Microfluidic cell sorter-aided directed evolution of a protein-based calcium ion indicator with an inverted fluorescent response†

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We demonstrate a simple, low cost and disposable microfluidic fluorescence activated cell sorting system (µFACS) for directed evolution of fluorescent proteins (FP) and FP-based calcium ion (Ca2+) indicators. The system was employed to pre-screen libraries of up to 10⁶ variants of a yellow FP-based Ca2+ indicator (Y-GECO) with throughput up to 300 cells per s. Compared to traditional manual screening of FP libraries, this system accelerated the discovery of improved variants and saved considerable time and effort during the directed evolution of Y-GECO. Y-GECO1, the final product of the µFACS-aided directed evolution, has a unique fluorescence hue that places it in the middle of the spectral gap that separates the currently available green and orange FP-based Ca2+ indicators, exhibits bright fluorescence in the resting (Ca2+ free) state, and gives a large response to intracellular Ca2+ fluctuations in live cells.

Introduction

The calcium ion (Ca2+) has a pivotal role in cell signaling, serving as the critical second messenger for numerous signaling pathways and processes in essentially all cell types. The Ca2+ signaling system is highly versatile and operates with a diverse array of spatial and temporal patterns in different tissues.1,2 To interrogate Ca2+ dynamics, it is essential to develop technologies that enable visualization and quantitative measurement of intracellular Ca2+ signals. Compared to synthetic dye-based Ca2+ indicators, fluorescent protein (FP)-based Ca2+ indicators offer a number of advantages for Ca2+ imaging in live cells and tissues. Specifically, FP-based Ca2+ indicators are genetically encoded and can therefore be delivered to cells transiently in the minimally invasive form of the corresponding gene or constitutively expressed in transgenic animals.3 Furthermore, with appropriate gene promoters and trafficking signals, FP-based Ca2+ indicators can be expressed only in specific cell types or targeted to specific subcellular compartments.

Extensive protein engineering efforts have led to an expansion of the toolbox of FP-based Ca2+ indicators and introduced new members with improved or novel properties.4–8 One effective approach for developing improved FP-based Ca2+ indicators is the manual one-by-one testing of a small selection of variants rationally designed on the basis of structure-based hypotheses regarding the mechanism.5,9–11 Arguably, a more effective approach for discovering improved FP-based Ca2+ indicators is the use of directed evolution, where large libraries of randomly generated variants are subjected to a high-throughput screen to identify improved variants.6–8 Such directed evolution...
efforts have typically relied on manual on-plate screening of bacterial colony fluorescence with moderate throughput of up to $10^4$ variants every few days. In this way rare clones with improved or unanticipated novel properties, that would be unlikely to be identified by rational approaches, can be discovered.

A promising approach for increasing the size of FP-based libraries that can be screened, and thereby increasing the odds of finding improved variants, is to exploit the ability of fluorescently activated cell sorting (FACS) to rapidly evaluate and sort very large numbers of fluorescently labeled cells. Relative to the labor-intensive and time-consuming process of on-plate screening, the throughput of FACS is 3 orders of magnitude higher. Unfortunately, FACS equipment is expensive, requires a high level of technical skill to operate, and is typically insensitive for screening properties other than brightness, such as analyte-dependent fluorescent changes or photoconversion. Microfluidic FACS ($\mu$FACS) devices are a promising alternative to traditional FACS for screening FP and FP-based Ca$^{2+}$ indicators, as they are inexpensive and much more amenable to customization. Recently, a number of $\mu$FACS devices have been reported for screening fluorescently-labeled cells, which could potentially be customized for molecular evolution of FP and FP-based Ca$^{2+}$ indicators. Despite the potential of $\mu$FACS, application of $\mu$FACS devices for evolving useful FP indicators remains elusive. To achieve this goal, we have built a simple, cost-effective and disposable high-throughput $\mu$FACS device integrated with a piezoelectric actuator based on previous reports. We use this device for library enrichment prior to a manual screen, to accelerate the directed evolution of a single FP-based Ca$^{2+}$ indicator derived from the yellow FP mPapaya. Compared to other sorting mechanisms such as electroosmotic, dielectrophoretic, hydrodynamic sorting and optical switching, the integrated piezoelectric actuator offers several advantages including low cost ($<$ $100), low power consumption ($<$ 0.1 mW), precise control, fast response time (0.1–1 ms) and high cell viability. In addition, the $\mu$FACS device costs less than $10$ and is disposable, which eliminates the possibility of cross-contamination from bacterial samples. In contrast, FACS often requires harsh and complex cleaning procedures for de-contamination of bacterial samples. These features of our $\mu$FACS device are preferable for live cell screening.

