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Intact living-cell electrolaunching ionization mass spectrometry for single-cell metabolomics†

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While single-cell mass spectrometry can reveal cellular heterogeneity and the molecular mechanisms of intracellular biochemical reactions, its application is limited by the insufficient detection sensitivity resulting from matrix interference and sample dilution. Herein, we propose an intact living-cell electrolaunching ionization mass spectrometry (ILCEI-MS) method. A capillary emitter with a narrow-bore, constant-inner-diameter ensures that the entire living cell enters the MS ion-transfer tube. Inlet ionization improves sample utilization, and no solvent is required, preventing sample dilution and matrix interference. Based on these features, the detection sensitivity is greatly improved, and the average signal-to-noise (S/N) ratio is about 20 : 1 of single-cell peaks in the TIC of ILCEI-MS. A high detection throughput of 51 cells per min was achieved by ILCEI-MS for the single-cell metabolic profiling of multiple cell lines, and 368 cellular metabolites were identified. Further, more than 4000 primary single cells digested from the fresh multi-organ tissues of mice were detected by ILCEI-MS, demonstrating its applicability and reliability.

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Introduction

Single-cell metabolite analysis is a promising method to reveal cellular heterogeneity at the most basic and subtle small molecule level,^{1–5} especially living single-cell metabolite analysis,⁶ enabling the gathering of the most veritable biochemical information and reflecting the cell status with higher fidelity, including cell differentiation and division,^{7–9} communication,^{1,10} interaction with the environment,^{11–13} and stress response.¹⁴ This diverse information is of great significance for the development of cell biology and medicine and facilitates cancer diagnosis, prognosis, and treatment.¹⁵ Mass spectrometry (MS) is the principal method for single-cell metabolite analysis because of its ability to afford rapid, wide-spectrum detection along with excellent qualitative structure determination.¹² Different MS-based methods have been developed to profile single-cell metabolites.^{16–20} Secondary-ion MS (SIMS)¹⁶ and matrix-assisted laser desorption/ionization MS (MALDI-MS)^{18,19} require a high-vacuum environment and complex sample preparation,

preventing the analysis of living cells. Cytometry by time of flight (CyTOF)²⁰ can be used to detect metal-labeled proteins in living cells; however, it is incapable of measuring small-molecule metabolites that are difficult to label. Therefore, electrospray ionization MS (ESI-MS), which does not require labeling, is more suitable for living single-cell metabolite analysis.^{11,21–26} In the detection of living cells, cell samples are usually intact cells or a part of the native components of living cells. Therefore, due to the complex biological structure of cells, the liquid system is no longer a homogeneous solution. However, current studies on single-cell ESI-MS still use the electrospray mode developed for homogeneous solutions, resulting in some insurmountable problems related to the ionization of heterogeneous single-cell samples. The use of a large amount of sheath fluid and sheath gas in the electrospray process results in the severe dilution of samples. The use of nanoliters of solvent for dissolution or extraction can result in thousand-fold sample dilution, which is lethal for trace components in single cells with picoliter volumes. In addition, because the ionization process occurs before the sample enters the MS ion inlet, the sample is subject to diffusion, dilution, and loss in the atmosphere; this results in problems such as a low sample-acquisition ratio in the MS ion inlet, thereby reducing the utilization of single-cell samples.²⁷ These problems limit the sensitivity of single-cell MS detection.

Although many bioelectrospray techniques based on high-throughput single-cell manipulation have been developed for tissue engineering and regenerative medicine,^{28–34} most are not

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suitable for MS. Therefore, to achieve high-sensitivity and high-throughput single-cell MS, a novel ionization technique that can be used for complex living-cell samples is needed.

Herein, we propose an intact living-cell electrolaunching ionization MS (ILCEI-MS) method using a narrow-bore capillary [the inner diameter (I.D.) is slightly smaller than the average cell diameter] to achieve efficient single-cell separation and transport while avoiding sample dilution by the sheath fluid. The novel ILCEI technique can realize undiluted living single-cell flow MS. In the ILCEI method, the constant-I.D., thin-walled tip of a quartz tube serves as the outlet, which reduces the volume of droplets formed by the combined action of the applied electric field and the surface tension on the port. Based on observation with an online ILCEI-MS visualization platform, the emitted single-cell droplet contains only a small amount of thin-layer buffer, wherein one cell acts as one droplet. The cell components are almost undiluted. Moreover, the cells remain intact and alive during flight. Ions are generated after a charged single-cell droplet forms in the heated transfer tube of the mass spectrometer, eliminating the loss of sample ions during transport through the atmospheric environment and mass spectrometer ion entrance, which is not possible with ESI. Therefore, the utilization of single-cell samples is significantly improved. Because of the abovementioned characteristics, the single-cell MS total ion chromatogram (TIC) of B104 cells obtained with ILCEI-MS has an average S/N ratio of $\sim 20 : 1$; thus, the detection sensitivity is greatly improved. Moreover, a throughput of approximately 51 cells per min was obtained for single-cell MS. Using the ILCEI-MS method, we analyzed more than 482 A549 cells and identified 368 cellular metabolites in one experiment. Additionally, more than 2800 *in vitro*-cultured living cells were discriminated with exact cell-type attribution. Hence, the ILCEI-MS method shows satisfactory single-cell detection sensitivity and stability. The high-throughput detection of 4072 single cells from multiple mouse organs in a tumor-bearing mouse model of non-small lung cancer (NSCLC) demonstrated the general applicability of this method in practical complex biological samples. ILCEI-MS is an easy-to-use method with a simple structure and stable operation. Moreover, ILCEI-MS is highly compatible with commercial MS ion source equipment and has low operating requirements.

