FluidFM for single-cell biophysics

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ABSTRACT

Fluidic force microscopy (FluidFM), which combines atomic force microscopy (AFM) with microchanneled cantilevers connected to a pressure controller, is a technique allowing the realization of force-sensitive nanopipette under aqueous conditions. FluidFM has unique advantages in simultaneous three-dimensional manipulations and mechanical measurements of biological specimens at the micro-/nanoscale. Over the past decade, FluidFM has shown its potential in biophysical assays particularly in the investigations at single-cell level, offering novel possibilities for discovering the underlying mechanisms guiding life activities. Here, we review the utilization of FluidFM to address biomechanical and biophysical issues in the life sciences. Firstly, the fundamentals of FluidFM are represented. Subsequently, the applications of FluidFM for biophysics at single-cell level are surveyed from several facets, including single-cell manipulations, single-cell force spectroscopy, and single-cell electrophysiology. Finally, the challenges and perspectives for future progressions are provided.

KEYWORDS

atomic force microscopy, fluidic force microscopy, single-cell manipulation, single-cell force spectroscopy, single-cell electrophysiology

1 Introduction

It is increasingly evident that investigating the physiological/pathological processes at single-cell level is of remarkable significance for unveiling the mysteries of life activities. Traditional biological assays are based on ensemble measurements performed on cell populations, which are undoubtedly powerful tools for the identification of components and interactions within complex metabolic, signaling, and transcriptional networks [1]. Nevertheless, it should be noted that cell-to-cell variability is not apparent in population-averaged assays [2]. It is widely known that single cells in a population display variable behaviors [3] and even genetically identical cells grown in homogeneous environments exhibit different phenotypes [4]. The stochasticity in molecular processes such as stochastic gene expression [5] has been found to be related to the cell-to-cell phenotypic variability. Particularly, for tumors, intratumor (within a tumor, subclonal diversity exists) and intercellular (within a subclonal population of tumor cells, tumor cells exhibit different behaviors) heterogeneity have been widely observed [6, 7], which are key factors that may lead to drug resistance and pose challenges for the design of clinical trials [8]. Hence, investigating life activities at single-cell level significantly benefits the reveal of the novel underpinnings regulating physiological/pathological processes, which has been inaccessible for a long time due to the lack of study tools. In the past years, various single-cell biochemical methods have emerged to profile genetic, epigenetic, spatial, proteomic and lineage information in individual cells [9], including single-cell ribonucleic acid (RNA) sequencing [10], single-cell deoxyribonucleic acid (DNA) methylation profiling [11], single-cell western blotting [12], and so on. With the use of these single-cell biochemical assays, unforeseen processes and patterns in life sciences have been revealed [13], such as intratumoral heterogeneity in primary glioblastoma [14], the random monoallelic gene expression revealed [13], such as intratumoral heterogeneity in primary glioblastoma [14], the random monoallelic gene expression in mammalian cells [15], and the identification of somatic mutations in single cell [16].

Investigating single-cell mechanics has emerged as a promising way for understanding the biological processes. Life is not only a chemical process, but also a mechanical one [17]. In fact, mechanical forces are increasingly recognized as major regulators of cell structure and function. The mechanical properties of cells are essential to the mechanisms by which cells sense external mechanical cues and transduce them into chemical signals for regulating various biological processes at the molecular and cellular level [18], such as gene expression, adhesion, migration, and cell fate [19]. Abnormal alterations in cell mechanics often influence, and are influenced by, the onset and progression of human diseases such as cancer [20]. For example, it has been found that in addition to genetic and external environmental factors, the physical interactions of cancer cells with their microenvironment, as well as their modulation by mechanical forces, are key determinants of the metastatic process of tumor [21]. Therefore, studying cell mechanics is meaningful for identifying the physical biomarkers in physiological/pathological processes [22], which significantly complements traditional biochemical assays for thoroughly understanding life activities. Particularly, single-cell biochemical assays require various pretreatments of cells, such as lysis, fixation, staining, and labeling [14–16, 23], meaning that cells
lose biological activities after these treatments. On the contrary, single-cell mechanical measurements are directly performed on living cells and the biological functions (such as invasion and metastatic capabilities [24]) of the same cells can further be examined by using biochemical assays, which facilitates investigating the direct association between cell mechanics and cell functions at the single-cell level and therefore is of fundamental significance for revealing the heterogeneous behaviors of single cells.