The design of the single FP-based Ca$^{2+}$ indicator engineered in this work is similar to that of pericam and GCaMP. In this design, the C- and N-termini of a circularly permuted FP (cpFP) are fused to calmodulin (CaM) and a peptide binding domain that modulates its fluorescent properties. The large conformational change associated with binding of CaM to Ca$^{2+}$ followed by binding of Rsf20 to Ca$^{2+}$-bound CaM, results in a change in the FP chromophore environment that modulates its fluorescent properties. Single FP reporters have been developed from engineered versions of Aequorea GFP, Clavularia sp. FP, Discosoma RFP and Entacmea quadricolor RFP. In this work, we have chosen to use a cp version of the monomeric Zoanthus sp. yellow FP-derived variant known as mPapaya0.4, the best mPapaya available at the beginning of this work, which has a unique emission wavelength around 540 nm. We reason that a Ca$^{2+}$ indicator that retained the fluorescence hue of mPapaya would fill the spectral gap between GCaMP and O-GECO in the visible wavelength palette of available FP-based Ca$^{2+}$ indicators and provide new opportunities for multicolor Ca$^{2+}$ imaging.

Results and discussion

Construction of the Y-GECO prototype and initial screening for improved variants

To construct a potential yellow FP-based Ca$^{2+}$ indicator, we replaced the cpGFP domain of G-GECO1.8 with the analogous cp version (N-term-148 to 238-GGTGGS-1 to 147-C-term) of mPapaya0.4. We designed 2 libraries in which one or two residues at the new N-terminus (residues 27 and 28 by Y-GECO numbering, Fig. S1, ESI†) and one residue at the C-terminus of cpmPapaya (residue 273, Fig. S1, ESI†) were randomized (Fig. 1A and B). The first version of the library had just residue 28 randomized at the N-terminus, while the second library had both 27 and 28 randomized. These two libraries were cloned into a customized vector that targets variants to the periplasmic space of E. coli cells. After plasmid transformation by electroporation, E. coli cells were plated on agar plates and formed individual colonies. Traditional on-plate manual screening of both libraries led to the identification of a variant (Y-GECO0.1) from the second library with an 88% decrease in fluorescence upon binding Ca$^{2+}$ (Fig. 1C). However, Y-GECO0.1 exhibited dim fluorescence and poor folding efficiency relative to mPapaya0.4. Two rounds of directed evolution for improved folding efficiency using manual screening led to the identification of Y-GECO0.5a and b (Fig. S1, ESI†) which were still dim but had better folding efficiency and similar dynamic range to Y-GECO0.1. At this point it was apparent that many more rounds of directed evolution would be necessary to improve Y-GECO to the point where it would be useful for live cell imaging. We reasoned that a $\mu$FACS-aided approach would accelerate this process and so we designed and applied a chip based on the work of Chen et al.22,25,36

Design and fabrication of $\mu$FACS platform

A simple, low cost $\mu$FACS platform is ideal for the directed evolution of FP-based Ca$^{2+}$ indicators as it greatly decreases the cost of fabrication and maintenance and can be readily adapted to the screen at hand compared to FACS. We elected to build a $\mu$FACS platform based on a polydimethylsiloxane (PDMS) microfluidic cell sorter chip with an integrated piezoelectric actuator. PDMS microchips containing 3 input channels and 3 output channels, each with a 5 cm channel for buffer mixing were fabricated using standard soft lithography techniques. For operation of this chip, cell samples were continuously flowed through the middle input channel and were confined to a single stream by sheath flow from the two side input channels. For a typical cell sample (~ $10^7$ cells per mL), the chip operated for more than 2 hours without obvious clogging. High signal-to-noise fluorescence detection of single bacterial
cells expressing Y-GECO variants was achieved using laser excitation at 488 nm with confocal detection using a photomultiplier tube (PMT) (Fig. 2D). The piezoelectric actuator integrated into the PDMS chip could be triggered for manipulation of fluid flow to achieve sorting into one of the side collection channels, with a 44% sorting efficiency of capturing bright cells that trigger actuation (Fig. 2A and Table S1, ESI†). The system was designed to have three output channels so that sorting of two different populations would be possible in principle. However, for the work described in this manuscript, we sorted all of the cells to just one of the two side outlets on the device.

To facilitate semi-automated cell sorting on chip, we designed a custom LabVIEW program for signal processing and sorting control. The signal-to-noise ratio (SNR) of cell fluorescence was calculated in real time based on the magnitude and background noise level of PMT signals. Fluorescent signals with SNR higher than a pre-defined threshold triggered sorting based on piezoelectric actuation. The delay between detection at the PMT and piezoelectric pump actuation (i.e., the response time of the system) was 2.2 ± 0.6 ms (25 trials). The time for an individual cell to travel to the sorting junction was matched to this response time by adjusting the position of the detection point within the channel. Response time was primarily limited by communication between the computer and the chip. An embedded field programmable gate array chip for a close-loop sorting control could significantly improve the response time.22

The major limiting factor that determines the throughput of the μFACS for library enrichment is the requirement that only one bacterial cell should be in the sorting junction at a single moment. As the dimension of the junction was about 150 μm × 150 μm × 50 μm, the excitation volume is roughly 1.1 nL. To avoid co-sorting of cells, there should be only one or less cell per excitation volume on average at a single moment. Given that the ratio of flow rate of sheath flow to sample flow was 10 : 1, the cell density of samples should be less than 1.87 × 10^7 cells per mL (see SI text and Fig. S3 for more detailed discussion, ESI†). When the total flow rate was between 10.5 μL min⁻¹ and 21 μL min⁻¹, the throughput of the system was between 150–300 cells per s. With this throughput, roughly 10^6 Y-GECO variants could be screened within 2 hours. In practice, we screened 5 × 10^5 to 1 × 10^6 variants in 2 hours with actual throughput around 75 cells per second to 150 cells per second.

