Single-cell analysis for drug development using convex lens-induced confinement imaging

Ndeye Khady Thiombane1, Nicolas Coutin1, Daniel Berard2, Radin Tahvildari2, Sabrina Leslie2 & Corey Nislow*1

KEYWORDS
CLiC • dose–response • methotrexate • phenotypic screen • single cell • yeast

1Pharmaceutical Sciences, 2405 Wesbrook Mall, University of British Columbia, Vancouver, BC, V6T1Z3, Canada; 2Department of Physics, 214 Rutherford Physics Building, McGill University, 3600 rue University, Montreal, QC, H3A 2T8, Canada; *Author for correspondence: corey.nislow@ubc.ca

ABSTRACT
New technologies have powered rapid advances in cellular imaging, genomics and phenotypic analysis in life sciences. However, most of these methods operate at sample population levels and provide statistical averages of aggregated data that fail to capture single-cell heterogeneity, complicating drug discovery and development. Here we demonstrate a new single-cell approach based on convex lens-induced confinement (CLiC) microscopy. We validated CLiC on yeast cells, demonstrating subcellular localization with an enhanced signal-to-noise and fluorescent signal detection sensitivity compared with traditional imaging. In the live-cell CLiC assay, cellular proliferation times were consistent with flask culture. Using methotrexate, we provide drug response data showing a fivefold cell size increase following drug exposure. Taken together, CLiC enables high-quality imaging of single-cell drug response and proliferation for extended observation periods.

METHOD SUMMARY
In this study, we use convex lens-induced confinement (CLiC) microscopy, which was originally developed for the study of single particles and biomolecules, and apply it to live cell imaging to detect fluorescently labeled cell compartments and compare the system to conventional imaging. We also performed an extended duration live-cell assay in CLiC by confining Saccharomyces cerevisiae yeast cells in well-defined pits in a flow cell and observing single-cell proliferation for 8 h. Finally, we assessed the effects of the antifolate drug methotrexate on cell morphology and mitochondria integrity by observing cells during drug exposure.

One of the driving forces in systems biology research is the emergence and development of new data-acquisition tools and technologies to better understand biological and biochemical mechanisms at the single-cell level. At present, most practical molecular and cell biology methods capture the average response of a cell sample population. However, results from these approaches can be misleading due to cellular heterogeneity. Such variation may arise from diverse genetic and nongenetic factors, including noise in gene expression. Factors that substantially define the amplitude of such noise include regulatory dynamics, transcription rates, genetic factors and intracellular copy numbers of molecules involved in the genetic network [1]. Single-cell heterogeneity is indeed a well-known phenomenon in microbial resistance and evolution studies, as well as in cancer research. For instance, most cancer cell lines display genomic alterations, as well as other forms of heterogeneity such as chromosomal instability [1,2]. Stochastic cellular process fluctuations, including the state, location, activity and concentration of polymerases, transcription factors and gene expression regulators, may also lead to diversification in an initially uniform cell population [3]. These variations can have a significant impact on the growth and environmental stress response of cells and cannot be captured using conventional cell biology methods. For instance, conducting a drug response analysis in yeast often requires recording its...
optical density (OD$_{600\text{nm}}$) at set time intervals over time. However, the recorded data are a bulk representation of the entire population and, as a consequence, may overlook fluctuations at the individual cell level that could lead to drug resistance [4].

The awareness of cell population heterogeneity and the importance of characterizing it have led to the emergence of a number of single-cell analysis tools such as fluorescence-activated cell sorting, microfluidic lab-on-a-chip and cellular microarrays. Microfluidics are a powerful, relatively recent innovation that are being applied for rapid antibiotic and antifungal susceptibility testing at a single-cell level, which can allow one to determine an isolate’s drug susceptibility in less than 30 min [5,6].

A range of antibiotic and antifungal drugs (e.g., penicillins such as mecillinam, amoxicillin-clavulate ampicillin, sulfamethoxazole, and so on) have been investigated using microfluidic lab-on-a-chip, and the results emphasize the utility of this technique to identify individual cells that maintain their fast growth rate against a background of drug susceptible cells [5]. The impact of single-cell analysis techniques on drug discovery and drug resistance studies in recent years underscores the clear need for more techniques that are robust in capturing not only the cell-to-cell variations, but also to chart their real-time proliferation and the evolution of the same individual cells for long periods.