The advent of fluidic force microscopy (FluidFM) [25] provides a powerful multifunctional toolbox for the studies of mechanics at the single-cell level. FluidFM is derived from atomic force microscopy (AFM) [26]. AFM uses a sharp tip mounted at the end of a microcantilever to probe the specimens immobilized on the support, and the unique advantage of AFM is that it is able to work under aqueous conditions with nanometer spatial resolution and millisecond temporal resolution [27], making it particularly suited for the studies of living biological specimens in their native state. Besides resolving the topography at diverse imaging modes [28], AFM has achieved unprecedented success in detecting the mechanical properties of biological specimens in force spectroscopy mode [29], significantly contributing to the field of mechanobiology. In addition to AFM, various emerging techniques are available for measuring the mechanics of single cells, including microfluidics, micropipette aspiration, micropost arrays, magnetic twisting cytometry (MTC), optical stretching, parallel-plate rheometry, particle-tracking microrheology (PTM), and cell monolayer rheology (CMR) [30–34], each of them with its advantages and limitations [30–34]. In practice, AFM is the most widely used tool for single-cell mechanical analysis [35–38]. However, regular AFM probe can only perform simple mechanical motions on cells in the horizontal plane, such as touching, compressing, indenting, pushing, stretching and cutting [39]. By mounting a nanoneedle [40] or nanotube [41] probe onto the AFM head, molecules could be delivered into single cells, but such protocol required the covalent conjugation of target molecules onto the surface of AFM probe and was dependent on reversible chemical reactions, which decreased the fidelity of target molecules and caused limited controllability. Inspired by the glass micropipette [42] which is a widely used tool for cell manipulation, FluidFM takes advantage of microchanneled AFM cantilevers connected to a pressure controller [43]. In this way, FluidFM integrates the advantages of micropipette and AFM, since FluidFM can not only perform three-dimensional manipulations of single cells, but also tune liquid delivery to single cells with high precision (femtoliter) and perform mechanical measurements on single cells with nanometer spatial resolution, providing new possibilities for the studies at single-cell level. Recently, Saha et al. [44] have surveyed the applications of FluidFM on force measurements. Here, the advances of FluidFM in single-cell biophysics (including single-cell manipulations, single-cell force spectroscopy (SCFS), and single-cell electrophysiology) are comprehensively summarized and illustrated with examples highlighting the unique capabilities of FluidFM. The challenges and future progressions are also discussed.

2 Fundamentals of FluidFM

The principle of FluidFM is shown in Fig. 1. FluidFM is based on AFM. An AFM is commonly composed of a piezoelectric tube driver which is connected to a probe, a laser generator, a four-quadrant position sensitive detector (PSD), a signal processing module and the feedback control electronics [45]. AFM probe raster scans the surface of the specimen, and a laser beam is reflected from the backside of the cantilever to the PSD to detect the cantilever deflection. During X-Y scanning, changes in PSD signals are analyzed by the signal processing and input into the feedback module which controls the Z (vertical) movement of the piezoelectric tube driver in order to maintain the feedback parameters constant (e.g., cantilever deflection and amplitude of the vibrating cantilever), thus yielding a three-dimensional topography of the specimen. The basic components of FluidFM are the same as AFM, except that FluidFM uses a microchanneled cantilever which is connected to a pressure controller by tubing, as shown in Fig. 1(a). FluidFM thus has a continuous and closed fluidic channel that can be filled with an arbitrary chosen liquid, which can be locally dispensed through the nanoscale aperture at the extremity of the cantilever [25]. The design of FluidFM gives AFM exciting novel capabilities; for example, it allows AFM to be able to perform precise three-dimensional manipulations on individual living cells (e.g.,

Figure 1 FluidFM configuration. (a) Schematic of FluidFM. The microchanneled cantilever is fixed to the drilled AFM probe holder, which is immersed in liquid on top of an inverted optical/fluorescent microscope. (Reprinted with permission from Ref. [25], © American Chemical Society 2009). (b)–(j) SEM images of the different types of microchanneled probes. (b) The whole probe. (c) Cutaway view of the pyramidal tip with embedded microchannel. (d) Hollow pyramid after sectioning the probe with FIB. (e) and (f) Aperture at the apex (e)/side (f) of the pyramidal AFM tip. (g) Aperture formed in tipless cantilever. (h) Whole image and (i) magnified image of the hollow probe fabricated by SU-8. (j) Microsphere tip prepared based on the microchanneled cantilever. (b) and (d)–(g) are reprinted with permission from Ref. [48], © Elsevier Ltd. 2014. (c) is reprinted with permission from Ref. [49], © Springer Nature 2019. (h) and (i) are reprinted with permission from Ref. [50], © IOP Publishing Ltd. 2016. (j) is reprinted with permission from Ref. [51], © The Royal Society of Chemistry 2017.
adsorbing, extracting, transporting, placing, delivering, and injecting) [46] by controlling the pressure of the fluidic channel in the AFM cantilever. Besides, experiments have shown that even though the hollow cantilever is influenced by small changes in pressure, the overall system can suppress this effect in order to ensure constant interaction forces with the specimen at all times during an experiment [43], making the FluidFM probe a highly force-sensitive nanopipette.

Diverse FluidFM probes have been developed for biomedical applications. The FluidFM cantilever is rectangular shaped (~150 μm long and 25 μm long) with an embedded 1-μm high microfluidic channel supported by parallel rows of pillars and an aperture at the free extremity. First, hollow cantilevers are produced on a wafer scale combining photolithography for standard AFM probe fabrication and additional protocols based on the “sacrificial layer” principle for fluidic functionality [46, 47]. After that, an aperture on the AFM tip can be formed either by photolithography (parallel process) or by focused ion beam (FIB) (serial process) [25]. FluidFM probes are commercially available (Cytosurge AG, Switzerland), facilitating the FluidFM studies. With these methods, different types of microchannelled probes can be fabricated [48, 49], as shown in Figs. 1(b)–1(g).

