Nanoscale monitoring of drug actions on cell membrane using atomic force microscopy

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Knowledge of the nanoscale changes that take place in individual cells in response to a drug is useful for understanding the drug action. However, due to the lack of adequate techniques, such knowledge was scarce until the advent of atomic force microscopy (AFM), which is a multifunctional tool for investigating cellular behavior with nanometer resolution under near-physiological conditions.

In the past decade, researchers have applied AFM to monitor the morphological and mechanical dynamics of individual cells following drug stimulation, yielding considerable novel insight into how the drug molecules affect an individual cell at the nanoscale. In this article we summarize the representative applications of AFM in characterization of drug actions on cell membrane, including topographic imaging, elasticity measurements, molecular interaction quantification, native membrane protein imaging and manipulation, etc. The challenges that are hampering the further development of AFM for studies of cellular activities are also discussed.

Keywords: atomic force microscopy; single-cell technique; nanometer scale; drug-target interaction; cell mechanics; cell elasticity

Introduction

The pharmaceutical industry is currently facing unprecedented challenges, primarily due to the increasing cost of developing new drugs, the sharply rising attrition rates and the dissipation of proprietary products. Therefore, innovations are urgently needed for promoting the development of the pharmaceutical industry. Modern drug discovery is primarily based on the search for drug leads against a preselected therapeutic target followed by subsequent testing of the derived drug candidates. Approximately half of the approved drugs target human membrane proteins on the cell surface, such as G-protein coupled receptors (GPCRs), ion channels and antigen molecules. In the process of drug discovery (e.g., target validation, lead optimization and preclinical testing), researchers perform complex experiments to demonstrate the efficacy and safety of a candidate drug by using various biochemical approaches (e.g., Western blotting, X-ray crystallography and radioimmunoassay). Although valuable, these assays provide averaged information obtained on large ensembles of molecules contained in many cells, thereby masking the rare events that take place in single cells. Recent studies have shown that even when derived from a seemingly homogenous population, individual cells can exhibit substantial differences in their gene expression, protein levels, phenotypic output and function. It has become increasingly apparent that obtaining information at the single-cell level is of central importance for revealing the novel regulatory mechanisms of diverse cellular processes. Single-cell approaches also provide opportunities to investigate many important biological processes that cannot be assessed by ensemble experiments, such as monoallelic gene expression, lineage tracing during cell differentiation and biofilm formation in bacteria. Therefore, it can be reasonably anticipated that advances in single-cell techniques will help the pharmaceutical industry to partly overcome its current challenges. However, for potential practical application in drug discovery, single-cell techniques that can rapidly and automatically obtain the multiple parameters of individual living cells with high spatial resolution are needed.

The invention of atomic force microscopy (AFM) has provided an excellent means for observing, manipulating and exploring the functional components of single living cells with nanometer resolution. Compared with the commonly used biochemical methods for assessing drugs, AFM has clear advantages. For example, optical microscopy cannot visualize nanoscale cellular structures and electron micro-
copy requires that cells be fixed and dried. By contrast, AFM can be used to observe living cells in aqueous solutions with nanometer resolution, so that the dynamic changes of cellular ultra-microstructures on single cells in response to drugs can be monitored by time-lapse AFM.[14] Additionally, the drawback of Western blotting, polymerase chain reaction and X-ray crystallography is that they require the extraction of target molecules from cells, which destroys the intrinsic structures of the cells. By contrast, AFM can image the integral membrane proteins in their native membrane environment at submolecular resolution[15], allowing the direct observation of drug-induced morphological changes in target molecules[16, 17]. Traditional methods (eg, surface plasma resonance, radioimmunoassay) that quantify the binding constant of a drug to its target molecule cannot reveal the molecular dynamics taking place during the association and dissociation processes of individual drug-target interactions[18]. By contrast, AFM can directly reveal the detailed dynamics of the dissociation process (eg, how many energy barriers are overcome during dissociation) by simply pulling a drug molecule from its target molecule with different pulling rates[19, 20]; therefore, there is no need to model the dynamic processes of drug-target interactions using complex molecular dynamics simulations[21]. In addition, recent advances (eg, high-speed AFM[22], peak force tapping AFM[23]) in AFM have increased the power of the technique. Currently, AFM can simultaneously obtain multiple physicochemical parameters (such as morphology, elasticity, adhesion, deformation and energy dissipation)[23] of biological systems in a relatively short amount of time (approximately several minutes), allowing the nanoscale structures, properties and functions of cells/targets in response to drugs to be correlated.

In the past decade, AFM has emerged as a valuable addition to the pharmaceutical industry’s toolkit[24] and has been demonstrated to be useful in assisting drug evaluations and assays (eg, drug-target interactions[25], target molecules[26, 27] and drug delivery system characterizations[28, 29]). The applications of AFM in pharmacology have provided novel fundamental insight for understanding drug actions[30]. The increased knowledge obtained through AFM will be significant for making improvements to the current drug discovery process[31]. Here, based on the authors’ own research on AFM investigations of cell-drug interactions, the principle of AFM imaging and representative quantitative measurements will be first presented, followed by a summary of the typical applications (eg, topographic imaging, elasticity measurements, molecular interaction quantification, native membrane protein imaging and manipulation) of AFM in assessing drug actions. Finally, the current challenges and future directions of this field will be discussed.

