

On-chip continuous monitoring of motile microorganisms on an ePetri platform†

Seung Ah Lee,^{*a} Guoan Zheng,^a Nandini Mukherjee^a and Changhui Yang^{ab}

Received 24th January 2012, Accepted 13th March 2012

DOI: 10.1039/c2lc40090a

Self-imaging Petri dish platforms with microscopy resolution, which we term ‘ePetri’, can significantly streamline cell cultures and/or other longitudinal biological studies. In this paper, we demonstrate high-resolution imaging and long-term culture of motile microorganisms in a specialized ePetri platform by taking advantage of the inherent motion. By applying a super-resolution algorithm to a set of low-resolution images of the microorganisms as they move across the sensing area of a complementary metal oxide semiconductor (CMOS) image sensor chip, we can render an improved-resolution image of the microorganisms. We perform a longitudinal study of *Euglena gracilis* cultured in an ePetri platform, and image-based analysis on the motion and morphology of the cells. As a miniaturized and automated culture monitoring platform, this ePetri technology can greatly improve studies and experiments with motile microorganisms.

Introduction

Optical detection and monitoring is a key tool in interrogating the growth and behaviour mechanism of microorganisms. Various optical methods, such as optical density, speckle pattern, spectral analysis, colorimetry and bright field/fluorescence imaging are used to measure the changes in various motion and growth parameters of microorganisms in a non-invasive manner.^{1–5} Above all, image-based monitoring gives more direct and detailed information about the culture by visualizing the morphological and behavioural changes of each individual cell. In addition, current computational capabilities allow for automated longitudinal imaging and analysis, with low cost, reduced labour and better temporal resolution.^{6–9}

A microscope has long been a standard piece of equipment for conducting imaging-based studies. However, time-resolved imaging using an expensive microscope increases the cost of these experiments. In addition, numerical aperture limitation of the objective lens in a microscope prohibits high-resolution imaging over a large area without the use of additional actuation systems. Miniaturization of imaging tools can benefit many scientific researches, such as study on the swimming behaviour of flagellated protozoa^{1,10} and behavioural studies using *Caenorhabditis elegans* models.¹¹ In addition, it can lower the cost of many clinical applications such as sperm counting,¹²

toxin screening assays using fresh water microorganisms,¹³ and blood/water-born parasite diagnostics.¹⁴

In recognition of this need, low-cost on-chip imaging techniques, such as optofluidic microscopes (OFM),^{15,16} have recently been developed. In the OFM, the sample is carried along a microfluidic channel on a CMOS image sensor, during which light transmission through the sample is collected and the time-resolved transmission signal is converted into spatial information of the sample. In the case of the most recent version of OFM, referred to as sub-pixel resolving optofluidic microscope (SROFM),¹⁶ a pixel super-resolution algorithm is used to combine a sequence of low-resolution shadow images to a high-resolution image. OFM methods rely on translational delivery of the target object by electro-kinetic or laminar flow inside a microfluidic channel, which makes it tough to continuously monitor the same living specimen repeatedly over a long period of time.

Recently, we have reported a self-imaging Petri dish for biological studies - the ePetri dish platform.¹⁷ Here, the CMOS image sensor serves as the substrate of cell growth. The ePetri allows for continuous monitoring of live specimen in a large field of view over a long period of time, using minimal labour and at low cost. In our previous work, a sub-pixel sweeping microscopy (SPSM) method was used in order to boost the resolution beyond the pixel size of the sensor. SPSM uses raster scanning of a light source to create sub-pixel-shifted low-resolution shadow images, which are later combined into a single high-resolution image with a pixel super-resolution algorithm. This approach works well for biological targets that are relatively immotile, as the time required to acquire an adequate sequence of low-resolution shadow images is relatively long. For moving microorganisms, the SPSM ePetri is simply unable to perform good imaging.

