figures and legends



**Figure 1 | Timeline of key inventions, starting from the birth of atomic force microscopy (AFM) in 1986 to the latest AFM imaging modalities in molecular and cell biology.** Key inventions developed over the years include: (i) optical detection system and fluid cell enabling to operate contact mode AFM in aqueous solution (Bio-AFM); (ii) dynamic mode AFM (DM-AFM) which oscillates the AFM tip to reduce friction while contouring the biological sample; (iii) force-distance curve-based AFM (FD-AFM) contours the surface of a biological system while recording pixel-by-pixel a full FD curve; (iv) multiparametric AFM (MP-AFM) contours the sample while mapping multiple physical or chemical properties; (v) molecular recognition AFM (MR-AFM) images and maps specific interactions of biological samples; (vi) multifrequency AFM (MF-AFM) contours the sample while oscillating the cantilever tip at multiple frequencies, thus mapping various physical parameters; (vii) correlating advanced optical imaging and AFM (Opto-AFM) for the imaging of complex biological systems; (viii) high-speed AFM (HS-AFM) speeds up the image acquisition time by a factor of  $\approx 1,000$ , providing access to dynamic processes in biology. Most modalities cross-fertilized each other ultimately leading to combinatorial AFM. Images adapted from refs.<sup>22,41,74,77,78,125,149</sup> with author permission and from refs. xx, xx with permission from XXX.