Although our μFACS does not surpass FACS in terms of throughput and sorting efficiency, our devices cost less than $10 each for fabrication and are disposable. The whole μFACS system including all the components costs less than $10k, which is at least 50 fold reduction in cost compared to commercial FACS systems, such as MoFlo (~$500k).36 In addition, the μFACS system is amenable for customization to adapt to different purposes. The off-chip optical platform enables convenient installation of additional optical components including extra laser light sources for multicolor excitation, and extra dichroic mirrors and filters for spectroscopical separation of the emission light,18 etc. With an appropriate optical setup, researchers could use μFACS to enrichment variants based on more complex optical properties, such as photostability21,37 and efficiency of Förster resonance energy transfer.

**μFACS-aided directed evolution of Y-GECO**

We designed a μFACS-aided directed evolution strategy for improving Y-GECO variants with the focus on improving brightness (Fig. S2, ESI†). Briefly, *E. coli* cells expressing periplasm-localized Y-GECO variants were suspended in a Ca²⁺-free buffer (around 1 × 10^7 cells per mL) and loaded into the chip. The fluorescence signal of hundreds of cells was measured and analyzed to determine the threshold, which was the trigger to activate the sorting to collect the top 0.1% brightest clones.
Cells collected during the µFACS pre-screen were grown as colonies on agar plates and then manually screened in a manner identical to the traditional on-plate screen. Flow cytometric analysis of the initial and enriched libraries by µFACS (Fig. 2E) shows that our µFACS system could enrich bright variants and increase the fluorescence intensity of the dominant peak by a factor of 16. Unlike other GECOs, the fluorescence of Y-GECO variants is dim in the presence of Ca²⁺ and is bright in the absence of Ca²⁺. Therefore, Y-GECO variants targeted to the E. coli periplasm using the pTorPE expression plasmid should be in the bright fluorescent state in cells suspended in Ca²⁺-free buffer. Sorting of E. coli expressing Y-GECO libraries flowing on chip acquired by PMT detector. The SNR was defined as the ratio of the signal magnitude of cell fluorescence and the background noise level of PMT signals. Inset shows a bright variant. (E) Cytometric histogram for a library of Y-GECO before and after sorting using µFACS system. (F) Screening results of selected variants from round 1 to round 3 with and without aid of µFACS system. Best variants were combined from both methods between each round. Each point represents the performance (brightness × maximum Ca²⁺ response) of each variant relative to the best from the previous round. Gray boxes show the average of relative performance; n is the number of tested variants, larger n indicates more time and effort invested; p values were obtained from two-tailed t-test, showing µFACS gave much better performance in the first two rounds and similar performance with much less effort (n is smaller) in the third round.

Cells could come at the expense of indicator dynamic range. To overcome this problem, it is necessary to perform a secondary in vitro screen for Ca²⁺ response. Briefly, colonies were picked based on the colony brightness and grown in liquid culture. Proteins were extracted from overnight cultures and tested for Ca²⁺ response in microplate reader.

Starting from Y-GECO0.5a and b, which was prepared from the initial two rounds of directed evolution for improved folding, we performed three rounds of parallel screening using both manual on-plate screening and the µFACS-aided approach (Fig. S2, ESI†). To enable unbiased comparison of the two screening methods, after each round we combined the pool of the most superior variants found from both methods. The combined gene pool served as a template to generate a new library that was constructed by mixing gene variants generated by error-prone PCR with gene variants generated by DNA shuffling. The resulting library was then screened using both the traditional method and the µFACS-aided approach in parallel.
Compared to the traditional manual on-plate screen, the μFACS-aided approach identified improved Y-GECO variants using substantially less effort (Fig. 2F), owing to the 100 times higher throughput of μFACS. In each of the first two rounds, the overall brightness and Ca²⁺ response of Y-GECO variants selected by the μFACS-aided approach were significantly higher than variants selected by the traditional on-plate screen. For round 3 it was apparent that beneficial mutations had become exceedingly rare within our library, presumably as the template had already acquired the most highly represented beneficial mutations. For this round, both screening methods produced effectively equivalent results, however, we had to pick 44 variants to find one superior variant using the traditional approach, while the μFACS enrichment approach required only 10 variants.