**AFM imaging and measurements: principle and methods**

The main components of an AFM include a piezoelectric tube driver, which is connected to a flexible cantilever with a sharp tip mounted at its end, a laser photodiode detector and feedback electronics, as shown in Figure 1A. High-resolution imaging of the topography of active biological samples (eg, living cells, native molecules) is the most important application of AFM in the life sciences. The two modes of AFM commonly used are contact mode and tapping mode. In contact mode (Figure 1Ai), the tip scans over the sample surface while changes in the cantilever deflection are monitored by the photodiode detector (the detector monitors a beam of laser light reflected from the backside of the cantilever). The cantilever is driven vertically by the piezoelectric tube to maintain a constant deflection according to the feedback electronics, thereby maintaining a constant loading force on the sample. In tapping mode (Figure 1Aii), the cantilever oscillates near its resonance frequency and is scanned over the sample surface. In liquid, the driving frequency of the cantilever is often approximately 10 kHz. The piezoelectric tube drives the cantilever to move vertically to maintain a set constant amplitude.

![Figure 1](https://www.nature.com/aps)

**Figure 1.** Using AFM to investigate the drug actions on single cells. (A) The main components of an AFM. Contact mode (i) and tapping mode (ii) are the two commonly used imaging modes. (B) Measuring the elastic properties of cells using AFM indenting technique. The cellular Young’s modulus is calculated from the approach curve. (i) Optical and SEM images of a sphere probe[49]. (ii) Schematic diagram. (C) Measuring the binding force of a drug molecule to the target molecule by pulling a drug molecule from its target molecule[54]. The binding force is calculated from the retract curve which has a specific unbinding peak (denoted by the green arrow).
According to the feedback electronics. Presently, contact mode is the more widely used imaging mode. Contact mode is particularly suitable for obtaining high-resolution images from living adherent cell imaging and native protein imaging. A notable point regarding cell imaging using contact mode is that the scan force between the tip and cells should be low enough to prevent sample damage. For suspended mammalian living cells, however, tapping mode is adequate, because suspended mammalian cells cannot spread on the substrate and have soft surfaces. Tapping mode can significantly diminish the influence of lateral forces during the scanning, so that there is no scraping of the sample. Moreover, tapping mode is suitable for some special applications, such as high-speed scanning and multiparametric imaging. Under near-physiological conditions (e.g., cell culture medium, buffered solution), the dynamic changes of cellular ultra-structures following the addition of drugs can be visualized to directly assess their influence on individual cells. By immobilizing cell membrane patches onto the supported substrate (e.g., mica), the morphological changes of single target molecules in response to the drug molecules can also be investigated.

The elastic properties of samples are measured by controlling the AFM tip to indent the samples, as shown in Figure 1B. For measuring the elastic properties of cells, the tip performs approach-retract cycles on the cell surface. During the process, the deflection of the cantilever and the movement of the piezo tube are recorded to produce so-called force curves. Figure 1B shows a schematic diagram of a force curve obtained during the approach-retract cycles. A force curve contains two components: an approach curve and a retract curve. The approach curve is used to calculate the cellular Young’s modulus, and the retract curve is used to obtain the Young’s modulus of the sample. The contact point, which leads to measurements often being performed on the central region of the cell. Researchers have shown that the JKR model can be applied in the case of large forces between the contact surfaces (e.g., electrostatic forces, adhesion forces, and friction forces) and is therefore applicable when the forces between the contact surfaces are negligible compared with the maximum load. By contrast, the Johnson-Kendall-Roberts (JKR) model and Derjaguin-Muller-Toporov (DMT) model take into account the adhesion forces between the contact surfaces.

The JKR model considers the adhesion forces inside the contact area whereas the DMT model considers the forces outside the contact area. Researchers have shown that the JKR model can be applied in the case of large tips and soft samples with a large adhesion, whereas the DMT model is applicable in the case of small tips and stiff samples with a small adhesion. The conventional cone tip is sharp with a small opening angle of the tip, and the DMT model is applicable in the case of large tips and stiff samples with a large adhesion. The JKR model is applicable in the case of large tips and soft samples with a large adhesion. The DMT model is applicable in the case of small tips and stiff samples with a small adhesion. The conventional cone tip is sharp with a small opening angle of the tip, and the DMT model is applicable in the case of large tips and stiff samples with a large adhesion.

The Hertz-Sneddon model is based on the following expressions:

\[
F_{\text{cone}} = \frac{2E\delta \tan \theta}{\pi} \\
F_{\text{sphere}} = \frac{4ER^{1/2}\delta^{3/2}}{3} \\
\frac{1}{E} = \frac{1 - \nu_{\text{tip}}^2}{E_{\text{tip}}} + \frac{1 - \nu_{\text{cell}}^2}{E_{\text{cell}}} \\
\frac{1}{R} = \frac{1}{R_{\text{tip}}} + \frac{1}{R_{\text{cell}}} 
\]

where \(\nu_{\text{tip}}\) and \(\nu_{\text{cell}}\) are the Poisson ratios of the tip and cell, respectively, \(F\) is the loading force of tip, \(\delta\) is the indentation depth, \(E\) is the effective Young’s modulus, \(E_{\text{tip}}\) is the Young’s modulus of tip, \(E_{\text{cell}}\) is the Young’s modulus of cell, \(\theta\) is the half-opening angle of the tip, \(R\) is the effective radius, \(R_{\text{tip}}\) is the radius of tip and \(R_{\text{cell}}\) is the radius of cell. Because the Young’s modulus of living cells is much smaller than that of the tip, \(E_{\text{cell}}<<E_{\text{tip}}\), formula (3) can be rewritten as:

\[
\frac{1}{E} = 1 - \nu_{\text{cell}}^2 \frac{E}{E_{\text{cell}}} 
\]