Results and discussion

Configuration and visual characterization of the ILCEI-MS system

Based on our previous work on narrow-bore capillary chromatography,^{35–41} we developed a single-cell MS analysis method using a novel ILCEI technique. This analysis method aims to realize high-throughput single-cell online dispersion, dilution-free MS injection, and high-sensitivity MS detection. As shown in Fig. 1, the method is simple and requires only a narrow capillary and an in-house-built pressurized poly (methyl methacrylate) chamber. The capillary has a constant-I.D. throughout its entirety and a thin-walled tip. A freshly prepared cell suspension is loaded into a solution vial inside the pressurized chamber and introduced into the capillary driven by nitrogen

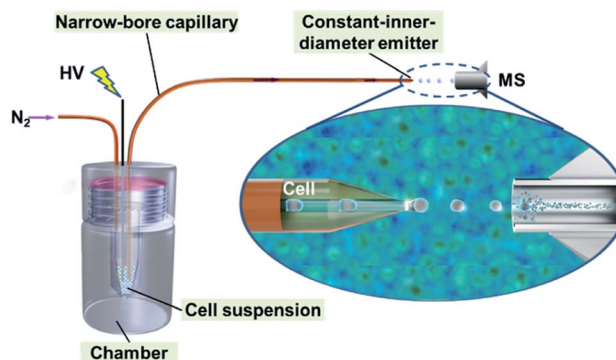


Fig. 1 Schematic illustration of the ILCEI-MS system.

gas for the injection of cells. The cells arrange themselves along the capillary axis to form a single-cell flow and are launched from the constant-I.D., thin-walled tip into the MS inlet for detection. A high voltage is applied directly *via* a stainless-steel needle to the cell suspension to launch single cells.

The necessity of each key strategy in the method was verified through a series of comparative experiments. A narrow capillary with an I.D. similar to the diameter of a cell is a prerequisite for separating single cells (Fig. S1†). A capillary wall thinned by etching is needed to generate a single-cell MS detection signal from a constant-I.D. capillary emitter (Fig. S2†). The tip with a constant-I.D. has a low risk of fluid clogging, enabling a stable flow rate and a long operation life, which are important factors for the MS detection of complex biological samples (Fig. S3†).

To objectively and comprehensively investigate the operation of the system, an online ILCEI-MS visualization platform was designed and built (Fig. S4†). The operation of the ILCEI-MS system was visually recorded using this platform. A single A549 cell moving in a narrow capillary propelled by a mobile phase was observed in real time and recorded at 10 000 frames per second (fps), as shown in Video S1.† The cell was slightly squeezed from its original spherical shape into a nearly cylindrical shape owing to the confinement effect as the cell passed through the narrow capillary (the I.D. was slightly smaller than

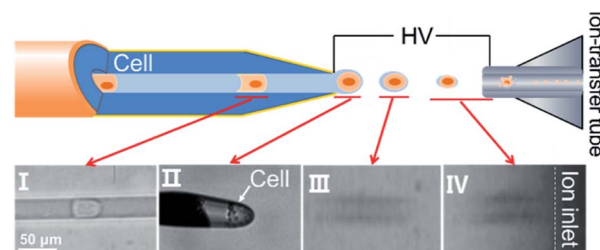


Fig. 2 Intact living-cell electrolaunching process in the ILCEI-MS system. Illustration (top) and corresponding high-speed screenshots (bottom) of different stages of the electrolaunching of a single cell: (I) motion of a single cell through a narrow capillary; (II) the cell reaches the front of the liquid cone of the emitter and is about to leave the emitter to form a single-cell droplet (arrows point to the true single cell); and (III) and (IV) the single-cell droplet flows in the atmosphere and is close to the MS ion-transfer tube inlet. HV: high voltage.



the cell diameter; Fig. 2, stage I). Additional video footage of a single cell being transported inside the capillary is provided in Video S2.† In this video, the single cell appears similar to a plug and is pushed along the capillary by the mobile phase. Once a single cell enters the capillary and begins its transport, its distance from the cell in front of it remains constant because the liquid is almost incompressible. Consequently, the single cells move in sequence. This observation is consistent with the expected behavior of cells in narrow-bore capillaries. Thus, the use of such capillaries is a simple and reliable method for single-cell dispersion and transport. Notably, even though the capillary was clogged with a few very large cells (due to cell heterogeneity), it could be flushed by applying a hydraulic pressure as low as 70–200 bar using a liquid chromatography system or simply a chromatography pump.

The ejection of an entire single cell from the constant-I.D.-emitter was recorded in real time at 50 000 fps (Video S3†). An intact individual cell was ejected from the capillary tip and carried a tiny amount of solvent to form the initial droplet that flew into the MS inlet. We termed this initial droplet the “single-cell droplet” (Fig. 2, stage II) and this process the “electrolaunching” (Fig. 2, stage II–IV). This unique launching phenomenon occurred at a certain voltage at the constant-I.D., thin-walled tip of the narrow capillary, which is key to resolving the dilution problem. We expected to achieve the same result when such a tip was used as an emitter. The results confirm that the ILCEI-MS method achieved single-cell injection rather than the injection of multiple cells simultaneously.

After synchronizing the high-speed microphotography with the single-cell MS data acquisition, three cells were successively electrolaunched and recorded using a high-speed camera at 50 000 fps (at this shooting speed, the camera could record up to 5 s); the corresponding mass spectra were collected simultaneously (Video S4†). The time interval of the electrolaunching of the three cells was consistent with the retention time of the MS peaks (Fig. S5 and Video S5†). This observation demonstrates that each signal peak in the TIC obtained by the ILCEI-MS system corresponds to a single cell; the peaks do not arise from the superposition of multiple cell signals. Further, we compared the ILCEI-MS results of the cell membrane debris sample and the intact living cell sample and found obvious differences in the MS peak shape and ion intensity between the two samples (Fig. S6†). Three samples were chosen to evaluate the signal stability of ILCEI-MS under complex solvent conditions: 40 mmol L⁻¹ ammonium formate aqueous solution (mobile phase); the cell lysate of lung cancer cells (the supernatant collected after centrifugation at 17 000 g to simulate the massive leakage of cellular components into the solution); and a liposome suspension with an average diameter of 200 nm (to simulate solutions containing a large number of cell membrane fragments). We collected 1000 scans, and the average relative standard deviation was less than 7%. The experimental results demonstrate that ILCEI-MS has good signal stability (Fig. S7 and S8†). The concern that cell debris or background noise may cause false-positive results in single-cell assays was ruled out. The visual characterization of the ILCEI-MS analysis system is shown in Fig. 2.

To further study the changes after the single-cell droplet was formed, we shifted the microscope observation field to the atmospheric section (Fig. 2, stage III) between the emitter and the MS inlet (Fig. 2, stage IV). The combined videos (Video S6†) show that the single-cell droplet did not undergo clear droplet splitting during its flight before it entered the ion-transfer tube. To further confirm that the cells remained intact before entering the MS inlet, we used a cover glass to obstruct and receive the single-cell droplet ejected by the emitter. Bright-field and fluorescence imaging revealed a distinct stereoscopic cell profile along with good single-cell dispersion and the absence of luminous cell debris, demonstrating that the cell entered the MS inlet in an intact state (Fig. S9†). To further verify that the ILCEI-MS system is a true living-cell MS detection system, we evaluated the viability of cells about to enter the mass spectrometer using CCK8 assay. The experimental details are provided in the Experimental section and Fig. S10.† Before entering the mass spectrometer, 87.22% of the cells were still alive, demonstrating that ILCEI-MS is truly a living-cell analysis method.