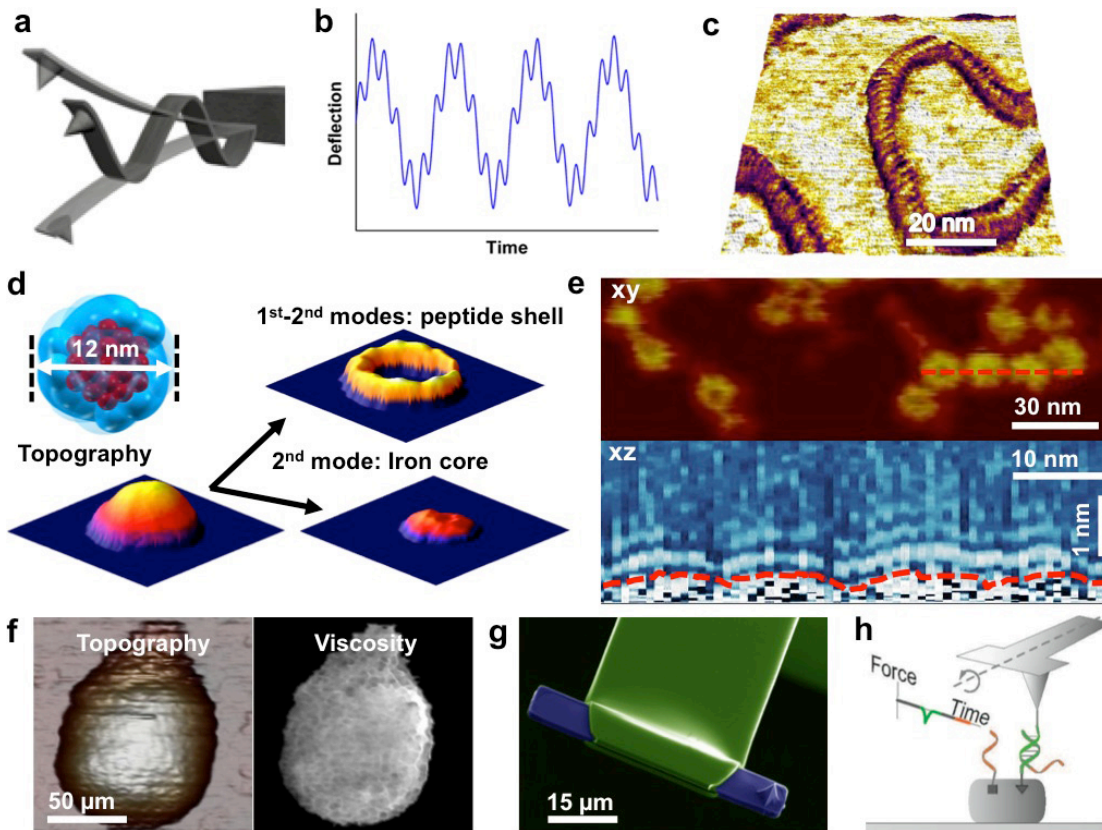


**Figure 2 | AFM-based imaging of native biological systems to molecular resolution.** **a**, Basic principles of contact (left) and dynamic (right) AFM imaging modes. In contact mode the cantilever deflection is kept constant (constant force) by adjusting the relative height between tip and sample. A topographic height change alters the cantilever deflection, which a feedback loop corrects by adjusting the tip-sample distance. The dynamic mode oscillates the cantilever close to or at resonance frequency. Height changes alter the cantilever oscillation, which is used to adjust the tip-sample distance. **b**, Contact mode AFM topographs. Left, cyclic nucleotide-regulated potassium channels (MlotiK1) reconstituted into lipid membranes. Middle, rows of rhodopsin dimers distributed in the native disc membrane extracted from rod outer segments of the eye. Right, image of a living SAOS-A2 cell bundling and pulling collagen fibrils coating a substrate. To maximize contrast, the exemplified image shows the deflection of the cantilever, which changes while contouring the sample. **c**, Dynamic mode AFM topographs. Left, an IgG antibody absorbed to mica and visualized with frequency modulation mode. Middle, single bromo mosaic viruses packed in a crystalline assembly. Right, circular plasmid DNA imaged in buffer solution by frequency modulation AFM. Images adapted from refs.<sup>24,40,41,150,151,152</sup> with permission from XXX.