After completion of the third round of directed evolution using both on-plate screen and the μFACS-aided approach, each method led to identification of one best variant that showed seemingly satisfactory brightness and Ca²⁺ response. Fluorescence measurement of the purified proteins revealed that the variant identified in the third round of μFACS platform-aided directed evolution was superior, relative to the variant identified in the third round of traditional screening, in terms of brightness and Ca²⁺ response (Table S2, ESI†). This variant, designated as Y-GECO1, has 17 mutations distributed throughout the protein sequence relative to Y-GECO0.1 (Fig. S1 and Table S2, ESI†), significant improvement on folding and brightness, a large ratiometric change, and an affinity to Ca²⁺ within the physiologically relevant range (see following section).

These results demonstrate that a μFACS-aided screen is more efficient for identifying improved FP-based Ca²⁺ indicators than the manual on-plate method. However, we believe that the full potential of this approach has not yet been fully utilized. The efficacy of directed evolution of FP-based Ca²⁺ indicators could be significantly increased by high throughput screening for both Ca²⁺ response and brightness simultaneously using a μFACS system. To achieve this, at least two detection points on both ends of the 5 cm main channel are required to facilitate detection of fluorescence changes of periplasm-targeted mutants before and after binding to Ca²⁺. In this new detection scheme, further optimization of experiment parameters, such as flow rate, cell density, buffer composition and optics, will be critical for the performance of the μFACS. Related work is currently in progress.

In vitro characterization of Y-GECO1

Spectroscopic characterization of the purified proteins reveals that the Ca²⁺ binding of Y-GECO1 leads to 95% decrease in fluorescence emission when using 525 nm excitation. Notably, a new fluorescence excitation peak around 413 nm appears upon binding to Ca²⁺ and fluorescence intensity increases 10-fold when exciting at this wavelength. Thus, Y-GECO1 is a ratiometric Ca²⁺ indicator with a 200-fold peak-to-peak ratiometric change (Fig. 3A) in fluorescence excitation. In this way, Y-GECO1 is analogous to the previously reported ratiometric-pericam³⁴ and

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**Fig. 3** Characterization and application of Y-GECO1. (A) Excitation and emission spectra of Y-GECO1, recorded with emission at 560 nm and excitation at 510 nm, respectively. (B) Comparison of brightness in the resting [Ca²⁺], of HeLa cells between Y-GECO1 (141 cells) and GCaMP6s (209 cells) using a common filter set for GFP. (C–E) Fluorescence images of HeLa cells expressing Y-GECO1 captured the moment of 10 μM histamine-induced intracellular Ca²⁺ rise. (C) Image acquired under 440 nm excitation. (D) Image acquired 480 nm excitation. Scale bar: 50 μm. (E) The ratiometric change \( \Delta R/R_{\text{min}} \) image (ratio \( R \) was calculated from dividing image C by image D; \( R_{\text{min}} \) was the minimum ratio of 440 nm/480 nm after adding ionomycin/EGTA. \( \Delta R/R_{\text{min}} = R/R_{\text{min}} - 1 \).) (F) Typical [Ca²⁺]; transients and oscillations of a select cell (enclosed by a dotted line in E) induced by histamine stimulations. The sampling interval was 3 s. (F, upper) A typical cell trace acquired under 480 nm (yellow line) and 440 nm (violet line) excitations. (F, lower) Excitation ratios, 440 to 480 nm. The right-hand ordinate calibrates [Ca²⁺], in pCa.
GEX-GECO1, but has a much larger dynamic range of Ca\(^{2+}\) response. In contrast to most previously reported GECO variants in which the pK\(_a\) of chromophore shifts forward to a lower value when bound to Ca\(^{2+}\), the pK\(_a\) of Y-GECO1 chromophore shifts from 6.3 to 8.9 upon binding to Ca\(^{2+}\) and results in loss of overall fluorescence intensity (Fig. S4B, ESI†). In vitro titration with Ca\(^{2+}\) indicates the value of K\(_{d}\) for Y-GECO1 is 190 nM (Fig. S4A and Table S2, ESI†).

Since mPapaya1 is superior to mPapaya0.4 in terms of brightness, photostability and maturation, we attempted to introduce some beneficial mutations of mPapaya1 relative to mPapaya0.426 into Y-GECO1, but none of them led to further improved variants (data not shown), which suggests Y-GECO1 and mPapaya1 evolved via different evolution paths. Interestingly, Y-GECO1 shares one mutation F223Y (F99Y by mPapaya numbering) with mPapaya1 (Fig. S1, ESI†), which is reported to dramatically improve the photostability of mPapaya0.6.26 According to the crystal structure of zFP538 (PDB ID: 2OGR), F223Y could enhance interaction with the internal z-helix of the FP and influence the chromophore environment, which may result in improved photostability. In the case of Y-GECO1, F223Y might also improve the brightness and Ca\(^{2+}\) response since it was the only new mutation found in the last round of screening (Fig. S1, ESI†).