The formation of single-cell droplets eliminated the dilution of single-cell samples during electrolaunching. Most of the space in single-cell droplets is occupied by a cell, and the solvent is present only as an ultrathin film on the droplet surface, which reduces the dilution factor by hundreds or thousands of times compared to traditional methods. Furthermore, less solvent means less interference from impurities and less charge competition.⁴² The above two points are key to improving the sensitivity of single-cell metabolite analysis.

Formation mechanism of the single-cell droplet

We speculated that the formation of the single-cell droplet could be attributed primarily to the flow field and electric field created by the narrow-bore, constant-I.D., thin-walled emitter. The following experimental investigation was carried out based on this speculation.

Impact of the flow field. To evaluate the effect of the flow field in more detail, an emitter model was established, and the fluid dynamics of cell motion at the emitter outlet were numerically simulated (Fig. 3). For a constant-I.D. emitter under a constant flow rate (1 μL min⁻¹), a smaller capillary I.D. corresponds to a greater flow velocity of the mobile phase and a greater concentration of the high-velocity area immediately behind the cell at the emitter tip (Fig. 3a1–a3 and Table S1†). This means that the cells ejected from a narrower emitter are subjected to greater forward thrust and more easily reach the front end of the liquid cone. The cells shift downward under gravity after leaving the outlet, causing them to move away from the tip of the liquid cone. Decreasing the wall thickness of the emitter can diminish the liquid cone volume under an adequate flow rate. At this point, the cell must move to the cone tip because there is no space left for it to move near the side of the liquid cone (Fig. 3b1 and b2). As demonstrated by the above simulation and the data shown in Fig. 3c and d, when the volume of the liquid cone generated from the emitter is small, the cell easily comes in contact with the front side of the cone.



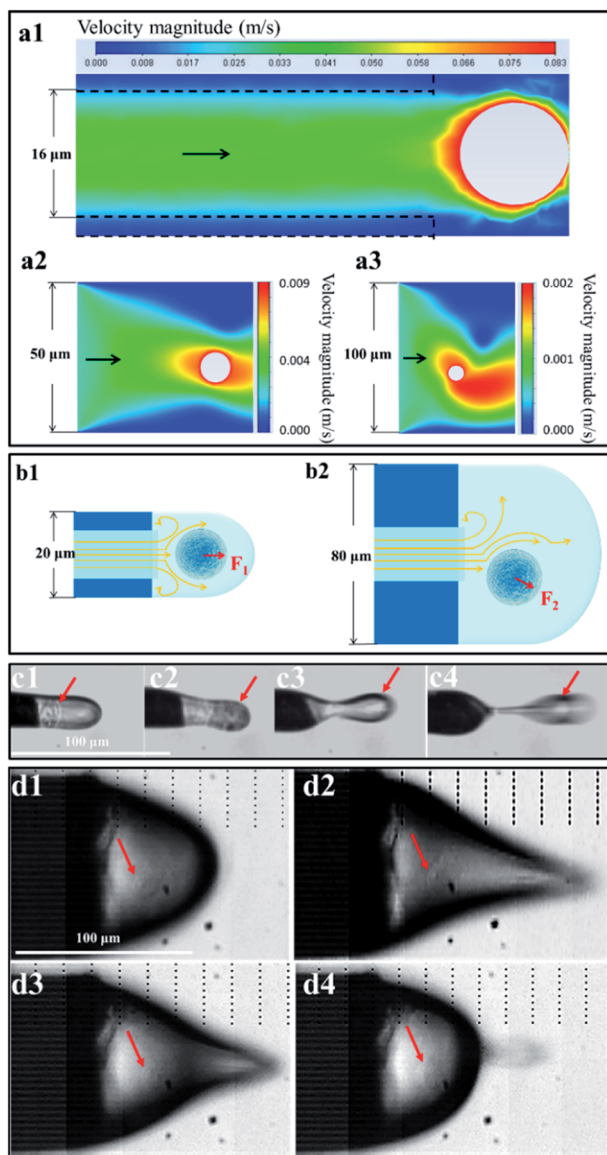


Fig. 3 Flow field generated by single-cell electrolaunching. (a) Velocity distribution diagram of the liquid surrounding a cell near the outlets of emitters with I.D.s of 16 μm (a1), 50 μm (a2), and 100 μm (a3). (b) Stress diagram of the liquid associated with a cell at the outlets of emitters with an outer diameter (O.D.) of 20 μm (b1) and 80 μm (b2). F_1 and F_2 are the directions of the resultant force on the cell. (c) and (d) High-speed screenshots of one launching circle when the emitter O.D. was 20 μm (c) and 80 μm (d). The red arrows marked in (c1)–(c4) and (d1)–(d4) represent the presence of a real single cell.

After reaching the front side of the cone, the cell, along with the mobile phase, is pulled forward by the force of the electric field. As the liquid cone is elongated (Fig. 3c3), the surface tension gradually increases and applies a force to only the backside of the cell because of the flexibility of the cells; this causes the liquid cone to contract inward until it is cut off (Fig. 3c4). At this time, a charged single-cell droplet (initial droplet) forms and is subjected to MS along the electric field line. Hence, from the perspective of fluid dynamics, to generate a single-cell droplet, a single cell must successfully reach the front side of the liquid

cone. A narrow-I.D. and thin emitter wall were conducive to satisfying this requirement. Moreover, the plugging and the high back pressure at the exit of the pulled conical emitter could be avoided by using a constant-I.D. emitter. This ensures the long-term stability of electrolaunching, which is important for high-throughput single-cell MS analysis (Fig. 3c and d, derived from Video S7†).

Impact of the electric field. To investigate the influence of the electric field on single-cell electrolaunching, the launching voltage was varied in the range of 1–2 kV. Two groups of MS detection experiments were carried out: one group with a KB cell suspension as the cell sample and a second group with adenine aqueous solution as a standard sample. All other experimental conditions were identical. The launching/spray currents at the corresponding voltages were recorded simultaneously for both groups of experiments (Fig. 4a).