Based on this approximation, formulas (1) and (2) can be recast as:

\[
F_{\text{cone}} = \frac{2E_{\text{cell}}\delta^3 \tan \theta}{\pi (1 - \nu_{\text{cell}})} \\
F_{\text{sphere}} = \frac{4E_{\text{cell}}R^{1/2}\delta^{3/2}}{3(1 - \nu_{\text{cell}})} 
\]

Living cells are often considered to be incompressible materials; therefore, the value of \(\nu_{\text{cell}}\) is 0.5. According to Hooke’s law \(F=kt\), where \(k\) is the spring constant of the cantilever), the loading force \(F\) can be calculated from the cantilever deflection \(x\). After obtaining the loading force \(F\) and the indentation depth \(\delta\), the cellular Young’s modulus \(E_{\text{cell}}\) can be calculated according to formulas (6) and (7). It should be noted that the Hertz-Sneddon model is based on a number of assumptions applied to the specimen to be indented, including homogeneity, isotropy, linear elastic material properties, axisymmetry and infinitesimal deformation of the sample, as well as infinite sample thickness and a smooth sample surface. Studies have shown that the Hertz-Sneddon model is appropriate as long as the indentation into the specimen is no more than 10% of the specimen thickness. However, the indentations should be larger than 400 nm to avoid errors due to the uncertain determination of the contact point, which leads to measurements often being performed on the central region of the cell. The Hertz-Sneddon model does not consider the forces between the contact surfaces (e.g., electrostatic forces, adhesion forces, and friction forces) and is therefore applicable when the forces between the contact surfaces are negligible compared with the maximum load. By contrast, the Johnson-Kendall-Roberts (JKR) model and Derjaguin-Muller-Toporov (DMT) model take into account the adhesion forces between the contact surfaces. The JKR model considers the adhesion forces inside the contact area whereas the DMT model considers the forces outside the contact area. Researchers have shown that the JKR model can be applied in the case of large tips and soft samples with a large adhesion, whereas the DMT model is applicable in the case of small tips and stiff samples with a small adhesion. The conventional cone tip is sharp with a small opening angle of the tip, and the DMT model is applicable in the case of large tips and stiff samples with a large adhesion. The JKR model is applicable in the case of large tips and soft samples with a large adhesion. The DMT model is applicable in the case of small tips and stiff samples with a small adhesion.
stimulation of drugs\cite{51,52,53} to assess the drug actions from the viewpoint of cell mechanics.

By linking drug molecules to the tip, AFM can probe the target molecules on the cell surface. This technique is called single-molecule force spectroscopy (SMFS)\cite{54}, as shown in Figure 1C. By performing approach-retract cycles on the cell surface using drug-tethered tips, force curves can be recorded. If a drug molecule binds to a target molecule during the contact between the tip and cell, the drug-target complex is then pulled by the AFM tip during the retract process, leading to a non-linear unbinding peak on the retract curve (indicated by the green arrow in Figure 1C). The non-linear shape of the peak is caused by the stretching of flexible linker molecules, which have non-linear elasticity\cite{55}. The magnitude of the peak is equal to the binding force between a target molecule and a drug molecule (Figure 1C). There are several methods for linking molecules onto AFM tips\cite{55,56}, including physisorption, covalent coupling with organosilanes or ethanamine hydrochloride and the functionalization of gold-coated tips. Using indolent polyethylene glycol (PEG) molecules to tether drug molecules onto the AFM tip is the most common strategy due to its many advantages, such as allowing the drug molecules to freely reorient with respect to the target molecules on the cell surface, preventing the danger of drug molecules being compressed when the tip contacts the cell surface, and enabling the clear identification of the specific binding according to the unbinding peak on the retract curve\cite{51}. Traditional methods for characterizing the binding constant of drug molecules cannot reveal the dynamics taking place during the dissociation of a drug-target complex. However, because SMFS can reveal the dynamics taking place during the process of pulling drug molecules from target cells, it enables better understanding of the drugs and is useful for drug design. We can thus conclude that AFM is a powerful method that can be used to obtain multiple physicochemical parameters, such as morphology, elasticity, and drug-target interactions, on single cells and native proteins under the stimulation of drugs. In the following sections, we will summarize the recent progress of AFM as applied to the monitoring of drug actions on membranes.

**Dynamic changes of cellular ultra-microstructure and elasticity in response to drugs**

The morphology of a cell is closely related to its functions. When a cell becomes pathological, its morphology changes. For example, the shape of normal red blood cells is an oval biconcave disk, whereas those of anemia patients become sickle-shaped\cite{57}. Additionally, the morphology of a normal cell changes when it transforms into a cancerous one\cite{58}. For example, the nuclear volume of cancer cells is often larger than those of normal cells and the nuclear contour of cancer cells is often irregular\cite{59}. There is also difference in the cell surface brush between normal cells and cancer cells\cite{60}. In fact, because of the significant morphological differences between normal cells and cancer cells, the primary method for diagnosing cancers in the clinic is based on the observation of stained pathological tissue samples\cite{50}. Due to its 200 nm limitation in resolution, the traditional observation of cellular morphology using optical microscopy cannot reveal cellular nanostructures. Thus, the detailed morphological changes of cells after the stimulation of drugs cannot be explored with optical microscopy.

