Lipid droplets as endogenous intracellular micro lenses

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Abstract

Using a single biological element as a photonic component with well-defined features has become a new intriguing paradigm in biophotonics. Here we show that endogenous lipid droplets in the mature adipose cells can behave as fully biocompatible microlenses to strengthen the ability of microscopic imaging as well as detecting intra- and extracellular signals. By the assistance of biolenses made of the lipid droplets, enhanced fluorescence imaging of cytoskeleton, lysosomes, and adenoviruses has been achieved. At the same time, we demonstrated that the required excitation power can be reduced by up to 73%. The lipidic microlenses are finely manipulated by optical tweezers in order to address targets and perform their real-time imaging inside the cells. An efficient detecting of fluorescence signal of cancer cells in extracellular fluid was accomplished due to the focusing effect of incident light by the lipid droplets. The lipid droplets acting as endogenous intracellular microlenses open the intriguing route for a multifunctional biocompatible optics tool for biosensing, endoscopic imaging, and single-cell diagnosis.

Introduction

With the demand in real-time monitoring of endoplasmic variations and rapid detection of extracellular signals, a great number of approaches to bioimaging have been developed, including electronic, X-ray, optical, ultrasonic, magnetic, thermal, and mechanical methods, to obtain the significant information about the physiological and pathological processes of cells¹–³. Among these approaches, optical microscope imaging techniques become preferable as they enable a straightforward and real-time visualization that is highly desirable for observation and diagnostics in vivo. To improve the discernibility to cells and tissues in different species, fluorescence signals are usually employed in optical microscope imaging techniques, which provides the possibility of monitoring and detection of specific organelles and cells⁴.

However, the fluorescence signals emitted from subcellular structures are generally very weak so that it becomes difficult to directly apply the fluorescent methods to intracellular imaging or detection for single-cell studies. Although the fluorescence intensity can be increased by increasing the power of excitation light, the risk of phototoxicity to cells becomes higher and the photobleaching effect becomes stronger at the same time⁵–⁷.

The past few decades have witnessed a dramatic progress in optical imaging⁸–¹⁰, especially with the emerging of microsphere-assisted techniques that enable real-time and super-resolution imaging at visible wavelengths with conventional optical microscope⁸,¹¹–¹⁶. The microspheres also provide the opportunities to take advantage of some photonic functions such as directional antennas, whispering-gallery modes, and photonic nanojets to enhance the optical signals including fluorescence¹³,¹⁷, upconversion emission¹⁸, Raman scattering¹³,¹⁹, and backscattering¹⁷,²⁰. However, most of the microspheres in current strategies are in solid and artificially synthetic materials (e.g., SiO₂, polystyrene, BaTiO₃, TiO₂) and so...
they are of very low biocompatibility, which becomes a fundamental aspect of biomedical applications when contact with the sample is needed, for example in single-cell analysis in vivo or endoscopy\textsuperscript{21}. Fortunately, lessons from nature have shown that some bio-components and already-existing objects (e.g., red blood cells, yeast cells, spheroid cells, spider silks) can interact with light and are able to take the functions as real optical elements (i.e. microlenses, resonators, waveguides, gratings or tools for photolithography)\textsuperscript{22–31}. The lensing effect of some biological samples indicates the possibility to construct fully biocompatible microlenses with proper focal length and magnification to achieve real-time monitoring and detection of optical signals of a single cell\textsuperscript{23–25}.

In this work, we demonstrate that lipid droplets, a sort of dynamic structures that naturally exist in cells\textsuperscript{32}, can act as intracellular microlenses for real-time monitoring of subcellular structures and detection of extracellular signals. With a spherical shape and a refractive index higher than cytoplasm and periplasm\textsuperscript{33,34}, the lipid droplets exhibit the lensing effect to efficiently converge both the excitation light and the fluorescence signals. Driven through optical tweezers, we show that single lipid droplets can be moved inside the cell to perform an enhanced intracellular fluorescence imaging of cytoskeleton. Moreover, we demonstrate that due to the refractive index contrast of the lipid droplets in respect to cytoplasm, the excitation light is suitably focused for enhancing the mitochondrial fluorescence signals from cancer cells in the extracellular fluid. Thus, we proof that the lipidic microlenses are very useful and real-world tools able to assist fluorescent imaging.