Y-GECO1 is an excitation ratiometric Ca\(^{2+}\) indicator because, unlike Y-GECO0.1, the Ca\(^{2+}\) bound form exhibits long Stokes shift fluorescence due to its ability to efficiently undergo excited-state proton transfer (ESPT). That is, the protonated form of the fluorophore, which exhibits maximal absorbance at 413 nm and predominates in the presence of Ca\(^{2+}\), can become deprotonated in the excited state and emit fluorescence from the anionic form. ESPT has been observed in a number of other FPs and FP-based indicators including Aequorea victoria GFP,39 mKeima,40 ratiometric pericam,44 GEX-GECO1, and GEM-GECO1.8

We characterized the kinetics of the Ca\(^{2+}\) response by stopped-flow fluorescence spectroscopy. The k\(_{off}\) rate of Y-GECO1 was around 1.40 s\(^{-1}\) at 20 °C, which was slightly slower than the fastest version of GCaMP6, known as GCaMP6f, which has a k\(_{off}\) of 2.32 s\(^{-1}\) at 20 °C (Fig. S4E, ESI†). Although the kinetic response of Y-GECO1 should be sufficient for most of the applications, it has limited capability for resolving Ca\(^{2+}\) spikes and firing frequency variations in neurons4 and in some subcellular locations, such as the cytoplasm of dendrites and dendritic spines, where physiological Ca\(^{2+}\) can rise within milliseconds.41,42 Sun et al. have reported mutations that disrupt the CaM-RS20 interaction and increase the k\(_{off}\) of GCaMP3.4 Preliminary results indicate that analogous mutations improve the k\(_{off}\) of Y-GECO1. Specifically, Y-GECO1 ΔG13 (analogous to GCaMP3 ΔG47) has a k\(_{off}\) = 9.75 s\(^{-1}\) at 20 °C (Fig. S4E, ESI†), but a 60% decrease in dynamic range (Table S2, ESI†). Efforts to further optimize this fast Y-GECO1 variant are ongoing.

Performance of Y-GECO1 in mammalian cell and tissue culture

To determine how Y-GECO1 performs in typical live cell imaging experiments, we recorded the response of Y-GECO1 to changes of the free cytoplasmic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_i\)) in HeLa cells using established protocols.43 We used a CFP/YFP FRET filter set (438/24 nm excitation and 542/27 nm emission) for long Stokes shift yellow fluorescence and a FITC filter set (480/40 nm excitation and 535/50 nm emission) for observing short Stokes shift yellow fluorescence and GCaMP6s green fluorescence.5 While these filter combinations are not perfectly matched to the fluorescence spectra of Y-GECO1, these sets are widely available and commonly used in cell imaging experiments. As expected, we observed opposing oscillations in the fluorescence intensities of both emission channels when cells were treated with 30 μM histamine (Fig. 3F). Using this filter combination, the maximum intensity change of Y-GECO1 in HeLa cells was determined to be 6 ± 1-fold and the maximum ratio change was 13 ± 3-fold (Table S3, ESI†). In comparison, the maximum intensity change of GCaMP6s was 20 ± 7-fold intensiometric change (Table S3, ESI†). The ratiometric nature of Y-GECO1 facilitates quantitative measurement of Ca\(^{2+}\) concentration (Fig. 3F) relative to intensiometric indicators such as GCaMP6 and R-GECO1.5 Although the FITC filter set did not match well with the excitation and emission of Y-GECO1, Y-GECO1 in resting state was about 220% as bright as GCaMP6s (Fig. 3B). Presumably, the performance and brightness of Y-GECO1 could be further improved using a custom filter set with 425/40 nm and 500/40 nm excitation filters, and a 560/50 nm emission filter.

Y-GECO1 is capable of detecting Ca\(^{2+}\) transients in dissociated hippocampus neurons in wide-field fluorescence imaging with performance comparable to the current popular FP-based Ca\(^{2+}\) indicators GCaMP6s and R-GECO1. As shown in Fig. 4 (see also Movie S1, ESI†), Y-GECO1 successfully detected spontaneous cytosolic Ca\(^{2+}\) oscillations in both soma and processes with large ratio change and excellent SNR. Y-GECO1 was able to reveal different [Ca\(^{2+}\)]\(_i\) waveforms in the soma and processes (Fig. 4A and B). In order to estimate the maximum signal change in neurons, KC1 was added to a final concentration 50 mM to depolarize neurons and induce a large Ca\(^{2+}\) influx via voltage-activated Ca\(^{2+}\) channels.44,46 Using the filter set described above, the maximum change of Y-GECO1 was 8 ± 2-fold in ratio and 2.6 ± 0.5-fold in intensity of the FITC channel. In comparison, the maximum change of intensiometric indicator GCaMP6s was 8 ± 5-fold and R-GECO1 was 3 ± 1 fold (Fig. S5, ESI†). While the intensity change is smaller for Y-GECO1 than GCaMP6s, the ability to give ratiometric sensing is a major advantage.