Here we introduce a novel approach for single-cell studies using the convex lens-induced confinement (CLiC) instrument [7] combined with flow cells containing embedded pits. CLiC is an imaging system that was primarily developed for the isolation and visualization of single molecules in a flow cell made of two glass layers (Figure 1A & B). The principle of CLiC is as follows: using a lens-rod perpendicular to the focal plane, pressure is applied downward directly onto the flow cell. The resulting deformation of the flow cell confines molecules between the smooth glass surfaces and can load them into features such as pits, which are embedded in the surfaces (Figure 1B) [8,9]. This technique had been shown to be able to successfully confine and isolate diffusing [5] and interacting molecules [10–12] for long periods and improve fluorescent signal detection [7]. To date, CLiC has primarily been used to study materials, biomolecules and particles, with only limited information available regarding its suitability for live-cell investigations [13].

In this study, we used the yeast Saccharomyces cerevisiae, which is a well-known and extensively studied model organism used in the investigation of diverse eukaryotic cell biology. The insights gained from yeast are simplified but can be extrapolated into mammalian systems. Due to the high degree of conservation existing between yeast and mammalian genes (over 50% for essential genes), yeast is often used to understand the mechanisms of action of small molecules and drugs and to acquire preliminary data for drug
discovery efforts [14–16]. Additional advantageous criteria make yeast an ideal model organism to benchmark the CLiC cell assay. These criteria include: its low level of genomic instability compared with cultured cells, its simple and low-cost growth requirements, and its short doubling time, which allows faster drug effects and pathway perturbation assessment. We therefore used yeast to validate the CLiC instrument for single-cell analyses, specifically comparing the CLiC imaging platform to a conventional microscope for fluorescent cell imaging, drug response studies and cell-proliferation observations.

**MATERIALS & METHODS**

**Yeast cell propagation & staining**

For localization experiments with fluorescent probes, we used BY4743 diploid cells (MATa/his3Δ1/his3Δ1 leu2Δ0/leu2Δ0 lys2Δ0/+ met15Δ0/+ ura3Δ0/ura3Δ0) generated from a cross between BY4741 and BY4742 [17].

**Vacuolar staining**

We inoculated BY4743 cells in 3 ml YPD media and grew them overnight (~16 h) at 30°C. We subsequently adjusted the cell suspension to an OD_{600 nm} of 10, added 4 μM FM4–64 vacuolar stain in YPD and incubated on an interval mixer for 30 min at 30°C. We then removed the excess probe by centrifuging at 500×g for 1 min, aspirating the supernatant and washing cells twice in YPD before suspending in 1 ml YPD.

**Nuclear DNA live cell staining**

We stained BY4743 cells in mid-log phase with an OD_{600 nm} of 0.5 using 2.5 μg/ml final concentration of 4′,6-diamidino-2-phenylindole (DAPI) on an interval mixer for 30 min at 30°C. We then harvested the cell pellet by centrifugation at 500×g for 1 min, washed with 1× phosphate-buffered saline and resuspended in YPD for a final OD_{600 nm} of approximately 10.

**Mitochondrial staining**

Upon treatment of BY4743 cells with 2 mM methotrexate or 2% dimethyl sulfoxide (DMSO) as a vehicle control for 20 h, we pelleted and resuspended cells in prewarmed (30°C) staining solution containing YPD media and 200 nM Mitotracker Red CMXRos. We subsequently incubated the cells at 30°C with shaking for 30 min in a thermomixer, pelleted and resuspended them in fresh YPD media prewarmed at 30°C.

**CONVEX LENS-INDUCED CONFINEMENT**

The CLiC instrument is composed of five major components assembled to allow confinement of single molecules or cells into embedded wells in the CLiC flow cell, which we refer to in this paper as pits. (i) The first component is a mounting stage that perfectly fits into an inverted microscope’s main stage. (ii) The second component is a mounting chuck that holds the flow cell, enables fluid exchange and is inserted onto the mounting stage. (iii) The third and main part is the flow cell made of two thin glass coverslips. The bottom coverslip contains embedded wells and is attached to the upper flat coverslip by a 30-μm thick double-sided adhesive [11]. Cell samples can be loaded between the two layers through a fluidic access port. (iv) The fourth component is a piezo, which controls the y-axis of the lens rod. (v) Last is the lens-rod, which consists of a convex lens, mounted curved-side down, on a narrow rod. When lowered, the lens-rod pushes on the top coverslip, deforming it downward, which consequently restrains cells into the...
embedded wells. This instrument does not tether or stick down cells, but simply confines them. Cells can then be imaged following the microscope’s standard protocols.