From the scanning electron microscopy (SEM) images of the fabricated hollow cantilevers, the microchannels in the cantilever are distinctly discernible (Figs. 1(c) and 1(d)). The aperture can be formed at the apex of the tip (Fig. 1(e)), at the side of the tip (Fig. 1(f)), or directly in the tipless cantilever (Fig. 1(g)) for different experimental purposes. Besides, polymer-based hollow probes (Figs. 1(h) and 1(i)) can be fabricated by using the epoxy-based photoresist SU-8 [50], offering a new alternative to conventional silicon-based hollow cantilevers with more flexibility in terms of complex geometric design and surface chemistry modification. Particularly, based on FluidFM, AFM tips with special shapes can be prepared in a direct way; for example, attracting single microspheres to the aperture of the probe by applying underpressure yields the microsphere probe [51] (Fig. 1(j)). A notable point is that the hollow probes need to be functionalized with an antifouling coating prior to experimentation to avoid biofouling (biofouling is the non-specific adsorption of biomacromolecules to the probe surface, which can cause the clogging of the flow through the channel in the probe) [52]. The hydrophilic copolymer poly(L-lysine)-graft-poly(ethylene glycol) (PLL-g-PEG), the gold standard for nonfouling surface coating [53], has been extensively used to confer antifouling properties of the FluidFM probes [52]. Recently, researchers have developed a simple protocol for the antifouling modifications of FluidFM probes, relying on the protein resistant copolymer called PACAm-g-(PMOXA, NH2, Si) [49, 54, 55]. For this method, FluidFM probes are first treated by oxygen plasma and then immersed in a PACAm-g-(PMOXA, NH2, Si) solution for 90 min at room temperature. The FluidFM probes can then be used for experiments after rinsing with pure water.

3 FluidFM for single-cell manipulations

3.1 Picking and placing single cells

By approaching the FluidFM probe onto a cell in contact mode and applying a underpressure in the fluidic channel [48], the cell can be tightly attracted to the aperture of the cantilever (I) in Fig. 2(a) in the vertical direction and therefore can be picked up detaching from the surface. By reversing the pressure, the cell can be released onto a desired spot (II) in Fig. 2(a)). For adherent cells which can attach to and spread on the substrate, the prerequisite of picking up the cell is removing the molecular anchors between cell and substrate. Taking advantage of the FluidFM multifunctionality, trypsin solution can be loaded in the channel and locally delivered to individual adherent cells [56] (III in Fig. 2(b)), which activates the selective detachment (rounding) of the targeted cell while neighboring cells remain adherent (III in Fig. 2(b)). The detached cell can then be easily lifted up and manipulated by the FluidFM, providing an efficient way for precise isolation of chosen cells in a cell population. In this way, we can transfer cells to targeted areas to observe cell behaviors (Fig. 2(c)) and we can also sort different types of cells under the guidance of fluorescent labeling (Figs. 2(d) and 2(e)). Besides, FluidFM is able to assist in the formation of cell patterns [57] (Fig. 2(f)). Patterning of cells is a critical physiological process for regulating the life activities such as embryonic development and tissue morphogenesis [58, 59] and investigating cell patterning behaviors benefits the understanding of the physiological/pathological processes. With FluidFM-based single-cell manipulations, we can not only place cells at specific positions on the same substrate or even of another substrate, but also can remove unwanted cells to facilitate forming desired cell patterns [57], which is useful for the studies of patterning at single-cell level.

3.2 Sucking colloidal beads as indenters

FluidFM opens the way to exchangeable colloidal AFM for single-cell mechanical assays. To probe the mechanical properties of cells, spherical AFM tips [60, 61] are commonly used, since the conventional conical tips only detect the local mechanics of the cell. Spherical tips have well-defined geometry, which facilitates theoretical modeling of the indenting process [62]. Besides, spherical tips prevent the local damage of the cell membrane caused by the serial poking of AFM sharp tips during indentation assays, for thus more reliable and safer measurements are obtained using spherical tips [63]. In the case of FluidFM, a single sphere can easily be aspirated to the aperture by applying a negative pressure [64]. Both tipless (I) in Fig. 3(a) or pyramidal FluidFM probes (II) in Fig. 3(a) can be used to prepare spherical tip, and specifically pyramidal probes are suited for preparing spherical tips with smaller...
sizes. Under the guidance of optical microscopy, the FluidFM probe is moved to touch individual spheres deposited on the substrate ((III) and (V) in Fig. 3(a)). Subsequently, applying adequate underpressure to the fluidic channel of the probe allows the adsorption of the sphere to the aperture of the probe ((IV) and (VI) in Fig. 3(a)) [51]. The spherical probes prepared by FluidFM can then be used for experiments such as probing the mechanics of single cells (Fig. 3(b)). Experimental results have significantly shown the capabilities of the FluidFM-prepared spherical tips in sensing the mechanical properties of cells [65]. A notable point is that the sphere sucked to the FluidFM cantilever can be exchanged in the working medium when probing cells. Applying positive pressure yields the release of the sucked sphere, and then moving the probe to the region where the colloidal beads had been previously deposited and applying negative pressure will adsorb a new clean one. This is particularly useful for probing the specific molecular interactions which require the functionalization of the tip (e.g., linking specific molecules to the surface of the tip [66]). For example, we can firstly approach a cell with a bare sphere as control, and then detect specific molecular interactions on the cell surface with an ad-hoc chemically functionalized sphere.