^aDepartment of Electrical Engineering, California Institute of Technology, Pasadena, CA, 91125, USA. E-mail: salee30@caltech.edu;

Fax: +1(626) 3958475; Tel: +1(626)3952258

^bDepartment of Bioengineering, California Institute of Technology, Pasadena, CA, 91125, USA

† Electronic Supplementary Information (ESI) available. See DOI: 10.1039/c2lc40090a/

Fortunately, we believe that the inherent motion of the microorganisms can be used as a means to obtain the sub-pixel shifts that are required for pixel super-resolution image reconstruction. In this paper, we report a strategy that combines key ideas of SROFM and ePetri to appropriately perform microscopy imaging of motile microorganisms. We name this technique sub-pixel motion microscopy (SPMM). Using SPMM, We first demonstrate high-resolution imaging by taking advantage of the swimming behaviour of motile protozoa *Euglena gracilis*, which is a well-known model organism for cellular tracking studies and many biological assays. Then we conduct a longitudinal study of *Euglena gracilis*, demonstrating microorganism counting, tracking and statistical analysis capabilities using the described ePetri system. Through the proof-of-concept experiments, we show that the ePetri platform using SPMM method allows for inexpensive, miniaturized and minimum-labour culture monitors that can benefit clinical and scientific research.

Results and discussion

Operation principle and device structure

The SPMM ePetri dish platform takes a very simple geometry; a bare CMOS image sensor and a poly(dimethylsiloxane) (PDMS) well mounted on top. As the microorganisms are cultured in the ePetri dish, the shadow images of the cells are continuously captured through the sensor. The resolution of the raw images is fundamentally limited to be equal to twice the pixel size of the image sensor by the Nyquist criterion (the pixel size is 2.2 μm in our experiment). The use of the SPMM method improves the optical resolution of the final images beyond this limit. SPMM is based on the pixel super-resolution algorithm,^{18,19} where low-resolution images of the same object with sub-pixel shifts are combined to a higher resolution image. For motile microorganisms, their inherent motion allows themselves to be scanned on the sensor's pixel grid, providing the sub-pixel shifts between each low-resolution frame that are essential for the image reconstruction. Collected low-resolution images are redistributed into a denser grid of a single high-resolution image, with the redistribution vector determined by the sub-pixel displacement of the cell at each frame with respect to the first frame. We use minimum of n^2 low-resolution frames for a single high resolution image reconstruction, where n is the enhancement factor, defined as the size ratio between the original low-resolution pixel and the virtual high-resolution pixel.

The ePetri system is shown in Fig. 1. Each ePetri chip containing the sample is loaded into a camera board. Customized software controls the camera through a frame grabber and performs subsequent image processing for pixel super-resolution reconstruction. We used Aptina MT9P031 5 megapixel image sensors with the pixel size of 2.2 μm , which has the total imaging area of 5.7 mm \times 4.8 mm. As in OFM systems, the optical resolution of the ePetri is the highest at the floor of the image sensor. In order to keep the microorganisms as close to the sensor surface as possible, we dispensed a small amount (<5 μL) of culture medium containing the specimen into an ePetri chip, and confined the vertical location of the microorganisms to less than 100 μm from the sensor's surface. A drop