**WIDE-FIELD MICROSCOPY**

**Image acquisition on microscope slide & processing**

We used a 63× 1.4 NA oil-immersion objective mounted on a Zeiss (Oberkochen, Germany) Axiovert 200 m microscope, an Excelitas (MA, USA) X-Cite 120 LED light source and a Zeiss AxioCam HRm Rev.2 camera to capture images of cells in bright-field channel exposed for 10 ms under 4.0-V transmitted light. We used excitation and emission filters appropriate for FM4–64 (Cy3, excitation/emission: 515/640 nm), DAPI (excitation/emission: 359/461 nm) and Mitotracker Red (Cy3, excitation/emission: 579/603 nm) with 830 ms, 125 ms and 150 ms of exposure, respectively. Images were captured using the multi-dimensional acquisition setting on the AxioVision 4.8.2 software (Zeiss).

**Image acquisition in flow cell & processing**

For cell imaging using the Scopesys (QC, Canada) CLiC instrument, we loaded 5 µl of the stained cell suspension into the flow cell-containing embedded pits and imaged with the 63× oil-immersion objective as described above. We captured images in the brightfield channel using 100 ms exposure under 11.0 V transmitted light and in the DAPI or Cy3 fluorescent channels with acquisition settings identical to those used for imaging cells on a microscope slide.

**Microscopy image analysis & fluorescent signal quantification**

ImageJ software (with the Fiji plugin) was used to measure the ‘area’, ‘integrated intensity’ and ‘mean gray value’ of fluorescent cells [18,19]. We first selected the regions of interest (ROI) delineating every single cell on channel 1 (bright-field-transmitted channel) and subsequently performed fluorescent signal measurements on channel 2 (fluorescent channel). For normalization, we measured an empty background area on the field of view and determined the corrected total cell fluorescence (CTCF) using the following formula:

$$\text{CTCF} = \frac{\text{Integrated Density} - \langle \text{Area of selected cells} \times \text{Mean fluorescence of background readings} \rangle}{10,000}$$
Figure 3. Corrected total cell fluorescence of DAPI-stained cells under confinement in flow cell vs on microscope slide. (A) DAPI-stained BY4743 cells captured under Brightfield (BF) and fluorescent (DAPI) channels. Imaging performed under flow cell confinement (top) and on a microscope slide (bottom). (B) Column scatter plot of CTCF of n = 100 cells per condition. Image displays individual data (blue markers), their average (black horizontal lines) and standard deviation (blue error bars) for each condition. ***p < 0.05 under an unpaired two-tailed t-test.

BF: Brightfield; CLiC: Convex lens-induced confinement; CTCF: Corrected total cell fluorescence; DAPI: 4',6-diamidino-2-phenylindole.

Figure 4. Mitochondria fluorescent imaging of methotrexate versus dimethyl sulfoxide-treated cells. Mitotracker-stained BY4743 cells treated for 20 h with 2% DMSO vehicle control (top) or 2 mM methotrexate (bottom) and captured in flow cells under Brightfield (BF) and Cy3 Fluorescent (Mitotracker Red™) channels. BF: Brightfield; DMSO: Dimethyl sulfoxide; MTX: Methotrexate.
Cell proliferation under flask versus flow cell platform
For the growth comparison experiment, we used the dfr1/DFR1 heterozygous deletion strain, derived from BY4743 [20]. This strain is deleted for a single copy of the enzyme dihydrofolate reductase, which is the established target of methotrexate [21].

Yeast growth in flasks
We propagated a cell suspension, characterized by an optical density measured at 600 nm (OD\textsubscript{600 nm}) of 0.25, in 50 ml YPD in a flask at 30°C for three generations, and performed readings with an Eppendorf (Hamburg, Germany) Biophotometer every 15 min.

Yeast growth in flow cell
We loaded 5\,\mu{l} of a cell suspension with an OD\textsubscript{600 nm} of 5 into a flow cell, and grew the cells at 30°C. To monitor and control the temperature of the flow cell environment, we used a silicon heat tracing cable and temperature sensor wrapped around the 63× oil-immersion objective. The Zeiss Axiovert 200 m microscope captured images of proliferating cells every 15 min for 8 h.

Viability assay
To prepare the samples for the spot assay, we grew BY4743 cells in the presence of 2.0, 1.0, 0.5 or 0.25 mM methotrexate, or 2% DMSO vehicle control for 20 h, normalized them to an OD\textsubscript{600 nm} of 1 and transferred them to a 96-well plate. We subsequently performed fivefold serial dilutions on each sample, spotted them onto YPD agar by transferring 5\,\mu{l} of each suspension, and grew the cells at 30°C for 48 h prior to image acquisition.