3.3 Dispensing biomolecules on top of single cells

FluidFM is able to deliver not only trypsin, but whatever biomolecules as chemical stimuli to selected living cells on a Petri dish. Under the guidance of optical microscopy, the microchanneled probe filled with a particular chemical solution is gently approached onto a selected cell relying on the AFM feedback and maintained in contact with the cell membrane ((I) in Fig. 4(a)). Subsequently, by applying adequate overpressure, the solution inside the channel can be released to the cell under the probe. Taking a fluorescent dye solution as in [25], which is known to diffuse into the cytoplasm of the targeted cell, the obtained fluorescent images ((II) and (III) in Fig. 4(a)) clearly confirm the successful staining of the targeted cell. In addition to small molecules, single virus can be delivered to the targeted cell by FluidFM dispensing [67]. The microchannel filled with the solution containing a virus conjugated with fluorescein for visual recognition, is brought into gentle contact with an optically selected cell ((I) and (II) in Fig. 4(b)): Applying an adequate pressure results in the release of a single virus to the targeted cell ((III) in Fig. 4(b)). The viral infection process on the targeted cell can then be monitored by fluorescence microscopy ((IV) and (VI) in Fig. 4(b)). These
experimental results [25, 67] show that liquids can be locally delivered to single cells with volume control by the FluidFM, which is particularly promising for investigating the single-cell behavior in response to external chemical stimuli such as drugs.

### 3.4 Intracellular injection of single cells

Using a pyramidal tip with a side-hole preserving the sharpness of the apex, FluidFM is able to puncture the cell membrane for intracellular delivery. Membrane perforation (I) in Fig. 5(a) is indeed achieved by exerting a higher indentation force on the cell surface, which is visually confirmed by the abrupt peak in the recorded force curves [25]. Subsequently, by applying overpressure to the fluidic channel of the probe, the solution inside the microchannel can be directly injected to the interior of the cell. For example, experiments have clearly shown that the injected dye molecules (that cannot otherwise diffuse across the cell membrane) successfully stain the cytoplasm of the targeted cell ((II) in Fig. 5(a)). Besides, FluidFM can even directly deliver liquids into the nucleus of single cells [68]. Typical events during the piercing process can be determined from the force curves ((I) in Fig. 5(b)), including penetration of upper cell membrane, penetration at the top of the nuclear envelope, penetration at the bottom of the nucleus envelope, penetration of lower cell membrane, also confirmed by confocal fluorescent imaging [69] ((II) in Fig. 5(b)). With this method, plasmid DNAs encoding green fluorescent protein (GFP) have been injected into the cell nucleus, and two days after injection the cell successfully produces GFP confirmed by fluorescence (Fig. 5(c)) [68]. Notably, delivering biomolecules via membrane perforation (Fig. 5) is destructive, but this way allows fast delivery to targeted cells and thus facilitates improving efficiency. On the contrast, delivering biomolecules via dispensing (Fig. 4) is non-destructive to cells, but this way is dependent on the diffusion of the dispersed biomolecules, which is much slower than membrane perforation. Consequently, these two methods of biomolecules delivery (dispensing and intracellular injection) provided by FluidFM are complementary and can be used flexibly in practice for different experimental purposes.

### 3.5 Intracellular extraction of single cells

FluidFM allows quantitatively extraction of the intracellular content of single living cells with spatiotemporal control for downstream analyses [70]. The procedure of intracellular structure extraction on single living cells is schematically shown in Fig. 6(a). The procedure consists of controlling the tip of the microchanneled probe to insert into the targeted cell and then applying adequate negative pressure to extract intracellular structures, followed by withdrawing the probe from the cell and dispensing the extraction on the substrate for analyses.

With this generic method, we can not only extract from the nucleus of single living cell (Fig. 6(b)), but also from the cytoplasm on the living cell (Fig. 6(c)). Cells expressing GFP were used to facilitate visually observing the extraction process (Figs. 6(b) and 6(c)) and the extracted intracellular structures (Fig. 6(d)). Since the geometrical parameters of the hollow probe are known, the volumes of the extracted structures can then be precisely quantified according to the fluorescent image of the extraction (Fig. 6(d)). The subsequent cell viability assays showed that after the extraction of volumes up to 4.0 pL from the cytoplasm, 82% of the cells remained viable, indicating that the cells have the ability to withstand the loss of a large proportion of the cytoplasm. The cells were still viable after the extraction of volumes up to 0.6 pL from the nucleus, indicating that extraction from the cell nucleus was more critical than extraction from the cytoplasm. The extracted intracellular structures of the targeted cell can then be used for various biochemical analyses, including structure characterizations, gene expression analysis, enzyme activity assay, mass spectroscopy, and so on [70, 71]. We know that there are not only biochemical alterations but also mechanical changes.
understanding cellular processes.