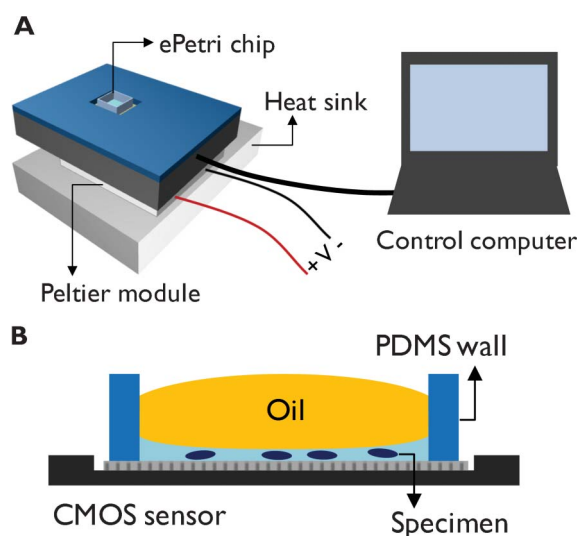


Fig. 1 (a) the ePetri dish imaging platform is composed of an ePetri chip, camera module and a control computer. A Peltier module and a heat sink are used to maintain the temperature of the chip. (b) Schematic diagram of a single ePetri dish chip. Sample is dispensed into a chip and a drop of oil is used to prevent evaporation of the medium.

of oil was put on top of the culture medium on order to prevent drying during the experiment.

SPMM imaging of motile microorganisms with ePetri platform does not require the use of a special light source, and can use the room lighting or natural lighting for illumination. We used a green LED lamp (0.2 W m⁻²) in all experiments as an illumination source to keep the phototaxis level identical in all cultures. Under these illumination conditions, 0.2 ms exposure for single frame was found to be adequate. Since the camera board can exhibit elevated temperature during the course of long term imaging, we used a Peltier module and a heat sink to dissipate the heat from the board.

SPMM imaging of *Euglena gracilis*

Pixel super-resolution image reconstruction in SPMM method requires a sequence of shadow images where the target object is continuously translated. The raw images captured through the sensor can be considered as undersampled images of the original object with a known sub-pixel shift between each frame. Upon reconstruction, the low-resolution images are enhanced by n times, meaning that the physical size that a single pixel represents decreases by a factor of $1/n$ in the reconstructed high-resolution image. This sub-pixel shift, which we denote as (d_{k_x}, d_{k_y}) for k^{th} image in the sequence in the x and y directions determines the vector in which n^2 different low-resolution shadows are redistributed within the denser pixel grid of a single high resolution image. However, unlike SPSM method where a scanning light source creates shadows that are incrementally shifted with the fixed amount of displacement along both x and y direction, The SPMM method is based entirely on the inherent motion of the microorganisms. This implies that we need to accurately trace and calculate each microorganism's position in every frame. As such, the sub-pixel shifts, here on referred to as motion vectors, of each cell are obtained by tracing the rough

position of each cell throughout the low resolution sequence. Details on the method by which we obtain motion vectors of the microorganisms and circumvent the reconstruction errors caused from the inadequate motion of the cells are discussed later in this paper.

We used *Euglena gracilis* for demonstration of the SPMM imaging technique. *Euglena gracilis* is an ellipsoidal protozoan with the dimensions approximately equal to $60 \times 10 \times 10 \mu\text{m}$. It is well known that its motion is propelled by a long flagellum beating around the body, and it moves in a rototranslatory motion with a speed of about $100\text{--}400 \mu\text{m s}^{-1}$.^{1,20,21} Its body is slightly tilted with respect to the axis of the movement.²⁰ Considering the motion parameters of *Euglena gracilis*, we set the imaging frame rate at 180 fps, where we typically used 16–25 images to reconstruct a single high resolution image with the enhancement factor $n = 4\text{--}5$. The lengths of the sequences represent the image of the cell during approximately 0.1 s, which typically gives $10\text{--}40 \mu\text{m}$ displacement of the microorganisms within the sequence and makes it safe to assume that the internal sub-cellular structures are static and the rotation perpendicular to the axis of translation is negligible. Due to the limitations in the data transfer rate of the camera, we used a smaller field-of-view (FOV) of 400×200 pixels.

Fig. 2 shows the comparison between the raw and the reconstructed high-resolution image of an *Euglena gracilis* with $n = 5$. The motion vector for each *Euglena gracilis* cell was calculated using a simple image processing technique. In the raw sequence, the centre locations of the organisms were found in each frame with the 0.5 pixel accuracy and then these values are smoothed throughout the sequence to estimate the cell location with sub-pixel accuracy. To reduce the amount of computation in the image reconstruction, each image is cut with a fixed window size around the target cell (17×17 pixels) and used the relative shifts ($d_{kx} - [d_{kx}], d_{kv} - [d_{kv}]$) for reconstruction. The raw

sequence used for this reconstruction and the measured motion vector is shown in Fig. S1, ESI†.