### Results

#### Materials characterization, experimental setup, and lensing effect analysis

The lipid droplets used in this work were those naturally existing in commercially available human adipocytes-visceral cells (Fig. 1a, see “Methods” for cell culture). The droplet generally maintains a smooth surface, micrometer size (0.5–40 \(\mu\)m), and a spherical shape\textsuperscript{35} (Fig. 1b; also see Fig. S1 and Video S1 of the Supplementary Information). Meanwhile, the droplets exhibit a high transparency at visible wavelengths (see the transmittance spectrum in Fig. S2 of the Supplementary Information) and a refractive index of \(\sim\)1.52 (see “Section” S1.1 of the Supplementary Information), higher than that of the cytoplasm (\(\sim\)1.36). With these morphology and optical properties, the lipid droplets are expected to act as a sort of endogenous lensing structures. The experimental setup was built around a scanning optical tweezing system (OTS) incorporated with an inverted optical microscope for detecting the bright-field and fluorescence signals (Fig. 1c1) (see “Methods” for the details of the experimental setup). The trapping light in the OTS was a laser beam (Gaussian beam) at the wavelength \(\lambda\) of 1064 nm that is within the biological window\textsuperscript{36}. The range of a single optical trap created by the trapping light of the OTS, i.e., the spot size...
of the beam, is \(\sim 1.3 \mu m\), and a single lipid droplet can be attracted and moved by a single optical trap at a time. With the assistance of such an OTS, the lipid droplets can be trapped and moved inside the cells, acting as transportable intracellular microlenses (Fig. 1c2).

In the case of intracellular imaging, the objects were generally underneath lipid droplets due to the intrinsic pressure inside the cell\(^{37}\) and so the droplets worked in a contact mode of microlenses\(^{8}\). To analyze their lensing effect in contact mode, the lipid droplets were extracted from the adipose cells to index matching liquid that was matched with the cytoplasm (\(\sim 1.36\)). Bright-field imaging was performed on a grating sample (width: 650 nm; spacing: 350 nm), and a magnification of \(\sim 2.1\) can be obtained by using the \(\sim 3.2 \mu m\) lipid droplet (see Fig. S3 of the Supplementary Information). A finer grating structure of a Blu-ray Disk (width: 200 nm, spacing: 100 nm) was also used as a sample for imaging in bright field (see Fig. S4a1, b of the Supplementary Information). Without the lipid droplet in the field of view of the optical microscope (objective \(\times 60, NA = 1\)), the gratings of the disk could not be resolved. With the presence of the lipid droplets, the adjacent gratings became discernible. The results indicate that in contact mode, a feature size of \(100 \text{ nm}\) can be resolved with the assistance of the lipid droplets. A possible mechanism of the gain in imaging ability is that the near-field evanescent waves that carry high spatial frequency information of the sample are coupled into the lipid droplets and transformed into propagating waves for microscopic observation\(^{38,39}\). Besides, the presence of the lipidic microlenses reduces the full width at half maximum of the point spread function of the imaging.

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**Fig. 2 Lensing effect of the lipid droplets.** a Images merged by bright-field and fluorescence images of the FNDs. a1 Bare FNDs on a glass slide. a2 FNDs with a lipid droplet (diameter: 9 \(\mu\)m) moved right below them. b Optical intensity distributions. c FOM as a function of the droplet diameter. The inset shows the optical intensity distributions of the FNDs with and without the 9-\(\mu\)m lipid droplet at different excitation powers (\(P_1\) and \(P_2\)) with the same detected maximum fluorescent intensity. d Energy density distributions of the emitters without (d1) and with (d2) the lipid droplet. e The corresponding emission patterns. f Enhancement factor (\(f\)) as a function of the diameter of the lipid droplets. The inset shows the position of the observation line for the calculation of \(f_1\).
In each of the measurements and imaging processes, an optical power (from 1 to 20 μm) were used to enhance the intensity of detected fluorescence and to obtain the fluorescent images of the FNDs (Fig. 2c). In each of the measurements and imaging processes, an optical power (from 1 to 20 μm) were used to enhance the intensity of detected fluorescence and to obtain the fluorescent images of the FNDs (Fig. 2c). In each of the measurements and imaging processes, an optical power (from 1 to 20 μm) were used to enhance the intensity of detected fluorescence and to obtain the fluorescent images of the FNDs (Fig. 2c).