To determine if Y-GECO1 is suitable for use with multiphoton excitation, we imaged neurons expressing Y-GECO1 using a 2-photon excitation laser-scanning confocal microscope. Rhythmic decreases in fluorescence could be easily observed using 990 nm 2-photon excitation with 35 mW output power (Fig. 4C and D), owing to the bright baseline fluorescence. The long Stokes shift fluorescence of Y-GECO1 could be detected using 850 nm 2-photon excitation (35 mW output power), however, it was much dimmer than the 990 nm excitation and the autofluorescence level was higher (Fig. 4E and F). Accordingly, excitation around 990 nm is preferred. Y-GECO1 is potentially an excellent inverted intensified Ca\(^{2+}\) indicator for two-photon imaging in vivo, and the bright baseline fluorescence of Y-GECO1 facilitates easier identification of transfected cells compared to intensiometric Ca\(^{2+}\) indicators which are dim in the resting state, assuming similar expression levels.
We tested the performance of Y-GECO1 in hippocampal neurons in organotypic brain slices at room temperature (20 °C) using confocal microscopy. The Y-GECO1 plasmids with Synapsin I or CMV promoters were delivered by ex vivo electroporation and expressed transiently in neural cells (Fig. 5A, C and D). The bright baseline fluorescence of Y-GECO1 facilitated identification of transfected neurons. We applied theophylline to excite the organotypic brain slices. Low millimolar doses of theophylline can evoke sustained rhythmic seizure-like activities in different neural networks including motor neurons and hippocampal neurons.47–50 Indeed, Y-GECO1 successfully detected evoked [Ca\(^{2+}\)]\(_i\) transients and oscillations with large signal magnitude (Fig. 5B and Movie S2, ESI\(^\dagger\)) as evidence of seizure-like discharge. Compared to cytoplasmic targeted Y-GECO1, processes could be satisfactorily visualized without saturation of the somata by tethering Y-GECO1 to the plasma membrane with the N-terminal peptide sequence of the Src kinase Lyn\(^\dagger\) (Fig. 5C and D). Lyn-Y-GECO1 expressed in glial cells was able to detect Ca\(^{2+}\) waves in various subcellular locations and process terminals (Fig. 5C). Although the FP body of Y-GECO1 belongs to Anthozoa-derived FPs, which are known to often form intracellular puncta in neurons,\(^51\) no bright intracellular puncta were found in neuronal cells expressing either cytoplasmic Y-GECO1 or membrane tethered Lyn-Y-GECO1. Our results suggest that Y-GECO1 can be a useful tool to study Ca\(^{2+}\) dynamic of neural cells in vivo with large signal magnitude.

In many cases, Y-GECO1 may be used as an intensiometric Ca\(^{2+}\) indicator with inverted fluorescence response to Ca\(^{2+}\) ("turn-off" indicator), as the dim fluorescence of the 410 nm excitation would not be useful if the background fluorescence was high. From a SNR perspective, intensiometric "turn-on" Ca\(^{2+}\) indicators with increased fluorescence response to Ca\(^{2+}\) would be superior for detecting Ca\(^{2+}\) oscillations compared to "turn-off" indicators with similar \(\Delta F\), mainly because the dim baseline fluorescence of "turn-on" indicators has less shot noise. However, in tissue samples, the dim baseline fluorescence of "turn-on" indicators may be buried by the high background fluorescence, whereas cells expressing "turn-off" indicators are more visible due to the bright baseline fluorescence. Furthermore, because most cells are in the resting state most of the time, the overall SNR of imaging of "turn-off" indicators would generally be better than "turn-on" indicators. In addition, Y-GECO could be combined with a different color of turn-on sensor that has similar \(K_d\) for ratio-metric Ca\(^{2+}\) imaging with significantly enhanced sensitivity.

Conclusions

We present a simple, low cost and disposable µFACS system with an integrated piezoelectric actuator for directed evolution of FP and FP-based indicators with throughput up to 300 cells per s. We developed a directed evolution approach that exploited µFACS for library enrichment, and ultimately led to improved variants of the single FP-based Ca\(^{2+}\) indicators Y-GECO. Three rounds of directed
The design of the microfluidic cell sortor was based on the work of Chen et al.22 Briefly, the device was fabricated using the conventional microreplica moulding technique22 in which device features were photolithographically defined by SU-8 photosist mold and then transferred to a PDMS substrate (Sylgard 184, Dow Corning). The actuation chamber of the PDMS substrate was created by manual cutting using a surgery knife, and the inlets and outlets were created using 2 mm punch. For device assembly, the PDMS substrate and glass were surface treated using a plasma generator (Electro-Technic Products Inc.). The PDMS substrate and the glass surface were placed into physical contact and the device was incubated at 65 °C for 20 min to allow the surfaces to bond. The stainless steel surface of the piezoelectric actuator (APC international) was aligned to the PDMS actuation chamber and put in contact with the PDMS substrate and sealed with commercially available syringe epoxy steel adhesive (Gorilla Glue). Finally, Teflon tubing was inserted into the inlets and outlets of the device for fluid flow introduction.