In principle, FlowFM single living cell extraction provides a feasible way for directly investigating the relationship between cellular properties and their behavior. Single living cells can be quantitatively extracted for biochemical assays \([70, 71]\), therefore providing a feasible method to study the properties of single living cells \([32]\), after which the intracellular content was extracted, respectively. \(d\) Different volumes of GMP-containing fluids were extracted in the hollow probes. (Reprinted with permission from Ref. \([70]\), \(\copyright\) Elsevier Inc. 2016).

4.1 Adhesion of eukaryotic cells

The adhesion forces between single living cells and substrates can be quantified by FluidFM. Traditional AFM-based SCFS \([74]\) is based on attaching a living cell to the AFM cantilever via complex chemical functionalization (for example, according to the established protocol in SCFS \([75]\), the cantilever is firstly coated by biotin-labeled bovine serum albumin, and then streptavidin is coated on the cantilever, which is followed by the coating of biotin-labeled concanavalin A), which is labor-intensive and requires specific expertise \([76]\). Particularly, chemically immobilizing the cell onto the AFM cantilever may inevitably influence the activities of the cell (e.g., cell surface damage and denaturation \([77]\)). With the use of FluidFM, single living cells can easily be physically immobilized onto the AFM cantilever by applying and maintaining adequate suction force \([78]\). FluidFM facilitates increasing the throughput and efficiency of SCFS assays \([78]\).

Based on FluidFM, in 2012, Potthoff et al. \([79]\) firstly showed the serial SCFS measurements using a single cantilever for sensing the adhesive interactions of multiple mammalian cells (HeLa cells and HEK cells). Besides, the adhesion forces of human umbilical vein endothelial cells (HUVECs) on different surface topographies were measured by FluidFM \([80]\), benefitting the reveal of the effects of surface textures on promoting endothelialization. In 2016, McGrath et al. \([81]\) investigated the adhesion interactions between single cells (mouse myoblast C2C12 cells) and different types of substrates \((I)\) in Fig. 7(a)). Two types of RGD-presenting surfaces (non-covalent surface and covalent surface) were analyzed. After adsorbing single living C2C12 cell with FluidFM probe, the cell probe was controlled to perform approach-retract movement in the vertical direction on the substrates (non-covalent surface or covalent surface), during which force curves \((II)\) in Fig. 7(a)) were recorded. From the force curve, several parameters reflecting the unbinding between cell and substrate can be obtained, including adhesion force \((III)\) in Fig. 7(a)), detachment distance \((IV)\) in Fig. 7(a)), and the binding energy \((V)\) in Fig. 7(a)), facilitating quantitatively understanding the adhesive interactions between cells and substrates. FluidFM-based SCFS has also been utilized to address various issues regarding cell adhesion, including the effect of electric current on cell adhesion \([83]\), intercellular adhesion forces in mature cell-cell contacts \([84]\), the adhesion of cancer cells at various stages of malignant transformation \([85]\), cell-sheet engineering \([86]\), cell-material interaction \([87]\), and single-cell adhesion assays with high throughput \([88]\), contributing much to the studies of single-cell adhesion in physiological/pathological processes.

4.2 Adhesion of microbial cells

FluidFM-based SCFS benefits the exploring of the underlying mechanisms guiding the adhesion interactions of individual microbial cells. Biofilms, a complex group of microbial cells that adhere to the exopolysaccharide matrix present on the surface of medical devices, play an important role in the pathogenesis of implant infections \([89]\). Originally microbes attach onto the surface of biomaterials, after which the intercellular interactions mediated by adhesins and cell wall proteins lead microbes to cluster together, forming microcolonies and biofilms \([90]\). Cell adhesion is mediated by a multitude of molecular interactions that are specific (i.e., molecular recognition between receptors and ligands) or non-specific (i.e., hydrogel bonding, hydrophobic, van der Waals, electrostatic, and hydrophobic interactions).
and macromolecular forces), but so far the precise mechanisms by which these interactions determine cell adhesion processes are not yet fully understood [91]. With the use of FluidFM, a single microscope can be easily immobilized to the AFM cantilever by applying underpressure, and then the adhesive interactions between microbes and substrates (or microbes) can be measured. In 2015, with the use of pyramidal microchanneled cantilever, Potthoff et al. [92] firstly showed that FluidFM is able to suck single bacterium and then measure the adhesion forces between bacteria and substrates. In 2017, Sprecher et al. [93] showed that FluidFM-based SCFS could benefit the investigation of the exopolysaccharide adhesins in bacterial surface colonization and biofilm formation. Candida albicans is the principal cause of opportunistic mycoses worldwide and the agglutinin-like sequence (Als) protein on the surface of C. albicans is closely related to the adhesion, aggregation and biofilm formation of C. albicans. [94]. In 2019, the studies by Dehullu et al. [95] with the use of FluidFM-based SCFS discovered a previously undescribed function (mediating amyloid-like homophilic adhesion) for C. albicans Als proteins. A yeast cell was immobilized on the aperture of the hollow cantilever by applying a negative pressure, and another yeast cell was mechanically trapped in porous membrane (II) in Fig. 7(b)). Staining the cell with live/dead indicator dye confirmed the integrity of the cell (II in Fig. 7(b)). The interactions between yeast cells at different conditions (e.g., altering the amyloid sequence or lowering the protein density on the cell surface) were then measured by performing SCFS to record force curves (III in Fig. 7(b)). The results significantly revealed a novel aggregation mechanism of C. albicans, in which force-dependent amyloid core sequences in Als proteins play a dual role, that is, in formation of adhesin nanoclusters and in homophilic bonding between amyloid sequences on opposing cells. Recently, the optimization of measurement parameters (e.g., setpoint, z-speed, z-length, pause time, and relative underpressure) when applying FluidFM-based SCFS on bacterial cells has been investigated [96]. As an alternative way, the bead whose surface is coated with chemical molecules can be attached to the aperture of the microchanneled cantilever, which is then used to probe the interactions between individual microbial cells (immobilized on the substrate) and bead surface [97]. In future, FluidFM may contribute to the identification of small peptide inhibitors [98] and titanium surface [99] for antiadhesion therapy.