A figure of merit (FOM) is thereby defined as FOM = (P2 – P1)/P2 × 100%. As shown in Fig. 2c, the FOM values were larger than 35% for the droplets of 1–20 μm, and a maximum of the FOM (73%) can be obtained at D = 9 μm. The results indicate that the presence of the lipid droplets significantly reduces the needed excitation power, thus decreasing the possible photobleaching and photo-damage to the sample.

In order to preliminarily evaluate the fluorescence enhancement by the lipid droplets in the contact mode, numerical simulations by finite element method of the field distributions were performed (see “Methods” for the simulation details). The FNDs were considered as point sources (emitters) that have both longitudinally and transversely polarized components. In the longitudinal polarization, the direction of the dipole was perpendicular to that of the objective, resulting in a very low collection efficiency by the objective (~10.7%). Therefore, the longitudinally polarized dipole is discussed here as an extreme circumstance (Fig. 2d1). Note that in the transversal polarization, as the direction of dipole was parallel to the direction of the objective, most of the light has been collected directly by the objective and the improvement in collection efficiency by the lipid droplets is much weaker than that in the longitudinally polarized dipole (see section S1.3). A lipid droplet of 9 μm was above and in contact with the emitter to act as a microlens (Fig. 2d2). The emission patterns for the emitters without and with the lipid droplet were then obtained by the simulated field distributions in the far field (50 μm) above the emitter, as shown in Figs. 2e1 and 2e2, respectively. The blue shades in Fig. 2e represent the collection angle (±47.3°) of the microscope objective used in the experiments. It can be seen that with the lipid droplet, the emission becomes highly directional (with a divergence angle decreased from 180° to about 110°) so that the emission is redirected to the objective. The collection efficiencies, defined as the ratio of the energy collected by objective to the total energy of the emission, were calculated to be 10.7% and 48.5% for the emitters without and with the lipid droplet (D = 9 μm), respectively. The enhancement factor f1 for the collection efficiency, defined as the ratio of the collection efficiencies with the droplets to those without the droplets (D from 1 to 20 μm), was estimated and presented by the purple dots in Fig. 2f. Since the fluorescence enhancement was also attributed to the focusing of the excitation light, the enhancement factor f2 for the excitation light is defined as f2 = I2/I1, where I1 and I2 are the average intensities at the observation lines (indicted by the blue dashed line in the inset of Fig. 2f), for the cases without and with the lipid droplets, respectively. In contact mode, the observation line was set in the near-field of lipid droplets. However, as the focusing region of the excitation light was beyond the near-field range of the lipid droplets, the values of f2, as presented by the blue dots in Fig. 2f, were much smaller than those of f1. The total enhancement factor f of the lipidic microlenses can then be...
calculated by $f = f_1 \times f_2$, and the results are presented by the red dots in Fig. 2f. It is indicated that the major contribution to the fluorescence enhancement was the increased collection efficiency by the lipid droplets acting as micro lenses in the contact mode. Therefore, the proposed strategy exploits improving the signal collection instead of increasing the excitation power, reducing the risk of photobleaching and photo-damage to cells in live-cell imaging. Furthermore, the lipid droplets can be extracted and patterned to form 2D micro lens arrays and moved rapidly to perform a scan with the OTS followed by a reconstruction of the image by stitching the collected signals (see Fig. S7 of the Supplementary Information), which is a feasible method to overcome the small field of view and the intrinsic spherical aberrations of micro lenses and to achieve high-throughput in scanning imaging and signal detecting in vitro.