RESULTS & DISCUSSION
Fluorescent signal detection comparison in flow cell versus microscope slide
To study the ability to obtain fluorescent signals from live cells on the CLiC instrument and compare the data to that obtained from conventional microscope slides, we captured images of BY4743 cells stained with 4-\muM FM4–64 vacuolar rim stain on a microscope slide and a CLiC flow cell under similar fluorescent settings. An unstained BY4743 negative control sample was also analyzed. Both platforms detected fluorescent signals consistent with vacuolar rims in stained cells (Figure 2A). From visual inspection, the CLiC instrument captured the stained vacuolar rims with high contrast. Quantitatively, the average signal-to-noise ratio with CLiC was enhanced compared with the microscope slide, 5.1 versus 3.1, respectively (Figure 2B). The CTCF distribution of the stained samples on the two platforms were not significantly different; however, the distribution obtained with CLiC extended over a larger dynamic range of signals, highlighting the CLiC ability to detect higher signals (Figure 2B).

Signal detection of cells stained with 2.5\,\mu{g/ml} DAPI for 30 min also demonstrated signal detection sensitivity differences between the CLiC flow cell versus the microscope slide. The average integrated fluorescence per stained cell was 10,517 for \( n = 110 \) cells visualized on a microscope slide compared with 17,192 for \( n = 104 \) cells imaged under confinement with CLiC. Under a student’s two-tailed t-test, assuming equal variances between the samples, the distribution of signals was significantly higher in CLiC flow cells (\( p < 0.05 \)) (Figure 3A & B).

The fluorescence signal detection comparison validated the ability of the CLiC instrument to image cellular compartments with high resolution and precision (Figures 2A & 3A). Additionally, DAPI fluorescent signal detection was greater with increased sensitivity in CLiC flow cells.
cells compared with microscope slides. This higher sensitivity could be explained by the decrease in the vertical volume of the cells when squeezed, allowing higher background signal rejection and better fluorophore detection [8].

Methotrexate effect on yeast in flow cells

After validating that the CLiC instrument was suitable for fluorescence imaging, this platform was used to study the effect of the antifolate drug methotrexate, a dihydrofolate reductase inhibitor, on BY4743 cells [21]. In yeast, the dihydrofolate reductase enzyme is encoded by DFR1 and is required for tetrahydrofolate biosynthesis. This enzyme is functionally conserved between yeast and human cells; that is, the viability and antifolate resistance of a yeast dfr1 mutant can be rescued by expression of the human enzyme DHFR [20]. Inhibition by methotrexate results in fitness defects of yeast cells, which is likely due to the adverse effects of the drug on nucleic acid and amino acid biosynthesis [21]. Dfr1 has previously been reported to be necessary for mitochondrial morphology maintenance in the presence of 10 μg/ml doxycycline [22]. We therefore hypothesized that inhibiting Dfr1 with methotrexate would have an impact on the mitochondrial structure of the cells.

BY4743 cells treated with 2.0 mM methotrexate for 20 h showed no major mitochondrial structural damage based on observations of Mitotracker Red stained cells (Figure 4). Methotrexate treatment did, however, result in a drastic morphology change in BY4743 cells. Treated cells were almost fivefold larger in volume than the 2% DMSO-treated cells, with an average volume ± SEM of 900.8 ± 35.41 μm³ (n = 100) compared with 185.9 ± 8.168 μm³ (n = 100), respectively. Most strikingly, methotrexate had a pronounced effect on cell cycle progression. When assayed for budding index [23], we found a 1.36-fold increase in the population budding index of n = 200 treated cells (87%) compared with the DMSO-treated control (64%) (Figure 5A & B). When treated with methotrexate, 73% of the cell population was found in the telophase/ anaphase stage of the cell cycle compared with 16% for the DMSO-treated control, suggesting that methotrexate either caused mitotic arrest or delayed mitotic exit [24].

To further investigate methotrexate’s effect on BY4743 cell viability, cells treated with 2.0 mM methotrexate for 20 h were spotted onto YPD agar for 48 h. To our surprise, despite their dramatic morphology change, budding alterations and growth inhibition (Figures 3, 4A & 4B), methotrexate treatment did not exhibit a major impact on the cell viability following recovery on YPD in the absence of methotrexate (Figure 6). This observation suggests that methotrexate is cytostatic and not cytotoxic under the conditions tested.