To clearly characterize the experimental results, we defined the extracted ion chromatogram (EIC) as the total intensity of the extracted ion signals during a period (area integration of peaks in the extracted ion chromatogram) and the TIC as the intensity of the total ion signal during a period (integration of peaks in the TIC). The detection sensitivities of the samples in the same period were evaluated based on the EIC/TIC. The experimental results showed that the relationship between the voltage and the MS signal differed between the KB cell suspension and adenine aqueous solution, although the two samples had nearly overlapping current curves. The highest mass spectral signal intensity and sample detection sensitivity were obtained at 1.3–1.4 kV for the KB cell suspension; however, for the adenine aqueous solution, the mass spectrometry signal intensity began to increase when the spray voltage exceeded 1.4 kV, and the optimum signal was obtained at 1.6–1.8 kV. According to other studies on electrospray theory,^{43–47} the voltage required for an adenine aqueous solution spray system to form a steady jet should be 1.6–1.8 kV. The KB cell suspension had the same mobile phase as the adenine aqueous solution and produced almost the same ionization current; however, it produced a completely different mass spectrum response signal, demonstrating that ILCEI is different from ESI. This phenomenon is interesting as it reveals a disadvantage of the electrospray mode in the steady jet state: the conventional ESI technique was developed based on a homogeneous solution model and thus cannot handle the characteristics of heterogeneous systems such as cell suspensions. In contrast, the ILCEI technique was developed specifically for heterogeneous systems.

Preliminary study on the ionization mechanism of ILCEI-MS

Based on observation with a high-speed camera, the volume of the initial droplets produced by ILCEI remained almost constant before entering the mass spectrometer (Fig. 2; also see Video S6† for details). This is completely different from the continuous splitting of the initial droplets into secondary droplets in ESI. To further show the difference between ILCEI and ESI, we compared the cell suspension and homogeneous solution during mass spectrometry injection (Fig. S11 and



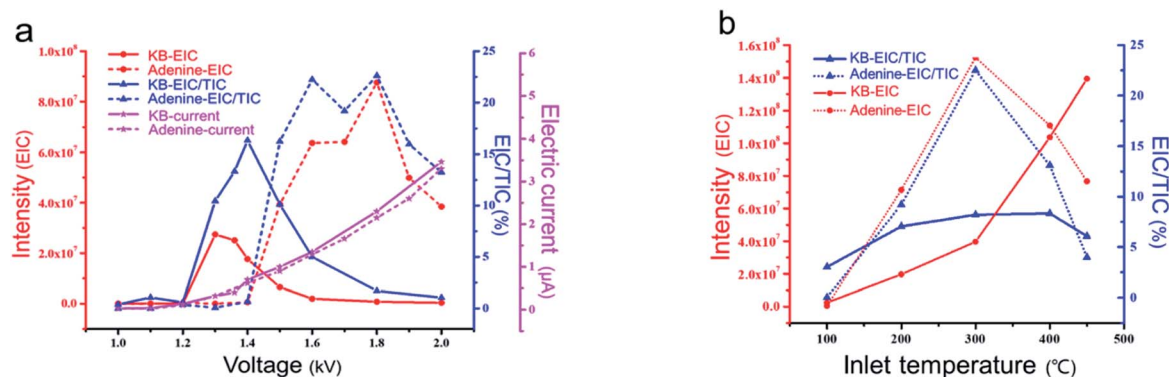


Fig. 4 Difference between single-cell electrolaunching ionization and electrospray ionization. (a) Influence of the launching/spray voltage (1–2 kV) on the single-cell/adenine MS signal intensity in the negative ion mode. (b) Influence of the temperature of the ion-transfer capillary entrance (100 °C, 200 °C, 300 °C, 400 °C, and 450 °C) on the single-cell/adenine MS signal intensity. KB-EIC: EIC of phosphatidylserine (PS 18 : 0/18 : 2; m/z 788.5446) in KB cells; adenine-EIC: EIC of adenine ($[M - H]^-$ m/z 134.0473); KB-TIC: total ion intensity from KB cells; adenine-TIC: total ion intensity of adenine; KB-current, adenine-current: the working electric current between the emitter and ion-transfer tube when detecting the KB cell suspension or adenine sample. Adenine was chosen because its MS behavior is similar to that of most polar metabolites in single cells.

Video S8†). The A549 cell suspension exhibited less droplet splitting (Fig. S11c2†) because of the presence of whole cells in the initial droplets of the cell suspension under electro-launching, which hindered droplet shrinkage. A large surface area was maintained, and the Rayleigh limit was not reached,⁴⁸ preventing the droplets from splitting. Because droplet splitting is accompanied by a random distribution of net charge, decreased droplet splitting results in more charge remaining on the surface of the mother droplet. In addition, a very small volume of the mobile phase means that very few impurity molecules are present in the mother liquid droplets. Thus, the cell components retained in the mother droplets will have a greater chance of gaining charge, thereby enhancing the ionization of the single-cell sample. The cells intercepted at the entrance of the mass spectrometer retained their structures and remained alive (see Fig. S9 and S10†). Thus, all cell components entered the mass spectrometer, avoiding sample loss in the atmospheric path and improving the sample collection rate. Hence, we believe that the ionization of single-cell components should occur in the ion-transfer tube of the mass spectrometer. Referring to similar studies on “inlet ionization”,^{27,49} we propose the following hypotheses: (1) after entering the MS inlet, the single-cell droplet experiences rapid boiling and explosion in a low-pressure, high-temperature environment; and (2) the sample molecules rapidly move to the gas phase and acquire random charges during the explosion. According to these assumptions, the proportion of sample molecules converted to gas-phase ions will increase as the intensity of the explosion increases.

The ionization processes that occurred after a single-cell droplet entered the ion-transfer tube could not be characterized visually using a high-speed camera. Therefore, we conducted a preliminary experimental verification. We increased the temperature of the ion-transfer tube from 100 °C to 450 °C in stages (the normal adjustable parameter range of a mass spectrometer) and recorded the changes in the MS signal intensity (EIC) and the MS sensitivity (EIC/TIC) for a KB cell

suspension and a 14 $\mu\text{mol L}^{-1}$ adenine standard solution (Fig. 4b). For both samples (*i.e.*, the KB cell suspension and the adenine standard solution), the EIC/TIC value gradually increased with increasing temperature below 300 °C; when the temperature exceeded 300 °C, the EIC/TIC value of the KB cell suspension continued to increase, whereas that of the adenine standard solution gradually decreased. Below 300 °C, the signal intensity of both samples increased with increasing temperature; this is because a reasonable increase in the temperature of the ion-transfer tube promoted the desolation of the sample and thus increased the intensity of the signal arising from the sample, regardless of whether ionization occurred in or out of the MS inlet. For a homogeneous solution, it is easy to reach the temperature required for complete evaporation. When the temperature continued to increase, the signal intensity decreased under the combined effects of background ion interference and the space-charge effect. However, single-cell droplets with larger volumes had more complex constituents than the initial droplets produced by homogeneous solution electrospray; thus, higher temperatures were required to complete the desolvation process quickly. These results corroborate our hypothesis that the cells are rapidly gasified in the ion-transfer tube to achieve ionization. As shown in Fig. 4b, the single-cell MS signal intensity obtained with our system can be further improved by increasing the temperature of the mass spectrometer ion-transfer tube to ≥ 400 °C. The main feature of inlet ionization is that it avoids the diffusion and annihilation of sample ions in the atmosphere, which greatly improves the detection sensitivity and sample utilization.