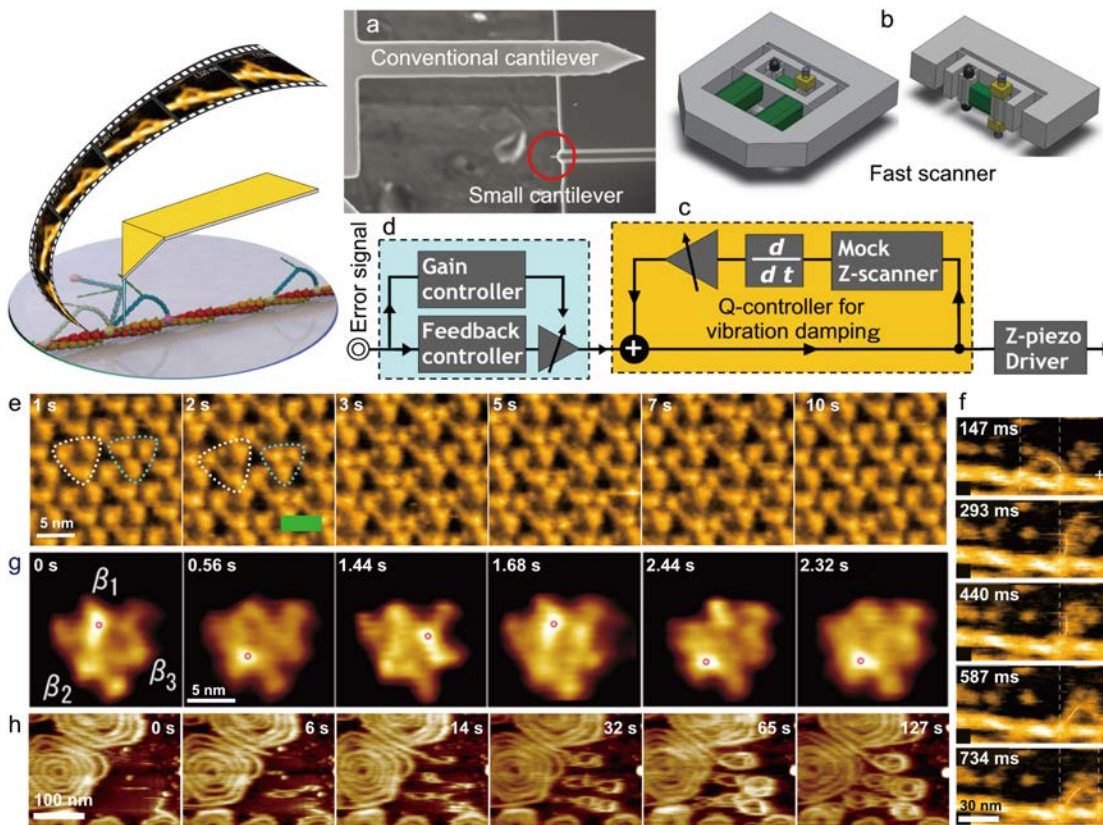


**Figure 3 | Force-distance curve-based AFM.** **a**, Principle of recording force-distance (FD) curves by approaching (blue) and withdrawing (red) the AFM tip from the sample. The tip of the cantilever is initially away from the sample (1) to which it is brought into contact (2). During retraction (3) of the AFM tip, adhesive events may occur at different distances due to non-specific (4) or specific (5) interactions between tip and sample. **b**, FD-based AFM imaging records pixel-by-pixel FD curves while contouring the sample topography. The indentation force  $F_i$  is controlled and parameters extracted include the tip-sample adhesion force  $F_{adh}$ , or elastic and electrostatic properties (by fitting the curve). Parameters can be displayed as colored maps and correlated to the topography. **c**, Example of multiparametric FD-based AFM imaging of the elasticity and adhesion of two dividing *S. aureus* cells. **d**, AFM force error and elasticity map of living HaCaT keratinocytes. **e**, Topography (brown colored) and stiffness map (upper right inset) of nuclear pore complexes (NPCs) from the cytoplasmic surface. The graph shows the stiffness as a function of tip-sample separation recorded close to the center of the cytoplasmic ring. **f**, Top left, topograph of human protease activated receptors 1 (PAR1) in proteoliposomes recorded with a SFLLRN-ligand functionalized tip. Bottom left, overlay of topograph (grey) and adhesive interactions (red) localizes individual receptors binding the ligand. Top right, force-distance curves exemplifying unspecific adhesion events (1 and 2) and specific ligand-receptor unbinding events (3 and 4) showing the stretching of the linker tethering the ligand to the AFM tip. Right bottom, free energy landscape of the ligand binding to PAR1 extracted from measuring the rupture force of the ligand-receptor bond at different loading rates. Images adapted from refs.<sup>77,78,99,153</sup> with permission from XXX.