Since the basic principle of imaging is similar to SROFM method, the ultimate optical resolution of SPMM is similar to that of SROFM, reported as $0.66 \mu\text{m}$ at best.²² The optical resolution of SPMM ePetri depends on the pixel size of the sensor, number of low resolution images used for the reconstruction and the motion of the microorganism. However, due to the erratic nature of the inherent motion of a microorganism, such as rotation, sudden change of direction and non-uniform speed, we can expect that the resolution would be poorer than the SROFM method. From the images we obtained with SPMM ePetri, we were able to observe a small bright feature with the full-width-half-maximum of $0.95 \mu\text{m}$, which is comparable with the resolution of a 0.30 NA $10\times$ objective lens ($0.92 \mu\text{m}$ resolution) used for comparison in Fig. 2(c). In addition, the resolution would be poor for larger and thicker microorganisms, due to the increased distance between the imaging plane and the sensor's surface. The lower limit to the size of microorganism that can be imaged is determined by the pixel size of the sensor; cells that are far smaller than the pixel cannot be recognized in the raw image, and thus are unable to be traced using the motion vector. With the $2.2 \mu\text{m}$ pixel sensor, we have successfully traced and imaged $0.5 \mu\text{m}$ features (microspheres) using the pixel super-resolution algorithm.^{17,22}

It is worth noting that there exist some criteria in the cellular motion for a proper image reconstruction. In order for the entire part of the cells to be scanned with sub-pixel displacement, the translation of the cells should be at a certain angle with respect to the sensor's pixel grid so that the sample is scanned with shifts both x and y directions. Assuming the linear translation of the cells within a sequence of N frames, the motion vector should satisfy following condition.

$$\min(|d_{Nx}|, |d_{Ny}|) > 1 \quad (1)$$

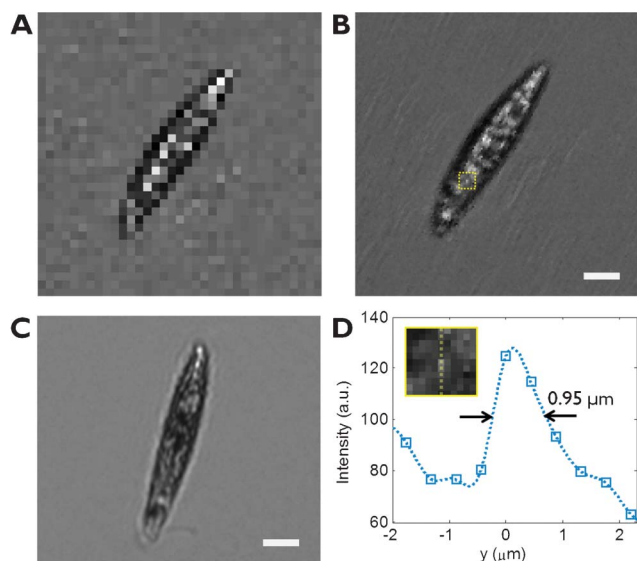


Fig. 2 (a) Raw and (b) reconstructed images of *Euglena gracilis* using SPMM method with $n = 5$. (c) A conventional microscope image of a *Euglena gracilis* taken with a $10\times$ objective lens. The scale bar indicates $10 \mu\text{m}$. (d) The line traces of a small bright feature in the reconstructed image. Inset corresponds to the area highlighted in (b).