Performance evaluation of ILCEI-MS for single-cell detection

The detection throughput and sensitivity of the ILCEI-MS system were tested using B104 cells as an example. A throughput of approximately 51 cells per min was obtained for single-cell MS (Fig. 5a), and the average S/N ratio of single-cell signal peaks in TIC (baseline intensity = 1×10^6) was



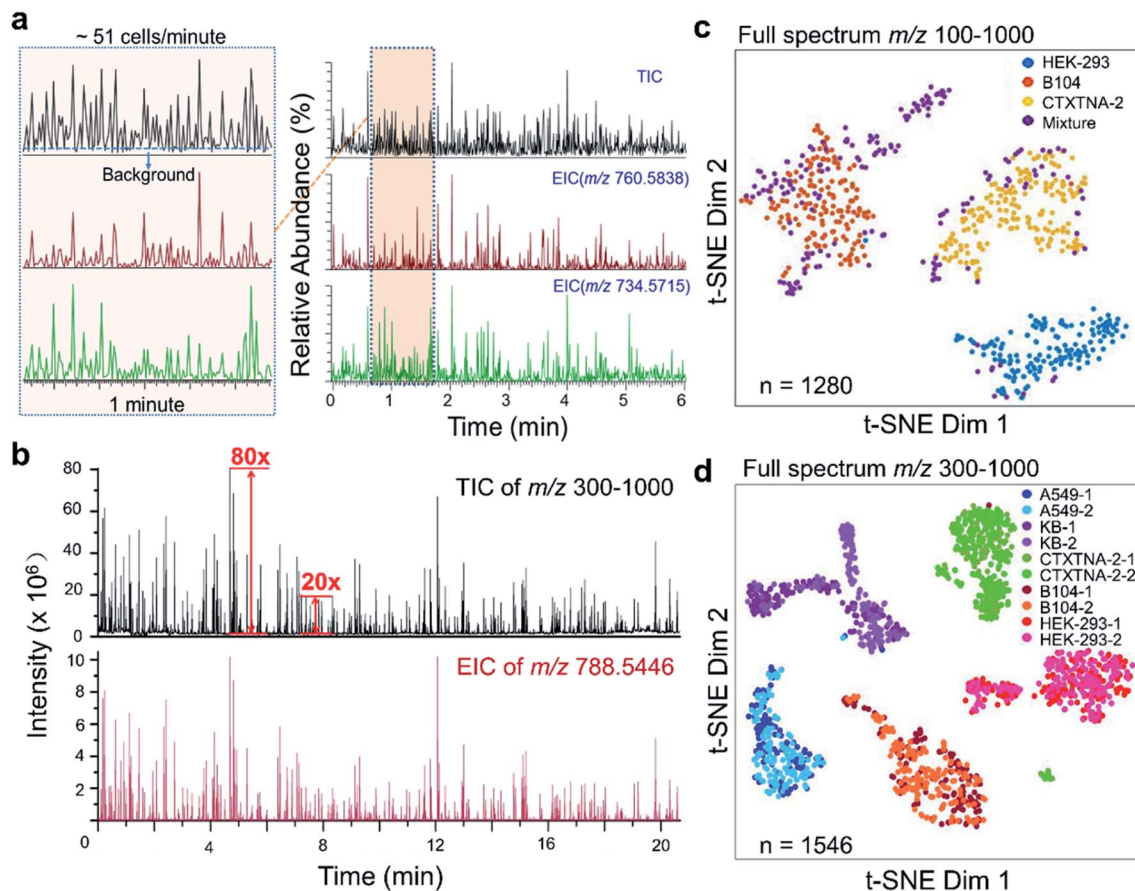


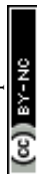
Fig. 5 Performance of the ILCEI-MS method for single-cell detection. (a) The detection throughput of A549 cells by ILCEI-MS reached approximately 51 cells/min. (b) TIC of B104 cells (black line) in the scan range of m/z 300–1000, and the EIC of phosphatidylserine (PS 18 : 0 / 18 : 2; m/z 788.5446) (red line). The average S/N ratio of the single-cell signal peaks in TIC (baseline intensity = 1×10^6) was approximately 20 : 1, and the highest S/N ratio was 80 : 1. (c) The t-distributed stochastic neighbor embedding (t-SNE) map of HEK-293, B104, and CTXTNA-2 cells and a mixture of these three cell types. (d) The t-SNE map of A549, KB, CTXTNA-2, B104, and HEK-293 cells based on their mass spectra measured at the different time periods (“-1” indicates measurement in the a.m., and “-2” indicates measurement in the p.m.). “n” is the number of single cells collected.

approximately 20 : 1, and the highest S/N ratio reached 80 : 1 in the scan range of m/z 300–1000 (Fig. 5b). We analyzed over 482 A549 cells and obtained approximately 5200 peaks in the m/z range of 100–1000. Among them, 368 cellular metabolites were assigned (Table S2†). On average, 800 ions and 745 ions (Fig. S12†) related to cellular metabolites in the positive and negative ion modes were detected from living single cells, respectively. The expanded coverage of single-cell metabolites demonstrates that ILCEI-MS has great potential for in-depth studies of cellular heterogeneity at the metabolic level. We also evaluated the cell discrimination capability and repeatability of ILCEI-MS. First, MS analysis was performed on HEK-293, B104, and CTXTNA-2 cells and a mixture of these three cell types. Fig. 5c shows that these three cell types were clearly discriminated, and the cell mixture was well resolved into these three cell types. Next, A549, KB, CTXTNA-2, B104, and HEK-293 cells were repeatedly detected by the ILCEI-MS system at different times of one day to evaluate the repeatability (Fig. 5d); the single-cell mass spectra of these five types of cell are shown in Fig. S13.† The cells of the same type were clustered together

in the t-distributed stochastic neighbor embedding (t-SNE) map, demonstrating the good repeatability and weak batch effect of the ILCEI-MS system.