Yeast cell growth study in flow cell

To investigate whether the CLiC instrument is useful for observing live cell growth, cells in log phase were re-suspended in YPD, adjusted to an OD₆₀₀ nm of 5 and loaded into a flow cell confined by CLiC. Observations recorded at 30°C using brightfield illumination (refer to ‘Materials & methods’ for temperature control). Confined single cells had an initial doubling time that ranged from 1 h 30 min to 2 h 30 min (Figure 7A & B), with an average of 2 h 15 min, which was consistent with the first doubling time recorded when the same strain is grown in flasks (Figure 7C & D, Supplementary Video 1). With the CLiC instrument and a cell suspension with an initial OD₆₀₀ nm of 5 prior to loading, we were able to reproducibly observe two cell doublings (Supplementary Videos 1–4). This assay validated the use of the CLiC instrument for single cell proliferation studies and could be used to observe the real-time drug response variabilities among a cell population.

This study validated the use of the CLiC instrument for accurate single-cell investigations. We observed that fluorescently labelled cells had improved background rejection (i.e., lower nonspecific fluorescence), which resulted in a higher signal-to-noise ratio within the flow cell compared with the microscope.
slides. The platform detected cellular DNA with higher precision and sensitivity, which also applied to other cellular compartments (vacuoles and mitochondria in this study). Most importantly, the CLiC instrument allows robust growth of yeast cells during single-cell confinement and observation in real time. This device could be used in future studies to observe the real-time emergence of resistant drug suppressors as well as yeast and bacterial persistence [4], a phenomenon whereby, upon drug treatment, a small surviving subgroup regrows into a population that has not acquired any resistance and is still as sensitive to the treatment. This device also holds great potential for routine use in fields such as cancer research, where single-cell variabilities and instabilities are especially important for developing effective therapies. While we focused on yeast cells as a model system in this work, previous studies performed on adherent Chinese hamster ovary (CHO-K1) cells demonstrated the feasibility of customizing the CLiC instrument with a cytoindentor, which allows precise indentation and imaging of live mammalian cells without significantly changing their oxidative stress or causing detectable impacts on cell fitness [13]. To manipulate cell adherence, the flow cells could be customized and coated with fibronectin as previously demonstrated [13]. Further optimization and validation studies will, however, be

Figure 7. Yeast cell growth comparison in convex lens-induced confinement instrument flow cell versus flask. (A) Timepoint 00:00 image of cells confined in convex lens-induced confinement flow cell and grown at 30°C in YPD for 8 h (refer to Supplementary Video 1 for complete growth cycle) and numeric labeling of pits containing cells. (B) Average doubling time of cells in Figure 7A numerically labelled pits. (C) Growth curve of cells grown in flask at 30°C with shaking in YPD, n = 2. (D) First, second and third doubling time of cells grown in flasks and deduced from Figure 7C.
that can be obtained from fluorescently labeled molecules should aid in the subcellular localization of ligands in single cells.

While generating arrays of cells within an imaging field is not a new idea [25], CLiC provides a flexible (the only constraints are in the initial flow cell design) means to achieve living cell microarrays that are unique in that no tethers are required to keep cells within the focal plane over extended periods. The next challenges with CLiC as applied to phenotypic screens will be to increase the capacity of the flow cells and to introduce automation, for example, for loading flows cells and liquid exchanges.

SUPPLEMENTARY DATA
To view the supplementary data that accompany this paper please visit the journal website at: www.future-science.com/doi/suppl/10.2144/btn-2019-0067

AUTHOR CONTRIBUTIONS
NKT, NC, DB, SL and CN designed and supported the microscopy experiments; NKT acquired and analyzed the data; RT developed and fabricated the flow cells. NKT, SL and CN wrote the manuscript.

ACKNOWLEDGMENTS
We thank ScopeSys (www.scopesys.ca) for access to the CLiC technology package. The project was supported by a grant from the CFI to CN.

FINANCIAL & COMPETING INTERESTS DISCLOSURE
SL acknowledges the NSERC Discovery, Accelerator, and Idea to Innovation programs as well as McGill University for research funding and support. SL and DB have a financial interest in ScopeSys, the company which is commercializing CLiC imaging. The authors have no other relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript apart from those disclosed.

No writing assistance was utilized in the production of this manuscript.

OPEN ACCESS
This work is licensed under the Attribution-NonCommercial-NoDerivatives 4.0 Unported License. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/4.0/

REFERENCES