AFM opens the door to *in situ* investigations of the nanoscale morphological changes in single cells after drug stimulation. Antimicrobial peptides are a promising class of antimicrobials that have demonstrated activity against antibiotic-resistant bacteria, parasites, viruses and fungi\cite{14}. In 2010, Fantner et al\cite{14} used AFM to image the dynamic nanoscale changes on single living bacterial cells in response to CM15 (an antimicrobial peptide drug), revealing that the cell surface changed from smooth to corrugated after the stimulation of CM15. Glyphosate is an extensively used herbicide that has been shown to increase the risk of cancer and quercetin is a well-known antioxidant\cite{61}. In 2012, Heu et al\cite{62} investigated the effect of glyphosate and quercetin on single living keratinocyte cells by AFM peak force tapping mode, showing that cells develop a filamentous cytoskeleton network after the stimulation of glyphosate, whereas the filamentous structures disappeared after the subsequent addition of quercetin. Daunorubicin (Dau) is a drug used to treat several types of cancers, such as leukemia and neuroblastomas\cite{63}. Dau intercalates in DNA to cause a local unwinding of the DNA helix. In 2013, Alonso-Sarduy et al\cite{64} applied time-lapse AFM to investigate the dynamic changes of DNA molecules in response to Dau in a physiological medium. The results clearly showed conformational changes in the DNA molecules after the addition of Dau. These studies have provided novel insights that are useful for understanding the drug actions and to assess drugs from the viewpoint of cellular nanostructures (e.g., cell surface, cytoskeleton, DNA).

Studies in the past decade have demonstrated that cell mechanics play an important role in maintaining cellular physiological activities\cite{53,63}. Cells sense external mechanical stimuli and convert them into intracellular chemical signals via proteins on the cell surface to regulate cellular physiological activities\cite{65}. For example, in the process of cancer metastasis, circulating tumor cells (CTCs) must undergo large elastic deformations to penetrate endothelial cell-cell junctions as well as overcome the effect of fluid shear to adhere to the vascular endothelium of distant organs\cite{65}. Researchers have shown that pathological changes can cause changes in cellular mechanical properties (such as changes in cell deformability and altered cytoadherence characteristics), which then result in changes in cellular physiological properties (such as protein expression and cell functions) and finally lead to the occurrence of cancer\cite{52}. In 2012, Plodinec et al\cite{66} comprehensively investigated the stiffness of three types of biopsy samples (normal breast tissues, benign breast tissues and malignant breast tissues), and the results showed that the stiffness profiles of breast tissues changed remarkably during the development of breast cancer, providing quantitative indicators in the clinical diagnostics of cancer with translational significance.
In recent years, there has been an increasing consensus that cell mechanics can serve as an effective label-free biomarker for the indication of cell states\cite{50, 66}.

Investigating drug-cell actions from the viewpoint of cell mechanics provides a novel way to evaluate drugs. The cytoskeleton is crucial for maintaining the cell shape and therefore some drugs, such as cytochalasin and taxol, influence the cytoskeleton to inhibit cell migration. In 2000, Rotsch et al\cite{70} investigated the effect of various drugs (cytochalasin, taxol, colchicine, and latrunculin) on fibroblasts by measuring the cell elasticity before and after the stimulation of drugs. The results showed that drugs (cytochalasin, latrunculin) that dis-aggregated the actin filaments could cause a distinct decrease in cell elastic modulus whereas drugs (taxol, colchicine) that stabilized microtubules could not significantly influence the cell elasticity. In 2007, Lam et al\cite{51} investigated the mechanical properties of leukemia cells (isolated from the blood of patients with diagnosed acute leukemia) in response to chemotherapy drugs (dexamethasone, daunorubicin), and the results showed that leukemia cell stiffness increased by nearly two orders of magnitude after exposure to dexamethasone or daunorubicin. N-methyl-D-aspartate (NMDA) specifically binds to the NMDA receptors in the cytomembrane that opens the ligand gated ion channel to facilitate Ca\(^{2+}\) influx into the cell\cite{71}. In 2014, Fang et al\cite{50} investigated the effect of NMDA on SH-SY5Y human neuroblastoma cells from the viewpoint of cell elasticity, and the results showed that the cell Young’s modulus significantly increased after the treatment of NMDA. These studies provide novel insight into the effects of drugs and also facilitate the development of new therapeutic strategies for combating diseases such as cancer. One important issue regarding measuring the elasticity of living cells using AFM is whether the consecutive probing by the tip itself causes the observed changes in cellular elasticity. Researchers have shown that prolonged poking (200 s) of the cell can lead to the remodeling of the cell cytoskeleton, and thus the probing time should be limited during the measurements to avoid this effect\cite{51}. In practice, when we investigate the effect of external stimuli such as drugs on the cellular elasticity, control experiments (e.g., monitoring changes in the cellular elasticity without the addition of drugs)\cite{53, 51} are often required to determine the influence of tip poking on the cellular elasticity.