**Intracellular imaging of subcellular structures**

By using the endogenous lipid droplets, intracellular imaging was then performed on several subcellular structures in living mature adipose cells. Within the life cycle of the cell, the endogenous lipid droplets vary in terms of their number, size, spatial organization, and the intracellular positioning. With the optical traps created by the OTS, the position in which the lipid droplet can be deployed is confined within a certain region into the live cells. The trapped lipid droplet thus can act as a movable intracellular micro lens, allowing us to achieve real-time imaging and detect the fluorescent signals of the field of interest (e.g. the subcellular structures in Fig. 1c2). With its microfilaments (F-actin filaments) labeled by SiR-actin (red, emission wavelength: 674 nm), a fluorescent living mature adipose cell containing a lipid droplet ($D = 7.7 \, \mu m$) was excited by green light ($540–580$ nm) at an optical power of $2 \, mW$ (intensity: $16 \, mW \, mm^{-2}$) (Fig. 3a1). Without the assistance of the lipidic micro lens, the targeted microfilaments cannot be resolved through the microscope imaging system even in the optimal focal plane. When the lipidic micro lens was trapped and then moved toward the targeted microfilaments by the OTS, the fluorescent signals were efficiently enhanced and the fluorescent image of the branch-shaped microfilaments was clearly visible (Fig. 3a2). Normalized intensity distribution was obtained along the observation lines through microfilaments without (blue dashed in Fig. 3a1) and with (red dashed in Fig. 3a2) the lipid droplet (Fig. 3b), indicating that the resolving ability of the imaging system was greatly improved by the lipid droplet. With the peaks of the intensity distributions through the microfilaments, the magnification ($M$) was calculated as $M = \Delta I_2/\Delta I_1 = 2$, where $\Delta I_1$ and $\Delta I_2$ are the separation between the peaks of the systems without and with the assistance of the lipid droplet, respectively. Also, the contrast $C$ of the fluorescent signals has been increased by the lipid droplet. The contrast $C$ can be expressed as $C = \Delta I/I_{\text{max}}$, where $I_{\text{max}}$ is the maximum fluorescent intensity of the peaks and $\Delta I$ is the drop from the peaks to the valley between them. Without the lipid droplet, the calculated contrast was 14.8%, below the Rayleigh criterion of 26.4%. Instead, with the lipidic micro lens, the contrast was increased to 62.5%.

Further experiments show that the lipid droplets can be used to improve the fluorescent imaging of microfilaments in both living and fixed cells (Fig. 3c, d), and a larger field of view can be achieved with the lipid droplets in larger sizes. Besides the microfilaments, the enhancement in the imaging by the lipid droplets is also exploitable for the lysosomes in living cells in both fluorescence (Fig. 3e1, e2) and the bright-field (Fig. 3e3) modes. The lysosomes were labeled by Lyso-Tracker (red) while the excitation power was $1.5 \, mW$ (intensity: $12 \, mW \, mm^{-2}$). Furthermore, with the feature of long-term existence (steady state) inside the mature adipose cells, the lipid droplets can be used to monitor the dynamic process and physiological state of the cell. The movement of the adenoviruses was monitored by the lipid droplet ($D = 8.1 \, \mu m$) inside the cell at an excitation power of $5 \, mW$ (intensity: $40 \, mW \, mm^{-2}$) (Fig. 3f), and the movement trajectories of the infection process of adenoviruses were recorded in 10 min (Fig. 3f3). The microfilament movement was also monitored by the lipid droplet ($D = 10.0 \, \mu m$) inside the cell at an excitation power of $2 \, mW$ (intensity: $16 \, mW \, mm^{-2}$) (Fig. 3g). The targeted microfilaments gradually moved from the edge to the center of the field of view covered by the lipid droplet in 10 min. Note that the fluorescence image of cells obtained by a conventional optical microscope is the 3D signals superimposed upon a 2D image. The overlap of signals on the $z$ direction leads to the difficulty in resolving the details of subcellular structures and consequently a low signal-to-noise ratio. As the lipid droplets naturally existing inside the cells are very close to the subcellular structures, most of the fluorescent signals from the subcellular structures can be collected by the lipid droplets, which allows us to focus on the physiological processes of the subcellular structures (Fig. 3a, c–e, g) or even virus (Fig. 3f) on a specific plane and suppress the noise signals from non-target regions.