4.3 FluidFM limitations for cell adhesion

The serial cell-adhesion protocol is without doubt the most straightforward one of those established with the FluidFM because of its broad validity. Nevertheless, it is not immune to drawbacks. If applied to mammalian cells, it is often observed that cell debris remain attached at the aperture edge despite the anti fouling coating, which then requires a cleaning step by immersing in an appropriate solution that can take place keeping the FluidFM probe mounted on the AFM head. It works although there is a risk that the cleaning solution may affect the anti fouling layer and thus jeopardizes its efficiency. On the other hand, if applied to bacteria, one has to be aware of the compromise between size of the aperture and adhesion...
strength. Bacteria need a small aperture to be efficiently attached at it, yet a small aperture means a small suction force (a few tens on nanoNewtons for an aperture diameter of around 300 nm and an applied underpressure of maximal 1 atmosphere [25]) sometimes not able to overcome the adhesion strength of the bacteria under investigation.

5 FluidFM for single-cell electrophysiology

5.1 Force-controlled patch clamp

FluidFM could inspire original developments for patch clamp. Ion channels are a diverse family of membrane-spanning proteins that lower the free energy required for ions to traverse the plasma membrane [100], and ion channels play an important role in regulating the life activities of cells [101]. Malfunctioning of ion channels has been implicated in human diseases (called channelopathies [102]) such as lysosomal storage disorders, neurodegenerative diseases and metabolic pathologies, as well as the progression of certain infectious diseases [103]. The patch clamp technique is an electrophysiological method that allows the recording of macroscopic whole-cell or microscopic single-channel currents in single cells using glass micropipettes filled with an electrolytic solution [104] and patch clamp has been the gold standard of the studies of ion channel for decades [105]. Combining FluidFM with patch clamp techniques allows precise monitoring of the mechanical forces exerted on cells while simultaneously sensing the ionic currents of the cell [106]. For doing this, an electrode was inserted into the reservoir of the FluidFM probe, and another electrode was placed in the bath solution (150 mM KCl) (Fig. 8(a)). A cylindrical aperture with a diameter of about 350 nm was milled by FIB at the pyramid apex (Fig. 8(b)). After selecting a cell on the dish, the FluidFM probe was positioned on top of it under optical microscopy guidance (Fig. 8(c)) and the probe was approached on the membrane of the targeted cell via the AFM force feedback. By applying a slight underpressure a seal was formed in the cylindrical aperture of around 150 MΩ allowing simultaneously monitoring ionic current and the force of the nanopipette (Fig. 8(d)). The contact between the tip and the cell is stable due to the automated force feedback of AFM, whereas the contact is prone to get lost in conventional patch clamping due to vibrations or cell volume changes. The currents in response to conventional voltage pulse were recorded and the superimposed whole-cell Na+ currents of the Na.v1.5 channel confirmed the specific electrophysiological characteristics of the Na.v1.5 ion channel (Fig. 8(e)). Based on the method, the whole-cell ionic currents and the contraction forces of mouse adult cardiomyocytes were simultaneously recorded (Figs. 8(f) and 8(g)), clearly showing the rhythmic movements of the cardiomyocyte as well as the changes of whole-cell ionic currents. Due to the unique advantages of the FluidFM-based patch clamp, it is particularly suited to investigate the behaviors of mechanosensitive ion channels [107, 108] which are activated by mechanical forces for revealing the principles of ion-channel-mediated mechanosensory transduction [109].