### Construction of Y-GECO gene libraries

The design of Y-GECO is based on well-established designs described by our group and others.8,33–35,53 Please see Supplemental experimental procedures for the detail of construction of Y-GECO gene libraries (ESI†).

### Screening of GECO gene libraries

The library screening of genetically encoded Ca\(^{2+}\) indicators using traditional colony-based manual screening method has been previously described in detail.8 For \(\mu\)FACS-aided screening, the screening protocol was modified accordingly. Briefly, a library of Y-GECO was used to transform competent E. coli and the transformed cells were plated on agar plates with roughly 30–100 times higher number of cells. It is critical that the freshly transformed cells should not be grown in liquid culture directly as many mutants that do not express protein will be enriched in culture due to faster growth times. This resulted in a high density of colonies (~10\(^7\) cells per plate) on the agar plate after overnight incubation at 37 °C. Cells were then collected by washing the colonies with M9 buffer supplemented with 100 \(\mu\)g mL\(^{-1}\) ampicillin, 0.1% glycerol, 0.002% L-arabinose and 1 mM EGTA. The cell density was measured by BD Accuri C6 flow cytometer (BD bioscience) and adjusted to 1–2 \(\times\) 10\(^7\) cells per mL accordingly.

The cell suspension was loaded into \(\mu\)FACS system via the middle input channel using a syringe pump (Harvard Apparatus). The sample flow was confined by the sheath flows (consist of the same M9 buffer) on both sides. The Y-GECO mutants that were flowing through the laser illumination point were excited by 488 nm argon ion laser beam, and the fluorescent emission was collected by a homemade confocal epifluorescence microscope using a 25× objective (numerical aperture (NA): 0.35, Leitz Wetzlar). A customized LabVIEW program was design for signal processing and chip control. PMT signals were acquired in sampling frequency 100 kHz. The noise level was calculated based on the variation of baseline PMT signals sampled in every 0.5 ms. The SNR of the fluorescence signal of variants was calculated in real time. The piezoelectric actuation would be triggered when the SNR signal of Y-GECO variants was larger than a pre-defined threshold that was determined based on statistical analysis of cell fluorescence signals in the samples prior to cell sorting. The sorted bright variants were collected via collection channels and grown on agar-plate overnight at 37 °C, picked in the next day and cultured in liquid LB medium at 37 °C overnight.

Proteins extracted from the overnight liquid cultures of the picked clones by either the traditional method or the \(\mu\)FACS-aided screening method was subjected to an additional screening in a Safire2 fluorescence microplate reader (Tecan). Briefly, cells were harvested by centrifugation at 13 000g for 2 min at 4 °C. Y-GECO variants along with other cytoplasmic proteins were extracted by suspension of cells using surfactant B-PER (Pierce) followed by centrifugation to pellet unwanted cell debris. The performance of each selected variant was evaluated in a fluorescence microplate reader. Their brightness and Ca\(^{2+}\)-dependent response was compared to the best variant from the previous round. Finally, the brightest functional variants were used as the library template for the next round of directed evolution.

### Protein purification and \textit{in vitro} spectroscopy

Please see Supplemental experimental procedures for the detail of protein purification and \textit{in vitro} spectroscopy characterization of Y-GECO (ESI†).

### Construction of mammalian expression plasmids

In this work, three types of mammalian expression plasmids were used. Please see Supplemental experimental procedures
for the detail of construction of mammalian expression plasmids (ESI†).

**HeLa cell culture imaging**

Please see Supplemental experimental procedures for the detail of preparation of HeLa cell culture (ESI†).

Wide-field imaging was done on a wide-field epifluorescence inverted microscope (Eclipse Ti-E, Nikon) equipped with a Photometrics QuantEM 512SC camera. The NIS-Elements Advanced Research (Nikon) was used for automated microscope and camera control. For determination of dynamic ranges in live cells, cells were imaged with a 20× air objective lens (NA 0.8). All imaging was performed at room temperature. For excitation of GCaMP6 and one channel of Y-GECO, the samples were illuminated with light from a 100 W mercury arc lamp that was passed through a 480/40 nm bandpass filter. The long pass dichroic mirror was 505 nm. The emission filter was 535/40 nm. For recording the long Stoke shift fluorescence of Y-GECO, 438/24 nm excitation filter, 458 long pass dichroic and 542/27 nm emission filter were used. 25% neutral density filter was used to decrease the strength of illumination for both indicators.

For imaging of histamine-induced Ca²⁺ dynamics, cells were imaged acquired with appropriate exposure every 4 s. Approximately 30 s after the start of the experiment, 100 μM histamine was added to reach a final concentration of 10 μM. Approximately 10 min after addition of histamine, 10 mM EGTA, 40 μM ionomycin in Ca²⁺- and Mg²⁺-free HBBS was added to the dish to reach a final concentration 1 mM EGTA, 4 μM ionomycin. At the end of the experiment, 20 mM CaCl₂, 40 μM ionomycin in HBBS was added to reach a final concentration 2 mM CaCl₂, 4 μM ionomycin.