**Non-contact mode and detecting extracellular signals**

In addition to the contact mode in which the collection efficiency is the major contribution to the fluorescence enhancement, the long focal length of the lipid droplets can extend the fluorescence enhancement strategies to non-contact mode, thereby increasing the working distance of imaging and detection of lipid droplets. In the non-contact mode, the concentration of excitation light becomes the
leading contribution so that the detection range of lipid droplets can be extended to the extracellular microenvironment and surrounding tissues. As indicated by the simulated field intensity distributions shown in Fig. 4a, with a plane wave as the excitation light (\(\lambda\): 473 nm), the focal length \(l\) becomes larger with the increase of the lipid droplet diameter. For the diameters larger than 2 \(\mu\)m, the excitation light is focused at \(l > 10 \lambda\) (Fig. 4b), and the intensity at the focus is also increased by the lipid droplet (Fig. 4c), suggesting a non-contact mode of such bio-microlenses in fluorescence. In this mode, the enhancement factor \(f\) of the lipid droplet is calculated given that the target of detection is at the focus of the microlens. It is found that the enhancement factor related to the excitation intensity increases with droplet diameter, but the factor related to the collection efficiency slightly contributes to the total enhancement factor as the near-field collection of the lipid droplet becomes ineffective due to the long distance between the lipid droplet and the emitter (see Fig. S8 of the Supplementary Information). The lensing effect over a long distance in the non-contact mode provides an opportunity to enhance the fluorescence from the extracellular environment surrounding the cells. As reported\(^{44}\), in addition to brain vessels, periadventitial adipose cells are distributed ubiquitously around the blood vessels throughout the body, especially around arteries. Therefore, we used the lipid droplet in adipose cells to monitor the blood vessels around the cells. In the experiments, a glass capillary (wall thickness: 4 \(\mu\)m; inner diameter: 45 \(\mu\)m) was employed to act as the blood vessel that was adhered to the surface of an adipose cell (see Fig. S9 of the Supplementary Information). The objective of the microscope was adjusted to be aligned with the centers of both the capillary and the lipid droplet (\(D = 20.0 \mu\)m) in the living cell. As the focal

![Fig. 3 Intracellular imaging of subcellular structures. a] Flourescence images of the microfilaments (F-actin filaments) of a living adipose cell. The weak fluorescent signals (a1) were efficiently enhanced by the lipid droplet (diameter: 7.7 \(\mu\)m) (a2). b) Normalized intensity distribution along the observation lines through microfilaments without (blue dashed in a1) and with (red dashed in a2) the lipid droplet. c) Fluorescence images of the microfilaments in a living adipose cell with a 12.5-\(\mu\)m lipid droplet. d) Fluorescence images of the microfilaments in a fixed adipose cell with an 18.6-\(\mu\)m lipid droplet. e) Fluorescence and bright-field images of the lysosomes in a living adipose cell with an 11.3-\(\mu\)m lipid droplet. f) Fluorescence and bright-field images of the adenoviruses in a living adipose cell with an 8.1-\(\mu\)m lipid droplet, the blue lines represent the movement trajectory of the adenoviruses in 10 min. The fluorescence images were focused on the surfaces of the cells (c1, d1, e1, and f1) and the virtual image planes (c2, d2, e2, and f2) formed by the lipid droplets. The bright-field image was focused on the virtual image plane formed by the lipid droplet (e3, f3). g) The microfilament movement monitored by the lipid droplet (diameter: 10.0 \(\mu\)m) inside the cell in 10 min.
length of the lipid droplet ($D = 20.0 \, \mu m$) is $\sim 40 \, \mu m$ according to the simulations, the detection range of the lipid droplet can cover the capillary in the radial direction, and the focusing effect of the lipid droplet is not affected by the presence of the capillary (see Fig. S10 of the Supplementary Information).

The excitation light was transmitted to the cell by the inverted objective and converged through the lipid droplet into the capillary (Fig. 5a1). The solution that contained blood cells and cancer cells (K562) flowed into the capillary. The mitochondria in the cancer cells were fluorescently labeled by Mito-Tracker (green) dyes as the targets of detection. The cancer cell flowing with the rightward fluid was gradually excited when it flowed through excitation light converged by the lipid droplet (Fig. 5a2, a3). Before reaching the effective region of the droplet, the fluorescence from the cancer cell was very weak (Fig. 5b1). As it arrived at the edge of the effective region, the fluorescence was enhanced a bit by the collection efficiency of droplet (Fig. 5b2). When the cancer cell reached the center of the effective region, the excitation light was highly concentrated, allowing us to obtain a much stronger fluorescent signal (Fig. 5b3, also see Video S2 of the Supplementary Information). As presented by the intensity mapping of the fluorescence from the cancer cell mitochondria (Fig. 5c), the signal strength of the fluorescence was greatly enhanced to improve the responsivity of the detection of cancer cells.