5.2 Force-controlled scanning ion conductance microscopy (SICM)

FluidFM can be also operated in simultaneous SICM mode. So far, it is still challenging for regular AFM to resolve the fine structures of living mammalian cells. First, the very soft and dynamic nature of the cell surfaces makes imaging living mammalian cells difficult [76]. Besides, the mechanical contact between AFM tip and cell surface usually deforms cellular surface molecules, which in turn results in the blurring of the recorded images [110]. To solve this issue, we would need a scanning probe technique in which the probe is not in direct physical contact with the cell surface, which is known as SICM [111]. Studies have shown that on very soft samples such as living cells, SICM can be advantageous to AFM in terms of imaging quality, accuracy in topography measurements, lateral resolution, and long-term imaging stability [112]. SICM uses an electrolyte-filled glass pipette (nanopipette) as a probe and relies on an ion current flowing between an electrode inside the nanopipette and another in an external bath solution [113]. The ionic current passing through the apex opening of the nanopipette is sensitive to the tip-sample separation, allowing non-contact topographic imaging of the specimens. Implementing SICM on the FluidFM allows simultaneously recording mechanical and conductance information of the specimens [114]. An electrode is inserted in the fluidic circuit of the FluidFM, and another electrode is in the bath solution (Fig. 9(a)). The ionic current through the microchannled cantilever is monitored with a patch clamp amplifier. Under this configuration, controlling the FluidFM probe (Fig. 9(b)) to scan the specimen allows recording simultaneously both the deflection of the cantilever and the ionic current. The topography images of specimens are obtained at non-contact...
mode by using the ionic current as the feedback, and the force images of specimens are generated simultaneously from the force feedback of the cantilever. Force spectroscopy experiments performed on substrates show the simultaneously obtained ionic curve and force curve (Fig. 9(c)), which delivers mechanical and electrical information. Experiments performed on standard grids (Fig. 9(d)) and living neurons (Fig. 9(e)) simultaneously show the topographic images and force images of specimens. The results [114] significantly show that FluidFM provides novel possibilities to SICM by integrating AFM force feedback, which will be particularly meaningful for investigating the structures and multiple properties (e.g., mechanics and electronics) of single cells.

5.3 Force-controlled scanning nanopore microscopy

The aperture at the pyramidal apex of the FluidFM probes can also be interpreted as a nanopore for the stochastic sensing of secreted molecules and the activity of ion channels in arbitrary locations both outside and inside a cell. Observing the activities of single molecules of single living cell in their native states is of critical significance for the investigations of cell signaling and molecular mechanisms [115]. Traditional single-molecule methods require the fluorescent labeling of the targeted molecules [116–118], which may influence the activities of molecules and cells. In fact, strictly speaking, the results obtained by single-molecule fluorescence microscopy only reflect the behaviors of the individual fluorescent spots but not the targeted biomolecules themselves [119]. Asymmetric glass capillary nanopore electrode has been used for single-molecule detection on single living cells [120], but the measurement precision is limited due to the lack of sensitive feedback. FluidFM can be used as scanning nanopore microscopy to sense the activities of biomolecules using simultaneous force and ion-current feedback [49]. For doing this, the apex of the tip is flattened by ion sputtering to produce a thin film of silicon nitride and then a small hole is formed in the middle of the thin film using focused He⁺ ions (Fig. 10(a)). We can shrink the pore to a desired size by raster scanning of the He⁺ ions across the pore (Fig. 10(b)). The nanopore provides the nanoscale confinement of biomolecules between the apex and the underlying surface (I) in Fig. 10(c)) and sets a barrier for the biomolecules before their entrance into the nanopore (III in Fig. 10(c)). The current–voltage (I–V) curve of the nanopore near the surface is recorded, which is dependent on the protein translocation through the nanopore, allowing the detection of single biomolecules. With FluidFM-based nanopore, not only the topographical image and the ionic current map of single living cell can be obtained (Fig. 10(e)), but also the proteins secreted by the targeted cell can be real-timely monitored via distinctive blockades and/or enhancement in the current traces (Fig. 10(f)). Besides, FluidFM-based nanopore enabled intracellular recordings at particular parts of the targeted cell (Fig. 10(g)). A large number of current spikes were observed (Fig. 10(h)), indicating the translocation of intracellular elements through the nanopore. Overall, the results [49] showed the great potentials of FluidFM-based scanning nanopore microscopy in addressing biological issues with unprecedented spatial resolution and force/ionic current sensitivity, which will contribute to the studies of single-cell electrophysiology at molecular level.

5.4 FluidFM limitations for electrophysiology

If the force feedback of the FluidFM is an advantage with respect to glass capillaries as far as the stability of the contact with the cell membrane, the tough challenge for the FluidFM is to show the capability of consistently obtaining the compulsory gigaseal. The pyramidal tip being not the most indicated for the establishment of the gigaseal due to the corners between the facets, it will be mandatory to explore the microfabrication of hollow tips with a cylindrical symmetry. Concerning the
nanopore mode, the promising aspect is surely related to the possibility to carry out the measurements close to or even within a cell; nonetheless the huge challenge is to reliably determine the identity of the biomolecules passing through the nanoaperture. Moreover, it has to be established whether translocating biomolecules are released by the approached cell or are from the cell culture medium [121].

6 Conclusion and future perspectives

In this review, the achievements in single-cell manipulations and detections using FluidFM in the past decade since its invention have been summarized, considerably demonstrating that FluidFM has become a powerful multifunctional toolbox for single-cell biophysics, including various single-cell manipulations (e.g., three-dimensional operation [48], cell sorting [56], cell patterning [57], exchangeable colloidal probe for cell indenting [65], drug dispensing & injection with very small volumes [25], cell nucleus injection [68], individual virus stimulation [67], and cellular extraction [70]), single-cell force spectroscopy (e.g., cell–substrate interaction measurements [82] and cell–cell interaction measurements [95]), and single-cell electrophysiology (e.g., simultaneous mechanical force and ionic current recording for patch clamp [106], force-controlled SICM [114], and single-molecule sensing based on scanning nanopore microscopy [49]), providing novel possibilities for investigating the behaviors of single living cells, which is significantly meaningful for the studies of revealing the underlying molecular mechanisms guiding cellular physiological/pathological processes.