Rituximab is a monoclonal antibody that targets the CD20 antigen on the surface of B-cell non-Hodgkin lymphoma cells. One of the primary functions of rituximab is that it can activate the classical complement pathway to lyse the cancer cells; this effect is called complement-mediated cytotoxicity (CMC). We have investigated the dynamic changes of cellular ultrastructures and their mechanical properties during the CMC process\cite{74}, as shown in Figure 2. A schematic diagram of the CMC process is illustrated in Figure 2A. The binding of rituximab to CD20 on a lymphoma cancer cell can activate the C1 complex, which then triggers the classical pathway by the cascade interactions of complement proteins (such as C2, C4, C3). The terminal product of CMC is the membrane attack complex (MAC) formed on the cell surface. Small molecules (e.g., water, ion) can enter into the cells freely, which causes them to swell and burst. Traditional methods for investigating the CMC effect are based on optical microscopy, which cannot resolve the detailed events taking place on the cell surface during CMC. We therefore used AFM to image the nanoscale morphology of lymphoma cells that exhibited the CMC effect, as shown in Figure 2B–2J. In the early stage of CMC, some micro-holes appeared on the cell surface (denoted by the arrows in Figure 2C), which were related to the formation of MAC on the cell surface due to the CMC effect. In the middle stage, we observed that the size of the holes increased (denoted by the arrows in Figure 2F) and that cytoplasm had escaped from the cell (denoted by the arrow in Figure 2E). In the final stage, the cells collapsed, leaving behind only cell membrane debris (Figure 2H–2J). Collectively, the AFM images clearly showed the dynamic processes of CMC from the nanoscale cellular topography. Additionally, changes in the cellular mechanical properties during CMC were also measured. The cells were PI-stained to identify the CMC effect (Figure 2K). Under the guidance of fluorescence, the AFM tip was moved toward the cells exhibiting the CMC effect (Figure 2I). The Young’s modulus of lymphoma Raji cells was measured at three intervals (before treatment, after treatment without CMC, and after treatment with CMC). The results (Figure 2M) revealed that during CMC the cellular Young’s modulus first decreased and then significantly increased, improving our understanding of rituximab’s CMC effect. The studies also indicated that the efficacy of drugs could be assessed by monitoring the mechanical properties of the cells. Both experimental\cite{50, 66} and theoretical\cite{71} studies have shown that cell mechanical properties are closely related to the uncontrolled division and migration of tumor cells; therefore, we can consider designing drugs that alter the mechanical properties of tumor cells. These drugs would likely not kill cancer cells, but may effectively hinder the propagation of the neoplasm\cite{74}.

**In situ quantification of individual drug-target interactions**

Drug-target interactions (such as the binding force) are closely related to the overall drug efficacy; therefore, investigating drug-target interactions is important for better understanding the drug action. Traditional methods for characterizing drug-target interactions require many purified target molecules that are isolated from cells. Studies have shown that the cell membrane plays an essential role in determining the functions of the membrane proteins\cite{74}. Therefore, the results obtained from purified proteins cannot faithfully reflect the real properties of the same molecules in vivo\cite{74}. The pioneering work of Gaub et al showed that AFM could be used to measure the binding force of individual receptor-ligand pairs\cite{74} and to investigate the unfolding dynamics of single molecules\cite{74}. Following his work, similar studies on different types of purified molecules emerged\cite{74, 78}, providing a better understanding of the molecular interactions. In 2000, Benoit et al\cite{74} investigated cell-cell adhesion interactions by AFM force spectroscopy, demonstrating the potential of AFM to measure
individual molecular interactions in situ. With AFM force spectroscopy, the binding force of individual drug-target pairs can be directly quantified by linking drug molecules to the AFM tip. Compared with traditional methods, AFM provides a novel approach to investigate the drug-target interactions and provides unique data that are useful for the improvement

Figure 2. Dynamic changes of cellular nanostructures and mechanical properties during CMC investigated by AFM. (A) Schematic diagram of CMC. (B–J) AFM images of lymphoma Raji cells after the treatment of Rituximab and human serum. (K) Fluorescence image of a PI-stained Raji cell. (L) Under the guidance of fluorescence, AFM tip was moved to the cell. (M) Dynamic changes of cellular Young's modulus during CMC. (B–M) were reprinted with permission from Ref. Copyright 2014 IEEE.
of drug design. Sodium-glucose co-transporter (SGLT) plays an important role in intestinal glucose adsorption and in renal glucose reabsorption in many organisms. SGLT is the target for a series of drugs used in the treatment of diabetes, such as canagliflozin\textsuperscript{[80]} and ipragliflozin\textsuperscript{[81]}. In 2006, Puntheeranurak \textit{et al}\textsuperscript{[82]} applied AFM force spectroscopy to measure the binding force of SGLT on living CHO cells by linking its antibody to an AFM tip. The results demonstrated the potential of AFM to study the presence and dynamics of plasma membrane transporters in intact cells at the single-molecule level. Vascular endothelial growth factor receptors (VEGFRs) are important targets for cancer treatment because of their crucial roles in vascular development, angiogenesis and lymphangiogenesis\textsuperscript{[83]}. In 2007, Lee \textit{et al}\textsuperscript{[84]} combined AFM imaging and molecular recognition to investigate the location and binding kinetics of VEGFRs on human umbilical vein endothelial cells (HUVEC) by linking monoclonal anti-VEGFR2 antibody to an AFM tip. The results demonstrated that AFM can provide spatiotemporal visualization of cell surface dynamics that regulate receptor-mediated behaviors. So far, researchers have applied AFM force spectroscopy to investigate a wide range of drug-target interactions on a series of cells, such as Herceptin-EGFR on HEK 293 cells\textsuperscript{[85]}, quantum dot-HeLa cell interactions\textsuperscript{[86]}, hormone-drug on HeLa cells\textsuperscript{[87]}, transporter-antibody on erythrocyte\textsuperscript{[88]}, and aptamer-glycoprotein on U251 cells\textsuperscript{[89]}. Although these studies improve our understanding of drug-cell interactions at the single-molecule level, they were performed on cell lines cultured \textit{in vitro}. In fact, the conditions used in research laboratories (eg, tumor cells are often selected based on their ability to grow rapidly \textit{in vitro} and occasionally their relative sensitivity to therapy) vary significantly from those used in the real-world clinical environment\textsuperscript{[90]}. Therefore, the results obtained using cell lines may not accurately reflect realistic drug-cell interactions.