### Discussion

As organelles, endogenous lipid droplets can persist in the cytosol for a time that is dependent with the physiological state and the type of the cells. Generally, the mature lipid droplets with a larger size have a more stable morphology, which benefits the function as intracellular microlenses. However, due to the crowded intracellular environment, a higher optical power is required for the trapping laser of the OTS to manipulate the larger lipid droplets inside the cells (see Fig. S11 of the Supplementary Information). In our experiments, we can move the lipid droplets with a diameter of less than $8 \, \mu m$ inside the mature adipose cells. In the solution, the freely suspended lipid droplets trapped by the OTS can move in synchrony with the motion of the trap, as long as the trapping force on the lipid droplet was not exceeded by a drag force from the flow (see Section S1.4 and Fig. S12 of the Supplementary Information). Due to the much stronger drag force caused by the crowded intracellular environment, the movement of the trapped lipid droplets to the target position inside the cells becomes much slower ($0.001–5 \, \mu m \, s^{-1}$). Note that an arbitrary shift of intracellular structures may affect the functions of the organelles and the interactions between subcellular components. To investigate the effect on cells by the moving of the lipid droplets, an experiment was carried out to indicate the cell viability after trapping and moving of the lipid droplets in adipose cells (see Fig. S13 of the Supplementary Information). In each cell, more than 3 lipid droplets were moved over an average distance of 5 $\mu m$. Untreated cells were included as a control group. The cells were then continuously cultured for 24 h and the survival rates were recorded every 8 h. In addition, the cell viability after cultivation for 8, 16, and 24 h was judged by staining the cells with the cell permeable nucleus counterstain (Hoechst 33342). The results show that the survival rates of treated cells were very close to those of the control group. Besides, no apparent morphological changes were observed after moving the lipid droplets, and the treated cells kept growing and storing lipids after 24 h.

The lipid droplets can also be applied to different species of cells by endocytosis (see Fig. S14a–d of the Supplementary Information). The free lipid droplets were cocultured separately with the macrophage, tumor, epithelial and myocardial cells, and then internalized by the cells through phagocytosis and pinocytosis. The transfer efficiency in the four types of cells for 24 h was obtained (see Fig. S14e of the Supplementary Information). The internalization of the lipid droplets did not impair the cell proliferation (see Fig. S14f of the Supplementary Information). There have been some methods to induce the formation and accumulation of lipid droplets in other cell types (non-adipocytes), such as myocardial cells, etc.
monocytes, astrocytes, and epithelial cells. The accumulation of lipid droplets in these non-adipocytes was mainly induced in biochemical ways. Actually, lipid overload occurs in nearly all types of cells when the cells are in the abnormal metabolism. Apart from adipocytes, hepatic stellate cells and the cone photoreceptors of some species are also rich in long-term storage spherical lipid droplets with diameters larger than 1 μm that can be used as microlenses. It is also worth noting that there are some other bio-components that can work similarly to the lipid droplets. For example, chloroplasts and red blood cells (RBCs) can act as microlenses and also enable the amplification in imaging as well as the enhancement in fluorescent signal detection. These microlenses in biomaterials can be applied to some scenarios different with those of the lipid droplets. Chloroplasts, as a sort of botanic organelles, can be employed inside the plant cells, while RBCs in the blood vessels of animals may act as adaptive optofluidic microlenses due to their intrinsic elastic properties. Compared with chloroplasts and RBCs, the lipid droplets exhibit higher robustness and stability so that they can act as microlenses in different biological environments and can be internalized into cells of different types. Besides, with a higher refractive index and a nearly spherical shape, the lipid droplets provide a better performance in magnification and fluorescence enhancement (see Section S1.5 and Fig. S15 of the Supplementary Information).