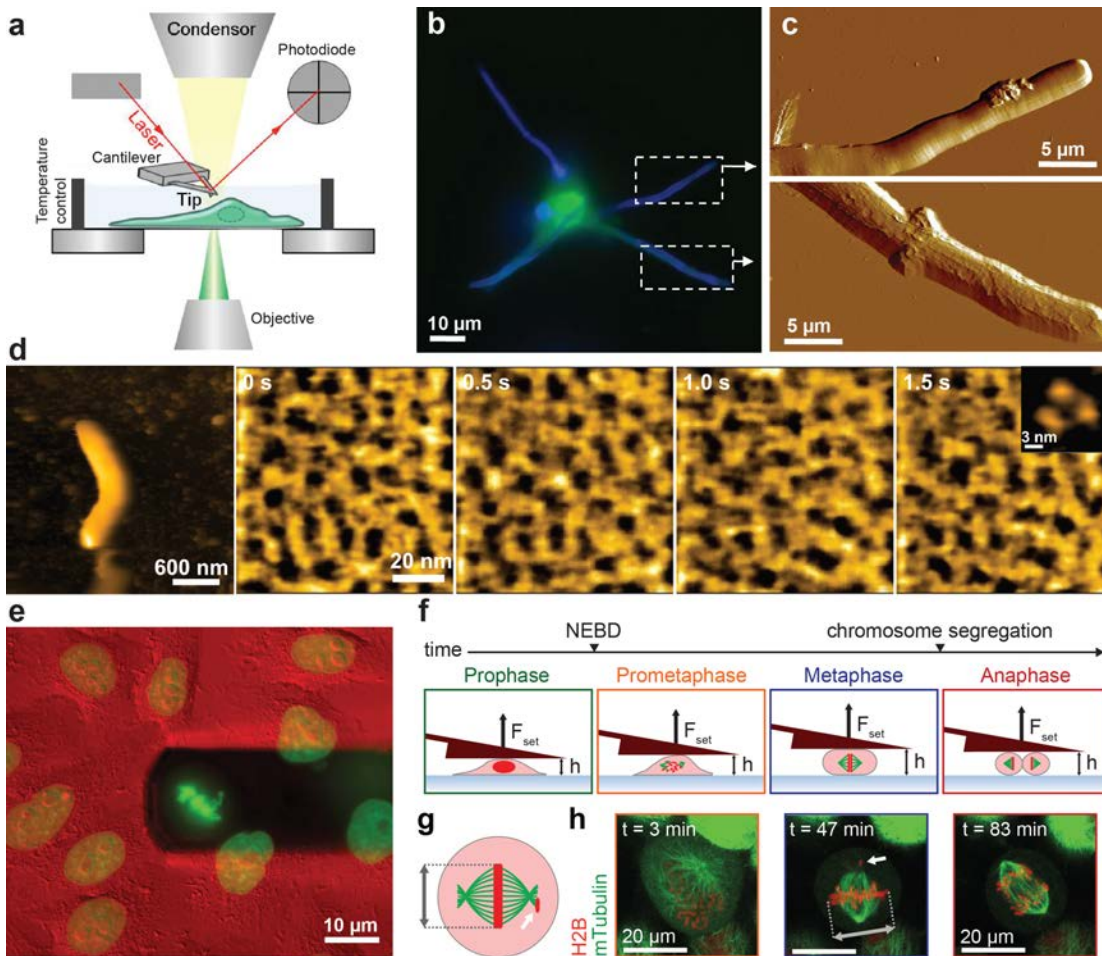




**Figure 4 | Multifrequency AFM.** **a**, Scheme of the deflection of the cantilever in bimodal AFM. **b**, Two eigenmodes of the cantilever are excited and detected. Observables associated with both eigenmodes are recorded to determine sample properties such as flexibility, deformation, and viscosity. **c**, Bimodal AFM image of a DNA in buffer showing the major and minor grooves of the double helix. **d**, Separation of short-range mechanical forces and long-range magnetic interactions in ferritin. The 1<sup>st</sup> eigenmode contours the topography while the 2<sup>nd</sup> eigenmode detects long-range magnetic forces. By combining both signals, the iron oxide core and the apoferritin shell is separated in the AFM image. **e**, The top panel shows an AFM topograph (*xy* frame) of GroEL proteins. The bottom panel shows a vertical profile (*xz* frame, taken along the red dashed line of the topography) of the hydration layers contouring four GroEL molecules. The dashed red line marks the surface of the GroEL molecules. **f**, Multifrequency flexural AFM of a bacteriophage  $\Phi 29$  mature virion. The virion topography is acquired simultaneously with multi-harmonic observables images, from which the viscosity map is shown. Images were recorded applying 100 pN. **g**, False color electron microscopy image of a T-shaped cantilever designed for torsional harmonics AFM. **h**, Multifrequency torsional harmonics scheme for probing chemical groups of a protein using DNA labels. A DNA strand attached to the tip interacts with target DNA strands. Complementary sequences have identical colors. Images adapted from refs.<sup>109,111,113,154</sup> with permission from XXX.



**Figure 5 | HS-AFM filming proteins in action.** **a-d**, Key devices and techniques for HS-AFM. **a**, Small cantilever with high resonant frequency and small spring constant. **b**, Fast scanner suppressing impulses generated by quick displacements of piezoelectric X- and Z-scanners. **c**, Active vibration damping based on Q-control with mock Z-scanner. **d**, Feedback controller with automatic gain tuning for low-invasive high-speed imaging without causing tip-parachuting. **e-h**, HS-AFM images of proteins in action. **e**, Bacteriorhodopsin in native purple membrane recorded under dark and illumination at 1 frames per second (fps). White triangles indicate bacteriorhodopsin trimers. Blue triangles indicate “trefoils” that comprises three bacteriorhodopsin monomers, each belonging to an adjacent trimer. Green light was illuminated at 2 s and switched off at 3 s. Upon illumination, bacteriorhodopsin trimers dilate outwardly, while bacteriorhodopsin monomers contact each other in trefoils. **f**, Myosin V walking unidirectionally along an actin filament, showing forward rotation of the leading lever-arm upon trailing head detachment from actin. **g**, Rotor-less F<sub>1</sub>-ATPase undergoing conformational changes. Red circles indicate the highest positions of the topographs. Since a nucleotide-free  $\beta$ -subunit protrudes higher than ADP- and ATP-bound ones, it is observed that the unbound state rotates counterclockwise. **h**, Spiral filament formation by polymerization of the ESCRT-III protein Snf7 on a supported lipid membrane. Images adapted from refs.<sup>125,126,127,128</sup> with permission from XXX.



**Figure 6 | AFM of cellular systems.** **a**, Schematic setup of an AFM combined with optical microscopy for the characterization of living cells. Fluorescence image (**b**) and correlative AFM images (**c**) of a macrophage (green) incubated for 3 h with cells from *Candida albicans* (blue). Images in panel **c** are enlarged views of the dashed areas shown in the fluorescence image. Internalized (bottom) and externalized (top) hyphae featuring major structural differences. **d**, HS-AFM topographs of the *E. coli* bacterium. The first topograph shows the entire bacterium, while the following images of the outer membrane show moving net-like structures formed by porin trimers. The last inset shows a single porin trimer. **e**, Mechanical confinement and morphological characterization of mitotic animal cells. The image shows overlaid differential interference contrast (DIC) and histoneH2B green fluorescent protein (H2B-GFP) images of mitotic HeLa cells. The cantilever (dark shadowed) confines a single mitotic cell in the metaphase to measure the force and pressure generated by the rounding animal cell. **f**, Wedged cantilever applied to confine a mitotic HeLa cell and to mechanically control mitotic progression. Depicted are mitotic phases, spindle microtubules (green), chromosomes (red), nuclear envelope breakdown (NEBD), time ( $t$ ), and force ( $F_{set}$ ) and height ( $h$ ) set by the cantilever. (**g**) Top view of spindle characterization scheme of a confined HeLa cell and (**h**) fluorescence snapshots of microtubules (mTubulin-GFP) and chromosomes (H2B-mCherry). Gray double arrows, metaphase plate width. White arrows, stray chromosomes.  $t = 0$ , NEBD. Images adapted from refs.<sup>54,131,135,143</sup> with permission from XXX.