To better understand drug-cell interactions, direct investigations of pathological cells from clinical patients are required. Experiments performed on patient cells can provide novel insights with translational medical significance. To this end, we have investigated drug-target interactions directly on cells from clinical lymphoma patients\textsuperscript{[91-93]}, as shown in Figure 3. The binding of rituximab to the CD20s on lymphoma cancer cells can lead to cell lysis via three mechanisms, including direct induced apoptosis, antibody-dependent cellular cytotoxicity (ADCC) and CMC (Figure 3A). To investigate the CD20-rituximab interactions directly on cancer cells from lymphoma patients, the cancer cells from the clinical biopsy samples should first be identified because in biopsy samples (such as bone marrow), cancer cells and normal cells are mixed together. Receptor tyrosine-like orphan receptor 1 (ROR1) is a specific cell surface marker that is highly expressed on B lymphoma cancer cells, but not on normal cells\textsuperscript{[94, 95]}. Therefore, we used ROR1 to identify the cancer cells in the bone marrow samples prepared from B-cell lymphoma patients. Then, the CD20s on the cancer cells could be probed using rituximab-tethered tips (Figure 3A). Under the guidance of ROR1 fluorescence recognition, the AFM tip was positioned on the cancer cell (denoted by the black arrow in Figure 3B). The obtained typical force curves contained a specific unbinding peak in the retract curve (denoted by the green arrow in Figure 3C), which was caused by the stretching of PEG spacer molecules. By obtaining thousands of force curves at different points on the cell surface, a histogram of the binding force was constructed (Figure 3D). Additionally, by obtaining 16×16 force curves on the local cell surface (500 nm×500 nm), gray maps that reflected the nanoscale distribution of CD20s were constructed (Figure 3E). For contrast, few gray pixels in the gray maps were obtained on normal cells (red blood cells) (Figure 3F) that did not express CD20s. To investigate the role of CD20-rituximab interactions in the rituximab clinical therapy, biopsy samples from three lymphoma patients were tested. The results showed that for the three patients, there was no significant difference in the binding force of CD20 on cancer cells (Figure 3G); however, there was significant difference in the distribution density of CD20 on cancer cells (Figure 3H). From the results obtained from combining AFM together with the clinical treatment information (Figure 3I), we were able to conclude that the distribution density of CD20 on the lymphoma cancer cells had a direct impact on the clinical efficacies of rituximab. Current biochemical research is typically performed on cells grown \textit{in vitro}, which leads to a large gap between the laboratory results and clinical requirements\textsuperscript{[54]}. The results shown in Figure 3 represent a new method to deepen the relationship between clinical treatment and laboratory research and also demonstrate the potential of using AFM force spectroscopy to evaluate drug efficacies from the perspective of clinical medicine.

**High-resolution imaging and manipulation of individual native target molecules**

Determining the structure of target molecules is crucial for understanding their interactions with drug molecules\textsuperscript{[28]}. Most of the targets are membrane proteins. The amphiphilic nature of membrane proteins makes the application of traditional structural biology methods (eg, nuclear magnetic resonance spectroscopy, electron microscopy and X-ray crystallography) toward their study challenging, especially with regards to protein expression, protein purification, and crystallization\textsuperscript{[86]}. AFM provides an alternative way to investigate the nanoscale morphology and mechanical properties of individual native membrane proteins. By reconstituting purified membrane proteins into a lipid bilayer to mimic their native environment, AFM can observe single native membrane proteins at work, which can thus provide different and complementary information to that obtained from traditional structural biology methods\textsuperscript{[28]}. Gap junction channels mediate communication between adjacent cells\textsuperscript{[97]}. In 2002, Muller \textit{et al}\textsuperscript{[87]} imaged the conformational changes of single native junction channels in response to ions in buffer solution by adsorbing the purified gap junction fractions to mica. The results showed that calcium ions could induce closing of the channels, as shown in Figure 4A and 4B. Potassium channels are tetrameric membrane proteins that facilitate the permeation of potas-

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Figure 3. Investigating the CD20-rituximab interactions on cancer cells in the biopsy samples prepared from clinical lymphoma patients by AFM force spectroscopy\cite{91-93}. (A) Schematic diagram of rituximab’s mechanisms and probing the CD20-rituximab interactions on lymphoma cancer cells. (B) Under the guidance of ROR1 fluorescence recognition, AFM tip carrying rituximabs was moved to cancer cell. (C) Typical force curve obtained on cancer cell with specific unbinding peak. (D) Histogram of binding force. CD20 distribution map obtained on cancer cell (E) and normal cell (F). (G) Histogram of binding force and (H) distribution density of CD20s on the cancer cells from three lymphoma patients. (I) Clinical treatment information of the tree patients. (B) was reprinted with permission from Ref\cite{91}. Copyright 2013 John Wiley & Sons, Ltd. (C–F) were reprinted with permission from Ref\cite{92}. Copyright 2013 Elsevier Inc. (G–I) were reprinted with permission from Ref\cite{93}. Copyright 2014 Royal Microscopical Society.
sium ions through the membrane with high specificity and high-throughput rates\cite{16}. In 2011, Mari et al\cite{16} visualized the conformational changes of a single MloK1 potassium channel mediated by ligands. The AFM images clearly showed that the binding of cAMP molecules could lead to the opening of the potassium channels, whereas the potassium channels closed in the absence of cAMP, as shown in Figure 4C and 4D. It has been established that traditional structural biology methods such as X-ray crystallography and electron microscopy can only provide the static structure of a membrane protein\cite{96, 98}. By contrast, AFM can image the dynamic conformational changes of single native membrane proteins in response to external stimuli such as drugs, providing direct evidence of the drug actions from the \textit{in situ} single molecules.