The lensing effect of the lipid droplets is also valid in bulky biological tissues. We explored the application of the lipid droplets to porcine adipose tissue, and the surface morphology (bright-field) of the tissue was magnified by the lipid droplets (see Fig. S16 of the Supplementary Information), which implies the potential of the lipid droplets for in vivo operations. Moreover, there are some methods that may further improve the performance of lipid droplets by increasing the utilization efficiency of incident light. For example, some carotenoid pigments for refining the spectral sensitivity and improving color vision naturally exist in the cone lipid droplets within the photoreceptors of vertebrates. The refractive index of these lipid droplets can be increased up to 1.6 by binding lipid-soluble pigments (see Fig. S17 of the Supplementary Information), which can be utilized to increase the index contrast and to improve the imaging ability. In addition, we found that some lipid droplets in nearly spherical

![Fig. 5 Detecting extracellular signals.](image-url)
shapes could also act as microcavities to support resonances in whispering-gallery modes (WGMs). With an optical fiber probe next to a spherical stained lipid droplet inside an adipose cell, 532-nm laser light was coupled into the lipid microcavity to generate the WGM effects (see Fig. S18 of the Supplementary Information). Numerical investigations show that when WGM excitation occurs, the high-order scattering mode can form comparatively narrow pattern in the virtual image plane, and the transformation of the evanescent waves to propagating waves can be enhanced by the WGM, both of which are crucial effects in super-resolution imaging50. Therefore, generating the WGM effects by the lipid droplets is also a possible approach to the further enhancement of the imaging resolution by microspheres.

To summarize, we employed endogenous lipid droplets in mature adipose cells to act as the intracellular biomicrolenses for fluorescence imaging and signal enhancement. In the contact mode, the collection efficiency of the emitted fluorescence signals was improved, which sufficiently reduced the optical power required for excitation light in fluorescence imaging of subcellular structures within living cells. In the non-contact mode, the excitation light was highly converged by the lipid droplets to enhance the fluorescence from the extracellular environment surrounding the cells in real time. The presented lensing effect of the lipid droplets is expected to find applications to fully biocompatible miniaturized tools for biosensing, endoscopic analysis, and single-cell diagnosis for endoscopy. The use of the lipid droplets as intracellular microcavities also provides opportunities to construct diverse endogenous photonic devices.

**Materials and methods**

**Cell culture and staining**

Human preadipocytes-visceral (HPA-v) cells (ScienCell, USA) were seeded at a density of $1 \times 10^4$ cells cm$^{-2}$ on glass-bottom Petri dishes (35 mm, Biosharp, Anhui, China) and allowed to proliferate in preadipocyte medium for 36–48 h at 37 °C and 5% CO$_2$. The fully confluent cells were then induced to differentiate. The preadipocyte medium was replaced by the preadipocyte differentiation medium and the replacement was conducted every 2 days. The differentiation to mature adipose cells was completed after 6–12 days (see Fig. S19 of the Supplementary Information), and mature adipose cells can be maintained in adipocyte medium up to 7 days. The K562 cells (Pro-cell, Hubei, China) were seeded at a density of $1 \times 10^4$ cells cm$^{-2}$ in T25 flask and allowed to proliferate in the complete medium (IMDM + 1% penicillin-streptomycin + 10% fetal bovine serum) for 30–48 h at 37 °C and 5% CO$_2$. The K562 cells were passaged at constant density (1:2-1:4) every 2 days. The *Escherichia coli* (*E. coli*) expressing green fluorescent protein were grown in Luria-Bertani medium at 37 °C for 12 h. The *E. coli* were harvested and washed gently in phosphate-buffered saline solution (PBS). The *E. coli* were fixed by mixing 300 μl of cells with 700 μl of a 4% (wt−vol$^{-1}$) solution of paraformaldehyde in PBS and the mixture was incubated on ice for 5 min. The *E. coli* were then washed twice in PBS for removing the fixative. Samples of 30 μl were placed on a poly-L-lysine-coated slide and air dried for 2 h.