Primary single-cell analysis

The cell suspension obtained from the digestion of fresh tissue is usually a mixture of different types of cell, such as epithelial cells, glial cells, and muscle cells. These cells are quite different in size, structure, physicochemical characteristics, *etc.*, presenting challenges in terms of the universal applicability of analytical methods. Therefore, we further studied the metabolic profiles of primary cells digested from fresh tissue. A549 cells were used to establish a tumor-bearing mouse model of NSCLC. As shown in Fig. S14,† the lung mass in the lung cancer mouse was obvious, and the liver was enlarged and hard, indicating successful cancer cell transplantation. ILCEI-MS was used successfully to detect 4072 single cells from the single-cell suspension (Fig. S15 and S16†) digested from tissues of hearts, livers and lungs. The original MS data are shown in Fig. S17–S22.† On average, 773 ions and 787 ions (Fig. S23†)



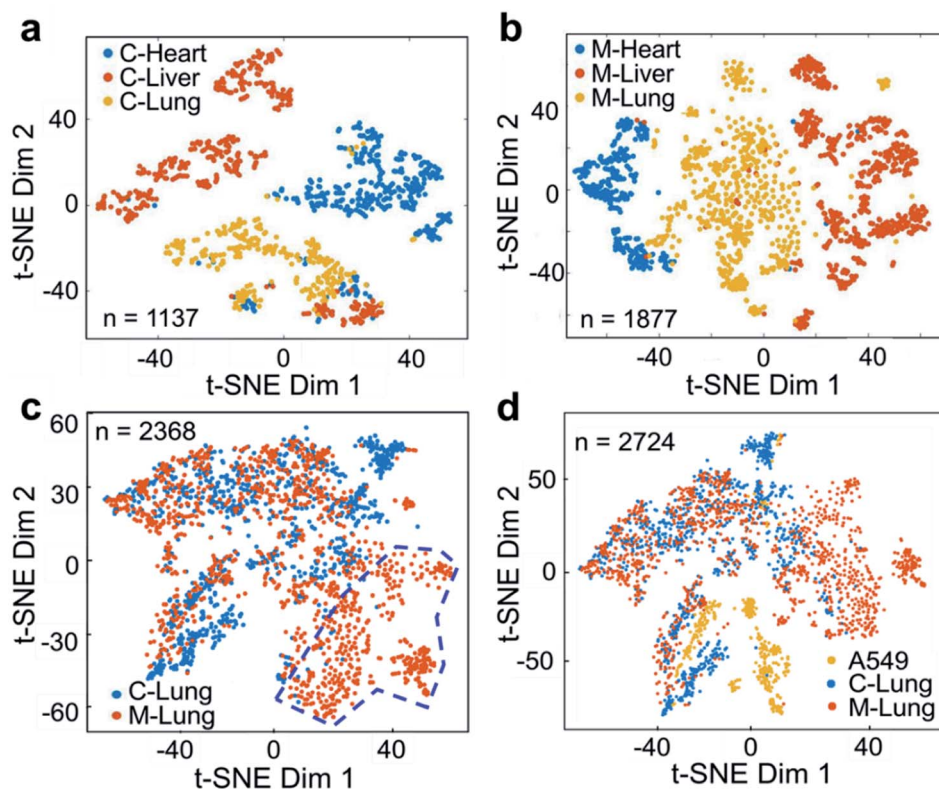


Fig. 6 Performance of the ILCEI-MS method for the detection of single cells obtained from the digestion of mouse tissues. (a) and (b) The t-SNE maps of single cells obtained from the heart, liver, and lung tissues of control mice ("C") and lung cancer model mice ("M"). (c) The t-SNE map of single cells obtained from digested lung tissues of control mice and lung cancer model mice. The blue dashed circle marks the specific cell subset of lung tissue cells in lung cancer model mice. (d) The t-SNE map of single cells obtained from the lung tissues of control mice and lung cancer model mice and cultured A549 cells *in vitro*. "n" is the number of single cells collected.

related to cellular metabolites were detected in the positive ion mode from 2368 individual single cells from lung tissues of the lung cancer model mice and control mice, respectively. To visualize the heterogeneity of cells in multiple organs, we performed a t-SNE analysis based on the single-cell metabolite profiles acquired by ILCEI-MS. Fig. 6a and b show that the differences in the heart, liver, and lung cells of the lung cancer model mice were more apparent than those in the control mice. Remarkably, the lung tissue cells from the lung cancer model mice exhibited greater metabolic differences (Fig. 6b), indicating more cell subpopulations, consistent with the literature.⁵⁰ Further comparison of the single cells from the lung tissues of the lung cancer model mice and control mice showed partial overlap and partial difference in the cell community. Interestingly, a specific subset of cells (marked by the blue dashed circle in Fig. 6c) appeared in the lung tissue cells of the lung cancer model mice. The metabolomic information contained in this cell subset may provide valuable information for further studies on biomarkers for NSCLC. The primary lung cells in the control and lung cancer model mice were clearly distinguished from cultured A549 cells *in vitro*, which were used to establish a lung cancer model (Fig. 6d). These results suggest that the human lung cancer cell line A549 underwent metabolic reprogramming after entering the mice to adapt to the micro-environment *in vivo* and meet the needs of their own growth.⁵¹

These results effectively demonstrate the reliability and universal applicability of the ILCEI-MS method.

Conclusion

Single-cell sequencing has contributed to great progress in the fields of single-cell genomics⁵² and transcriptomics.⁵³ However, metabolites cannot be amplified like DNA/RNA. Therefore, the development of single-cell metabolomics relies on high-sensitivity detection methods. At present, single-cell metabolite MS methods suffer from several problems. First, the detection sensitivity and detection coverage of existing methods are insufficient, restricting the investigation of cell heterogeneity. Second, many single-cell MS detection methods involve complicated systems, require difficult operation procedures, and have high manufacturing costs and poor stability. These problems arise because sample dilution, waste, and matrix interference in the electrospray process have not been well resolved. In addition, these problems become more prominent when heterogeneous cell samples (*e.g.*, living single cells) are ionized.