6.1 Technical challenges and developments

There is still considerable room for the advancement of FluidFM. Notably, at present, the three-dimensional manipulations by FluidFM (via controlling the pressure of the fluid channel in the FluidFM probe) are dependent on the guidance of optical microscopy. For the objects which are invisible for optical microscopy, the objects being manipulated by FluidFM need to be labeled with fluorescence for visual recognition; for example, when manipulating single virus, the virus was labeled with fluorescein [67]. Besides, when applying FluidFM to deliver/inject liquids to single cells [56] or to extract intracellular ingredients [70], fluorescent dyes are mixed in the solution or cells are engineered to express GFP for visually exact determination of the volumes of the delivered liquids or the extractions. The introduction of fluorescein may influence the activities of the cells or the objects being delivered and therefore the conditions are not fully native. Studies have shown the successful label-free optical imaging of 75-nm adenoviruses by submerged microsphere optical nanoscopy without the use of fluorescent labeling [122], which is inaccessible for conventional optical microscopy. Therefore, introducing the advanced microsphere-aided super-resolution optical imaging to the optical system of FluidFM will help improving the label-free manipulations of FluidFM. For FluidFM-based whole-cell patch clamping, the current seal resistances achieved are typically less than 150 MΩ [106] which is insufficient for reaching gigaseal, significantly influencing the measurement quality of patch clamping. Optimizing the geometry of the FluidFM probes will be mandatory to improve the seal resistance for patch clamp [52].

6.2 Future directions for FluidFM-based single-cell biophysics

Further explorations using FluidFM will contribute to answering fundamental issues involved in biomechanics at the single-cell
level. So far in practice for standard AFM-based studies of cell mechanics in response to external chemical stimuli, drug molecules are added into the dish perturbing all the cells on the dish [32], making impossible to estimate how many drug molecules are indeed acting on the targeted cell. With FluidFM, we can precisely deliver known volume of drug molecules to the selected cell on the dish [56] and then monitor the real-time mechanical changes of the targeted cell (specially, due to the fact that the hollow tip is not suited for AFM indentation assays, single microsphere can be immobilized to the aperture of FluidFM probe for measurement [51] prior cellular mechanical measurements). Particularly, since the delivery is locally for the targeted cell whereas the other cells are probably not influenced by the delivered drug molecules, it is very convenient for control experiments; for example, we can perform measurements on the cells that are not influenced by the drug molecules in the same dish. Since the measurements are performed in identical conditions (e.g., in the same dish, in the same cell culture medium, and in the same environment), it fully meets the requirements of comparative studies [123]. These experiments will be particularly useful for revealing the detailed dynamics of cell mechanics taking place after the treatment of known small volumes of drug molecules on single cells. Besides, with the use of FluidFM’s three-dimensional capabilities, cells can be picked up and placed on the different substrates with diverse properties (such as stiffness) for measurements, and therefore this will significantly facilitate examining the exquisite effects of substrate properties on the cellular mechanics (e.g., elasticity, viscoelasticity, and adhesion force [124]) at the single same cell level, which is quite meaningful for the studies of cell mechanics.

Due to the intrinsic geometry of the apex of the hollow probes, FluidFM is not suited for high-resolution morphological imaging of the samples. Combining FluidFM with complementary techniques will benefit the investigation of the activities of single cells with subcellular resolution. Based on the single-cell manipulation of FluidFM, intracellular substances of single living cells can be extracted for further biochemical assays [70], which can provide the chemical properties of the cell. Besides, we can use regular AFM high-resolution morphological imaging and mechanical analysis [28] to characterize these intracellular structures. So far studies of organelles by AFM were performed on organelles (e.g., cell mitochondria [125] and cell nucleus [126]) which are isolated from the lysis of the ensembles of cells and therefore the results only reflect the averaged behavior of organelles from cell populations. Combining FluidFM with regular AFM as well as biochemical assays, we can probe the activities of single living targeted cell, after which the structures and properties (mechanics and chemical properties) of the organelles of the targeted cell can be characterized, thus providing a feasible way for correlating the structures and properties of organelles with the behaviors of the same cell with subcellular resolution. The origin and spread of molecular mutations take place at multiple scales (from the organelle to cell and the human population) [127] and recently researchers have shown the great significance of investigating enzymatic activities in organelles for discovering novel mechanisms guiding critical cellular functions [128]. Consequently, combining FluidFM with complementary techniques to investigate cell behaviors at subcellular resolution will have general impacts on the studies of life sciences.

Taken together, the developments of FluidFM significantly provide novel possibilities for single-cell biophysics, which will have great impacts on the communities of biomedicine. Notably FluidFM for biomedical applications is still in its infancy and there is huge room for further advancements.

In the future, as more biological systems are investigated by FluidFM, particularly combined with other complementary techniques, we have much to look forward to.

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