**Figure 4.** Imaging and manipulating single native membrane proteins by high-resolution AFM and force spectroscopy\cite{16, 97, 99}. (A, B) Imaging Ca$^{2+}$-induced conformation change of the extracellular connexon surface of gap junction channels. (A) In Ca$^{2+}$-free buffer solution. (B) In the presence of 0.5 mmol/L CaCl$_2$. The insets are the average of the raw data. (C, D) Visualizing ligand-induced conformation change of the potassium channel. (C) In the presence of cAMP. (D) In the absence of cAMP. The insets are the average (top) and SD map (bottom) of raw data. (E, F) Single-molecule force spectroscopy of membrane protein NhaA. (E) Schematic diagram. (F) Force curve revealing the unfolding of a single NhaA molecule. (A, B) were reprinted with permission from Ref\cite{97}. Copyright 2002 European Molecular Biology Organization. (C, D) were reprinted with permission from Ref\cite{16}. Copyright 2011 National Academy of Sciences. (E, F) were reprinted with permission from Ref\cite{99}. Copyright 2006 Elsevier Ltd.
In addition to imaging the two-dimensional morphology, AFM can also reveal the unfolding dynamics of a single membrane protein by pulling one of its ends, as shown in Figure 4E. In 2006, Kedrov et al. investigated the unfolding dynamics of individual antiporter NhaAs that were reconstituted into the lipid bilayers by obtaining force curves on the protein. The obtained force curves shown in Figure 4F clearly revealed serial serrated peaks, each of which corresponded to the unfolding of one domain of the membrane protein. By applying AFM force spectroscopy, the unfolding dynamics of various types of membrane proteins have been investigated. In a recent study, high-speed force spectroscopy was developed to unfold single proteins at velocities previously only reached by simulations. This finding has enabled direct comparison of experimental results to those of simulations, which is of considerable significance in studies of molecular mechanical dynamics. These studies have demonstrated the ability of AFM to investigate the dynamic morphology and unfolding process of single native target molecules, which is of potential significance for pharmacology and drug development. AFM can be used to directly visualize drug effects on single target molecules as well as to obtain the unfolding dynamics of a single target molecule. Current molecular simulations are based on 3-D structural data obtained by traditional structural biology methods. The information obtained by AFM will provide complementary data, which may in turn improve the computer simulations of drug-target interactions.

**Challenge and outlook**

AFM provides a novel powerful instrument for researchers to investigate drug-target interactions at the single-cell and single-molecule levels under near-physiological conditions. It allows us to visually and physically probe drug effects at the nanoscale, which are inaccessible by traditional biochemical methods. Measuring the physical properties of active biological samples (such as living cells and native membrane proteins) probed by AFM is advantageous in that it does not require costly labeling or sample preparation. Single-cell assays based on AFM offer new opportunities for analyzing the heterogeneity of individual cells, which is masked by traditional ensemble experiments. The typical applications summarized here of AFM in the past decade (topography imaging, elasticity measurements, molecular interaction quantification, native membrane protein imaging and manipulation) for monitoring the physiological activities on single cells and single molecules after the stimulation of drugs have provided considerable novel insight into our understanding of drug actions, demonstrating the valuable role of AFM for pharmacology and drug development. However, there are still several challenges that need to be addressed to improve the current AFM-based research.

The first challenge is how to improve the spatial resolution and reliability of AFM imaging and force spectroscopy probing of living cells. Although AFM imaging and force spectroscopy can directly image and reveal the unfolding dynamics of single native membrane proteins, the current research has been performed on reconstituted lipid bilayers. Clearly, the mechanistic results derived from an *in vitro* simulacrum must be addressed at the cellular level. AFM has achieved great success in studies of microbial cells (such as bacteria and fungi) because of the well-defined and rigid nature of the microbial cell envelope. However, AFM imaging of mammalian cells remains limited to resolutions in the 50–100 nm range due to the soft, dynamic and highly heterogeneous nature of the cell surface, meaning that single proteins on living cells cannot be observed. For mammalian suspended cells in particular, the prerequisite of performing AFM experiments is to immobilize them onto the substrate. Current immobilization methods such as the micro-well array and micro-pillar array methods are effective for trapping the whole cell. However, they cannot eliminate the dynamic movement of the cell membrane, which prevents the acquisition of higher resolution images of local areas on living mammalian cells. Methods that can inhibit the dynamics of the cell surface are still lacking. Moreover, the vast majority of interactions detected during AFM force spectroscopy experiments are unspecific because of the high complexity of the cell surface (e.g., a cell membrane contains hundreds of lipids and a plethora of proteins, and also exhibits raft-based heterogeneity). When performing force spectroscopy on living cells, the AFM probe can be contaminated after recording only a few force curves. Therefore, current AFM methods (such as simultaneous topography and recognition imaging) are mainly adapted for chemically fixed cells, which allow us to obtain data with a higher signal-noise ratio. Studies that convincingly demonstrate the reliability of force spectroscopy experiments on living mammalian cells are still needed.