In this study, we developed a new ionization method that continuously introduces live single cells into a mass spectrometer. With a cell detection throughput of 51 cells per min, ILCEI-MS achieved high sensitivity for single-cell detection, and the average S/N ratio of single-cell signal peaks is about 20 : 1 in the



TIC. The average information coverage of a single cell is greater than 700 metabolite-related ions. These promising results can be attributed to the following two characteristics: first, the narrow-bore capillary exerts a significant spatial confinement effect on the cell, resulting in reliable single-cell high-throughput separation along with orderly long-distance cell transport while avoiding the dilution of cell samples by sheath solution and sheath gas. Second, a novel intact living-cell electrolaunching ionization process was developed based on a constant-I.D., thin-walled emitter at the end of a narrow capillary. This approach can realize single-cell MS injection with almost no mobile phase dilution, little matrix interference, and inlet ionization, resulting in greatly improved single-cell sample utilization. A series of single-cell detection experiments with a high sampling volume were tested, demonstrating that the ILCEI-MS system has good single-cell recognition ability and stability. The universal applicability of ILCEI-MS for actual biological samples was demonstrated by detecting 4072 primary single cells digested from fresh mouse heart, liver, and lung tissues.

Although the mechanism of the ILCEI-MS requires further study, it does not hinder its application in fields closely related to cell heterogeneity. For example, ILCEI-MS can be used in developmental biology to study the metabolic molecular mechanism of cell differentiation in the early embryo, related research has emerged in the field of genomics.⁵⁴ We expect that this method can also be applied to high-throughput single-organelle metabolite analysis in, for example, the nucleus and mitochondria.

Data availability

The data supporting the findings of this study are available within the article and in the ESI.† The raw data are available from the authors upon reasonable request.

Author contributions

X. Y. W. supervised the project. X. Y. W. and Y. L. S. designed the experiments. Y. L. S. designed and modified the instrument under the guidance of X. Y. W. and Yaoyao Z. Q. Z., Y. L. S., and Yingyan Z. performed the high-speed photography and the corresponding data analysis. Y. L. S., W. M. Z., Yaoyao Z., Y. X. L., and G. Z. Z. prepared the materials and performed the MS detection. Yaoyao Z., G. S. G., S. C. Z., Y. X. L., X. R. Z., W. M. Z., Y. L. S., H. Y., and H. S. Z. performed the data analysis. Y. L. S. designed and performed the simulations under the guidance of X. Y. W. The manuscript was written by Y. L. S., Yaoyao Z., Yingyan Z., W. M. Z., and X. Y. W. All authors discussed the results and commented on the manuscript. All authors approved the final manuscript.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 F. J. Hartmann, D. Mrdjen, E. McCaffrey, D. R. Glass, N. F. Greenwald, A. Bharadwaj, Z. Khair, S. G. S. Verberk, A. Baranski, R. Baskar, W. Graf, D. Van Valen, J. Van den Bossche, M. Angelo and S. C. Bendall, *Nat. Biotechnol.*, 2021, **39**, 186–197.
- 2 M. Fessenden, *Nature*, 2016, **540**, 153–155.
- 3 V. Pareek, H. Tian, N. Winograd, S. J. Benkovic and H. Tian, *Science*, 2020, **368**, 283–290.
- 4 Q. S. Huang, S. F. Mao, M. Khan, W. W. Li, Q. Zhang and J. M. Lin, *Chem. Sci.*, 2020, **11**, 253–256.
- 5 C. Seydel, *Nat. Methods*, 2021, **18**, 1452–1456.
- 6 M. H. Zhuang, Z. H. Hou, P. Y. Chen, G. L. Liang and G. M. Huang, *Chem. Sci.*, 2020, **11**, 7308–7312.
- 7 X. Xu, J. Wang, L. Wu, J. Guo, Y. Song, T. Tian, W. Wang, Z. Zhu and C. Yang, *Small*, 2020, **16**, e1903905.
- 8 M. Oginuma, Y. Harima, O. A. Tarazona, M. Diaz-Cuadros, A. Michaut, T. Ishitani, F. Xiong and O. Pourquie, *Nature*, 2020, **584**, 98–101.
- 9 S. Trefely and K. E. Wellen, *Science*, 2018, **360**, 603–604.
- 10 M. N. Artyomov and J. Van den Bossche, *Cell Metab.*, 2020, **32**, 710–725.
- 11 H. Zhu, N. Wang, L. Yao, Q. Chen, R. Zhang, J. Qian, Y. Hou, W. Guo, S. Fan, S. Liu, Q. Zhao, F. Du, X. Zuo, Y. Guo, Y. Xu, J. Li, T. Xue, K. Zhong, X. Song, G. Huang and W. Xiong, *Cell*, 2018, **173**, 1716–1727.
- 12 L. Zhang and A. Vertes, *Angew. Chem., Int. Ed.*, 2018, **57**, 4466–4477.
- 13 M. Xue, W. Wei, Y. Su, J. Kim, Y. S. Shin, W. X. Mai, D. A. Nathanson and J. R. Heath, *J. Am. Chem. Soc.*, 2015, **137**, 4066–4069.
- 14 P. J. Hines, *Science*, 2016, **354**, 843.
- 15 R. Zenobi, *Science*, 2013, **342**, 1243259.
- 16 Z. Y. Yuan, Q. M. Zhou, L. S. Cai, L. Pan, W. L. Sun, S. W. Qumu, S. Yu, J. X. Feng, H. S. Zhao, Y. C. Zheng, M. L. Shi, S. Li, Y. Chen, X. R. Zhang and M. Q. Zhang, *Nat. Methods*, 2021, **18**, 1223–1232.
- 17 L. Rappez, M. Stadler, S. Triana, R. M. Gathungu, K. Ovchinnikova, P. Phapale, M. Heikenwalder and T. Alexandrov, *Nat. Methods*, 2021, **18**, 799–805.
- 18 K. Scupakova, F. Dewez, A. K. Walch, R. M. A. Heeren and B. Balluff, *Angew. Chem., Int. Ed.*, 2020, **59**, 17447–17450.
- 19 E. K. Neumann, T. J. Comi, S. S. Rubakhin and J. V. Sweedler, *Angew. Chem., Int. Ed.*, 2019, **58**, 5910–5914.