**Imaging of rat hippocampal neuron culture**

Please see Supplemental experimental procedures for the detail of preparation of rat hippocampal neuron culture (ESI†).

The system setup for wide-field imaging was similar to that for HeLa experiment except of the following. A 60× oil objective lens (CFI Plan Apo NA: 1.4), or a 40× air objective lens (CFI Plan Apo NA: 0.9) was used for neuron imaging. For GCaMP6, 25% neutral density filter was used. For Y-GECO1, 12.5% neutral density filter was used. The exposure time for both indicators was 10 ms. For excitation of R-GECO1, the excitation filter was 545/30 nm, the long pass dichroic was 565 nm and the emission was collected through 620/60 nm filter. Neurons were imaged in the culture medium at room temperature 20 °C.

For confocal imaging, fluorescence signals were measured using a confocal microscope and software (20× XLUMPlanF1, NA: 1.00, water immersion objective; Olympus FV1000; Olympus Canada, Richmond Hill, Ontario, Canada). Similarly, Olympus FV1000 connected to a MaiTai DeepSee Ti:sapphire laser with a tunable excitation range from 710–990 nm (Spectra Physics, Santa Clara, CA, USA) was used for two-photon imaging. With reduced resolution and 2–4 μs per pixel scanning rate, the image acquisition was adjusted to be 1–3 frames per s to effectively detect Ca²⁺ oscillation of neurons expressing Y-GECO1.

**Preparation and transfection of rat hippocampal organotypic brain slices**

All procedures for brain slice experiments were carried out in compliance with the guidelines of the Canadian Council for Animal Care and with the approval of the University of Alberta Animal Care and Use Committee for Health Sciences.

Brains were dissected from 0-day-old (P0) Sprague Dawley rat in ice-cold HBBS containing 1.3 mM CaCl₂ and 1 mM MgSO₄. Horizontal 250 μm thick brain slices were generated using a vibrating microtome (Leica VT1000S, Leica Microsystems) as described previously⁵⁴ and were placed on a sterile 0.4 μm pore membrane cell culture insert (Millipore PICMORG50). The insert and slice were then placed in a Petri dish containing 1.5 mL of NbActiv4 (BrainBits) supplemented with 5% bovine serum (FBS) and penicillin-G potassium salt (50 units per mL), and streptomycin sulfate (50 μg mL⁻¹). Slices were cultured at 37 °C and 5% CO₂ for 24 h prior to transfection by electroporation. The insert and slice were then placed directly above a platinum plate Petri dish electrode (CUY700-P2E, Napa Gene), and the gap between the electrode and the membrane was filled with electroporation buffer (EB) (HBBS + 1.5 mM MgCl₂ + 10 mM glucose). Plasmids (pcDNA3.1, Life Technologies) for expression of the gene of interest were dissolved in EB at a concentration of 1 μg μL⁻¹ and sufficient volume was added to just cover the slice. A square platinum electrode (CUY700-P2L, Napa Gene) was then placed directly above the hippocampus slice and a power supply was used to apply five 20 V pulses (5 ms each, 1 Hz). The direction of electrical field was reversed and a second set of five pulses with the same settings was applied. The EB buffer was carefully removed from the slices and they were then put back and incubated at 37 °C and 5% CO₂.

**Confocal imaging of theophylline-induced Ca²⁺ elevations in rat organotypic hippocampal slices**

A modified version of protocol for Ca²⁺ imaging in brain slices⁵⁴,⁵⁵ has been used in this work. Immediately prior to imaging, the slices were perfused with superfusate containing (in mM): 120 NaCl, 3 KCl, 1 CaCl₂, 2 MgSO₄, 26 NaHCO₃, 1.25 NaH₂PO₄ and 10 D-glucose (pH adjusted to 7.4 by gassing with 95% O₂, 5% CO₂), at 5 mL min⁻¹ using a peristaltic pump (Watson-Marlow Alitea-AB) and kept at room temperature. Imaging was started within 10 min following activation of the perfusion system. Fluorescence signals were measured using a confocal microscope and software (20× XLUMPlanF1, NA: 1.00, water immersion objective; Olympus FV1000; Olympus). The cells were excited with 488 nm laser (Olympus) and emission was collected from 500–600 nm using a variable barrier filter.

Images were acquired at 2–3× digital zoom at a reduced frame resolution (256 × 256) and with 4 μs per pixel scanning rate, this allowed image acquisition to be 1–2 frames per s to effectively detect cytosolic Ca²⁺ oscillations of neurons and glial cells expressing Y-GECO1. For imaging of theophylline-induced Ca²⁺ dynamics, images were acquired every 0.5 s. Theophylline (10 mM, Sigma Aldrich, Canada) was dissolved in superfusate.
and bath applied via peristaltic pump approximately 30 s after the start of the acquisition. The drug was washed approximately 10 min later by switching back to control superfusate.

Accession numbers

The GenBank accession numbers for Y-GECO1 and Y-GECO1 ΔG13 are KJ193859 and KJ193860.

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