The second challenge stems from the low throughput of AFM experiments and the intensive labor required to perform them. Currently, AFM experiments are not automatic and must be performed manually. Under the guidance of optical microscopy (or fluorescence), an operator moves the AFM tip to the cell of interest to perform measurements (such as morphology imaging and force spectroscopy). After the measurements are finished, the AFM tip is directed toward another cell. This process leads to a very low throughput (the time of handling one cell is often several minutes). To enable practical applications such as drug screening and evaluation, the throughput of AFM should be considerably improved and thus automation strategies may be required. Current AFM measurement experiments (such as elasticity measurements and molecular interactions measurements) also require the off-line processing of force curves, which is time-consuming (typically thousands of force curves need to be analyzed). Recently, the advent of a novel AFM mode called peak force tapping mode has enabled analysis of the obtained force curves in real-time and directly provides the multiparametric images (such as the Young's modulus and adhesion force). This development has greatly improved the efficiency of AFM force measurements. We have presented a strategy for automatically performing AFM measurements by cell recognition via an image processing method, and the results demon-
strated that it could significantly reduce the measurement time. However, there are still several issues that need to be addressed to achieve fully automated AFM measurements, such as the integration of cell delivery techniques, automation of the system calibration and visualization of the detection results. Regarding cell delivery, microfluidic techniques\(^ {115}\) are of potential significance, whereas for data visualization, the 2-D scatter plots and gating approaches widely used in flow cytometry\(^ {467}\) will be useful. If fully automated AFM measurements, such as automatic monitoring of the changes in cellular elastic properties, can be achieved, then they can be applied to drug research. For example, we can imagine employing AFM to screen out drugs that can bind to cytoskeleton proteins and change the elastic properties of cells, which is of significance in promoting the development of the pharmaceutical industry and clinical disease therapy.

The third challenge is to improve the time resolution of AFM measurements. Currently, the time needed to acquire a high-resolution image (often several minutes) is much greater than the time scale at which dynamic processes usually occur in biology\(^ {22}\) (e.g., bacteria respond to antibiotics on the time scale of seconds\(^ {116}\)). Although high-speed scanning AFM has reduced the acquisition time for an image to less than one second\(^ {22,41}\), it is primarily limited to certain samples (such as rigid microbial cells\(^ {116}\) and isolated molecules adsorbed on substrate\(^ {41}\) and the scan size is often small (less than one micrometer). To make high-speed AFM applicable to living mammalian cells, which are large and have rough surfaces, several technical and conceptual obstacles need to be overcome\(^ {117}\). Moreover, so far AFM cannot access the interior of cells. A recent study has combined AFM with confocal fluorescence microscopy to measure in situ the elasticity of the cell nucleus directly on living cells via a needle tip (the tip can penetrate the cell membrane\(^ {118}\), demonstrating the potential of AFM for intracellular mechanics studies. However, this penetration method is in fact invasive; therefore, the influence of the manipulation process on the biological activity of cells needs to be clarified (e.g., whether the cell is still alive and whether it can be used for subsequent experiments). In addition, some applications (e.g., force spectroscopy for probing molecular interactions or cell-cell adhesions) require particular modifications of the AFM tip, such as linking specific molecules (e.g., antibodies, ligands, viruses) or a living cell to the AFM tip\(^ {119}\). However, the currently available protocols are labor-intensive, time-consuming and complex, and the lifetime of the functionalized tip is often short. Therefore, developing simple and standardized protocols for tip functionalization\(^ {22}\) will certainly contribute to advances in AFM force spectroscopy experiments.

In summary, the results achieved by applying AFM to the monitoring of drug actions on membranes at the nanoscale in recent decades have provided new opportunities for drug development. However, there are still challenges, as described above, which are hampering the further development of AFM. Overcoming these challenges will require the efforts of researchers from different disciplines (e.g., biology, engineering, physics, automation). Nonetheless, it is gratifying to see that in recent years important progress has been made to enhance the functions of AFM, such as high-speed imaging (to improve the scan speed)\(^ {41}\), multiparametric mode (to simultaneously obtain multiple parameters)\(^ {22}\), high-speed force spectroscopy (to improve the speed of obtaining force curves)\(^ {120}\), nanofluidics AFM (to allow the injection of chemical molecules or drugs into a single cell)\(^ {125}\), tomographic contact resonance AFM (to generate cross-sectional stiff images of samples)\(^ {121}\) and fast nanomechanical spectroscopy (to improve the speed of mapping the mechanical properties)\(^ {122}\). Moreover, AFM can also be combined with other techniques to obtain complementary information, and this will be an important aspect for future AFM studies. For example, the spatial resolution of optical microscopy can now reach the nanometer level via fluorescence\(^ {122}\). Under the guidance of ultra-resolution optical (fluorescence) microscopy, AFM may be used to probe individual structures such as cellular machinery on the surface of a living cell (e.g., to measure their mechanical properties). Patch-clamp is a commonly used technique in studies of ion channels. AFM has been successfully combined with patch-clamp to simultaneously obtain the mechanical dynamics and current responses of the ion channels on a living cell\(^ {124}\), providing novel insights into the mechanosensitive ion channels. Scanning ion conductance microscopy has been shown to be effective in imaging the surface of living cells, as it has a noncontact character\(^ {125}\). This technique has inspired us to improve the spatial resolution of imaging living cells by developing noncontact AFM scanning mode in liquids. Collectively, the exciting results obtained with AFM indicate that its further applications in the life sciences will bring about yet more changes in related fields such as pharmaceutical industry and clinical medicine.

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