- 20 Q. L. Liu, W. J. Ge, T. T. Wang, J. Y. Lan, S. Martinez-Jarquin, C. Wolfrum, M. Stoffel and R. Zenobi, *Angew. Chem., Int. Ed.*, 2021, **60**, 24534–24542.
- 21 W. F. Zhang, N. Li, L. Lin, Q. S. Huang, K. Uchiyama and J. M. Lin, *Small*, 2020, **16**, 1903402.
- 22 H. Yao, H. S. Zhao, X. Zhao, X. Y. Pan, J. X. Feng, F. J. Xu, S. C. Zhang and X. R. Zhang, *Anal. Chem.*, 2019, **91**, 9777–9783.
- 23 J. F. Cahill, J. Riba and V. Kertesz, *Anal. Chem.*, 2019, **91**, 6118–6126.
- 24 S. Xu, M. Liu, Y. Bai and H. Liu, *Angew. Chem., Int. Ed.*, 2021, **60**, 1806–1812.
- 25 Z. Z. Shen, H. S. Zhao, H. Yao, X. Y. Pan, J. L. Yang, S. C. Zhang, G. J. Han and X. R. Zhang, *Chem. Sci.*, 2022, **13**, 1641–1647.
- 26 L. Song, K. Chinglin, M. Wang, D. Zhong, H. Chen and J. Xu, *Anal. Chem.*, 2022, **94**, 4175–4182.
- 27 V. S. Pagnotti, N. D. Chubaty and C. N. McEwen, *Anal. Chem.*, 2011, **83**, 3981–3985.
- 28 S. N. Jayasinghe, P. A. M. Eagles and A. N. Qureshi, *Biotechnol. J.*, 2006, **1**, 86–94.
- 29 S. N. Jayasinghe, A. N. Qureshi and P. A. M. Eagles, *Small*, 2006, **2**, 216–219.
- 30 P. A. M. Eagles, A. N. Qureshi and S. N. Jayasinghe, *J. Biol. Chem.*, 2006, **394**, 375–378.
- 31 P. K. Odenwalder, S. Irvine, J. R. McEwan and S. N. Jayasinghe, *Biotechnol. J.*, 2007, **2**, 622–630.
- 32 S. Sahoo, W. C. Lee, J. C. H. Goh and S. L. Toh, *Biotechnol. Bioeng.*, 2010, **106**, 690–698.
- 33 D. I. Braghirolli, F. Zamboni, P. C. Chagastelles, D. J. Moura, J. Saffi, J. A. P. Henriques, D. A. Pilger and P. Pranke, *Biomicrofluidics*, 2013, **7**, 044130.
- 34 H. J. Chun, C. H. Park, I. K. Kwon, G. Khang, N. Maurmann, L. E. Sperling and P. Pranke, in *Cutting-edge enabling technologies for regenerative medicine*, Springer Press, Singapore, 2018, vol. 1078, pp. 79–100.
- 35 Y. L. Shao, Y. Y. Zhou, Y. Y. Wu, Q. Zhang, Z. H. Yu, G. S. Guo and X. Y. Wang, *Talanta*, 2020, **218**, 121096.
- 36 X. Y. Wang, V. Veerappan, C. Cheng, X. Jiang, R. D. Allen, P. K. Dasgupta and S. R. Liu, *J. Am. Chem. Soc.*, 2010, **132**, 40–41.
- 37 X. Y. Wang, L. Liu, Q. S. Pu, Z. F. Zhu, G. S. Guo, H. Zhong and S. R. Liu, *J. Am. Chem. Soc.*, 2012, **134**, 7400–7405.
- 38 L. Liu, V. Veerappan, Q. R. Pu, C. Cheng, X. Y. Wang, L. P. Lu, R. D. Allen and G. S. Guo, *Anal. Chem.*, 2014, **86**, 729–736.
- 39 L. Liu, V. Veerappan, Y. Z. Bian, G. S. Guo and X. Y. Wang, *Sci. China: Chem.*, 2015, **58**, 1605–1611.
- 40 R. N. Li, Y. L. Shao, Y. M. Yu, X. Y. Wang and G. S. Guo, *Chem. Commun.*, 2017, **53**, 4104–4107.
- 41 W. M. Zhang, L. Liu, Q. Zhang, D. T. Zhang, Q. Hu, Y. N. Wang, X. Y. Wang, Q. S. Pu and G. S. Guo, *Chem. Commun.*, 2020, **56**, 2423–2426.
- 42 A. C. Susa, Z. J. Xia and E. R. Williams, *Anal. Chem.*, 2017, **89**, 3116–3122.
- 43 G. I. Taylor, *Proc. R. Soc. London, Ser. A*, 1966, **291**, 145–158.
- 44 M. Cloupeau and B. Prunet-Foch, *J. Aerosol Sci.*, 1994, **25**, 10211036.
- 45 I. Hayati, A. I. Bailey and T. F. Tadros, *Nature*, 1986, **319**, 41–43.
- 46 A. Jaworek and A. Krupa, *J. Aerosol Sci.*, 1999, **30**, 873–893.
- 47 P. Nemes, I. Marginean and A. Vertes, *Anal. Chem.*, 2007, **79**, 3105–3116.
- 48 L. Rayleigh, *Philos. Mag.*, 1882, **14**, 184–186.
- 49 C. N. McEwen, V. S. Pagnotti, E. D. Inutan and S. Trimpin, *Anal. Chem.*, 2010, **82**, 9164–9168.
- 50 A. Prat, O. Karginova, J. S. Parker, C. Fan, X. P. He, L. Bixby, J. C. Harrell, E. Roman, B. Adamo, M. Troester and C. M. Perou, *Breast Cancer Res. Treat.*, 2013, **142**, 237–255.
- 51 D. F. Quail and J. A. Joyce, *Nat. Med.*, 2013, **19**, 1423–1437.
- 52 C. A. Lareau, L. S. Ludwig, C. Muus, S. H. Gohil, T. Zhao, Z. Chiang, K. Pelka, J. M. Verboon, W. Luo, E. Christian, D. Rosebrock, G. Getz, G. M. Boland, F. Chen, J. D. Buenrostro, N. Hacohen, C. J. Wu, M. J. Aryee, A. Regev and V. G. Sankaran, *Nat. Biotechnol.*, 2021, **39**, 451–461.
- 53 A. H. Rizvi, P. G. Camara, E. K. Kandror, T. J. Roberts, I. Schieren, T. Maniatis and R. Rabadan, *Nat. Biotechnol.*, 2017, **35**, 551–560.
- 54 G. Peng, S. Suo, G. Cui, F. Yu, R. Wang, J. Chen, S. Chen, Z. Liu, G. Chen, Y. Qian, P. P. L. Tam, J. D. J. Han and N. Jing, *Nature*, 2019, **572**, 528–532.