To stain the lipid droplets and the F-actin filaments, the mature adipose cells were washed twice with DPBS (Dulbecco’s phosphate buffer saline) and then fixed in paraformaldehyde for 15 min, followed by staining in Oil Red O (in 70% isopropanol; Shanghai Yuanye Bio-Technology, Shanghai, China) for 40 min and in PBS with Actin-Tracker (50 nM, Beyotime Biotechnology, Shanghai, China) for 30 min, respectively. After that, the cells were washed three times with 70% ethanol and DPBS to remove the excess dye. The nuclei, membrane, and microfilaments of living mature adipose cells were washed twice with DPBS were stained by PBS with Hoechst 33342 (5 μg ml$^{-1}$, Sigma-Aldrich, Germany), DIO (5 μM, Beyotime Biotechnology, Shanghai, China), and SiR-actin (0.5 μM, Cytoskeleton, USA) for 30 min, respectively (see Fig. S20 of the Supplementary Information). The K562 cells, which were suspended rather than adherent cells, were collected by centrifugation at 1000 r min$^{-1}$ for 5 min and then washed once with PBS to remove the complete medium. The cells were resuspended and the mitochondria were stained by PBS with Mito-Tracker (30 nM, Beyotime Biotechnology, Shanghai, China) for 30 min. After staining, all the staining agents were removed and replaced with phenol red-free complete medium.

The adenoviruses (VectorBuilder, Guangzhou, China) were incubated with amine-reactive Cy5 dyes (1 mg mL$^{-1}$ in DMSO, Shanghai Yuanye Bio-Technology, Shanghai, China) in a carbonate buffer (pH 9.3) at 25 °C for 2 h. The virus mixture was centrifuged at 15000-20000 g for 1-2 h, to remove the unbound Cy5 dyes. The viral aggregates were removed with 0.22 μm pore size filters before experiments. The adipose cells were incubated with the viruses at 37 °C and 5% CO$_2$ incubator for 30 min. Cells were washed with DPBS for 2 times before experiments51,52.

**Experimental setup**

The experimental setup consisted of an inverted fluorescence microscope (Eclipse Ti, Nikon) and a scanning optical tweezing system (OTS) (Tweez 250si, Aresis). The trapping laser (1064 nm) in the OTS was passed through an acousto-optic deflector (AOD) and a beam expander, and was refocused by the inverted objective onto the target. For bright-field imaging, a halogen light source (D-LH/LC:12 V, 100 W)
was used to illuminate the sample from the top, and the image was transmitted through the inverted objective (×60, DIC, WD = 2.0, water-immersion, Nikon) and received by a charge-coupled device (CCD) (MG-100, Mshot, Guangzhou, China). For fluorescence imaging, the excitation light from the LED was transmitted through the dichroic mirror and converged by the objective to illuminate the sample. The fluorescent signals were then collected by the objective and received by the CCD and a spectrometer (SHIS VNIR-520-20-S, Wayho Technology, Guangzhou, China). The filtrated bands for exciting the blue (Hoechst 33342), green (DiO, Mito-Tracker and Actin-Tracker), red (FNDS and Oil-red O), and deep red (SiR-actin) markers were 361–389 nm, 465–495 nm, 540–580 nm, and 590–650 nm, respectively.

Simulations
The electric field distributions of the lipid droplets were simulated with a finite element method (COMSOL Multiphysics, 5.5). The lipid droplets were set as isotropic microspheres. The refractive indices of the lipid droplets and surrounding (cytoplasm) were set as 1.52 and 1.36, respectively. Triangular meshes with a minimum element size of 1.5 mm and a maximum of 100 mm were set for the simulated region (200 μm × 200 μm) with scattering boundary conditions. To calculate the collection efficiency of the lipid droplets, a point source emitter (λ = 685 nm) was set in contact with the surface of the lipid droplet. The average Poynting vector flux was computed over a spherical surface (radius: 50 μm) above the droplet, with a center at the emitter. The collection angle over a spherical surface (radius: 50 μm) above the droplet was set as ± 47.3° in correspondence to the microscope objective NA (1.0). To numerically analyze the converging ability of the lipid droplet in the non-contact mode, a plane wave (width: 200 μm) at the wavelengths of 473 nm was set as the incident light.

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Author contributions
BL supervised the project. Y.Z., Y.L., P.F., and BL conceived and designed the research; X.C., T.W., and Z.G. performed the experiments; J.G. provided the experimental materials; Y.Z., Y.L., and P.F., and BL analyzed the data; X.C. and X.L. carried out the theoretical modeling and calculations; X.C., Y.Z., Y.L., P.F., and BL wrote and revised the paper. All authors discussed the results.

Conflict of interest
The authors declare no competing interests.

Supplementary information
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