When such condition is not satisfied, the sample is not entirely scanned and the reconstructed image shows empty pixels in a form of stripes (horizontal when d_v , vertical when d_x is not enough.) with the period of the enhancement factor n (Fig. 3(A2)). In order to prevent missing information in the image, we obtain the low resolution sequence for longer amount of time than 0.1 s, find the sequence where the angle of motion satisfies the condition in Eqn (1) with minimum N (Fig. 3(A3) and Fig. S2†), or use more frames to satisfy Eqn (1).

Rotation of the cells can also cause artefacts in the reconstructed image. Although the rotation in a typical motion of a *Euglena gracilis* during the imaging period of 0.1 s is small compared to the translational component, the rotational frequency may increase depending on the condition of the cell¹ and a cell may change its direction abruptly upon incidence with an obstruction. Fig. 3(B2) and 3(C2) show reconstructed images with periodic rotation artefacts at the tip of the microorganisms. To take the rotation of the cell into an account for the pixel super-resolution reconstruction, it is required to measure the changes in the angle of orientation of the cells in each frame. We used two approaches to measure the angle of a cell; one approach calculates the angle from the shape of the cells in the

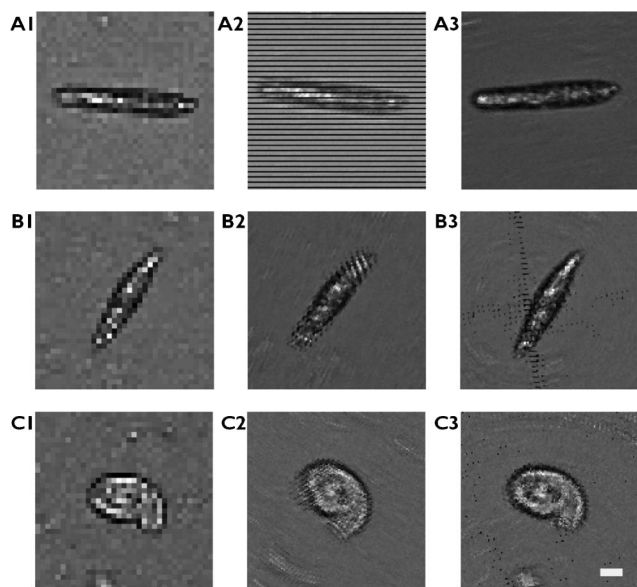


Fig. 3 Artifacts in image reconstruction and improved images. (A) Cells moving in horizontal direction show missing information in y direction (A2) and a different sequence of the same cell with adequate motion vector showing a well-reconstructed image (A3). (B–C) Cells with large rotational movement show artifacts at the two ends of the cell (B2, C2). Reconstructed image with rotation compensation (B3, C3). The scale bar indicates $10\ \mu\text{m}$.

low resolution images and the other approach calculates it from the displacement of cells between each subsequent frames. For the elongated microorganisms, the rotation angle can be easily measured directly from the raw images (Fig. 3(B), Fig. S3†). For the round cells, we can use the fact that the cells are directed towards the direction of motion and calculate the angle from the motion vector (Fig. 3(C), Fig. S4†). The rotational motion of the microorganisms can be compensated by re-rotating each frame by the angle $-\theta_k$ during the reconstruction process. This rotation process can be described as below.

$$i_{kr}(I_x, I_y) = i_k(I_x \cos \theta_k - I_y \sin \theta_k + nd_{kx}, I_x \sin \theta_k + I_y \cos \theta_k + nd_{ky}) \quad (2)$$

where i_k represents k^{th} low-resolution image up-sampled by the factor of n and i_{kr} represents k^{th} low resolution image up-sampled by n , shifted and rotated for reconstruction. With the same set of low resolution sequences used for Fig. 3(B2) and 3(C2), we can reconstruct rotation-compensated images (Fig. 3(B3) and 3(C3)) where the stripe-patterned artefacts are effectively removed. However, it is worth noting that the image blur caused by the cell rotation perpendicular to the axis of translation cannot be computationally removed. We minimize this rotation-induced blur by maximizing the frame rate such that the cell rotation within the low resolution sequence is negligible.

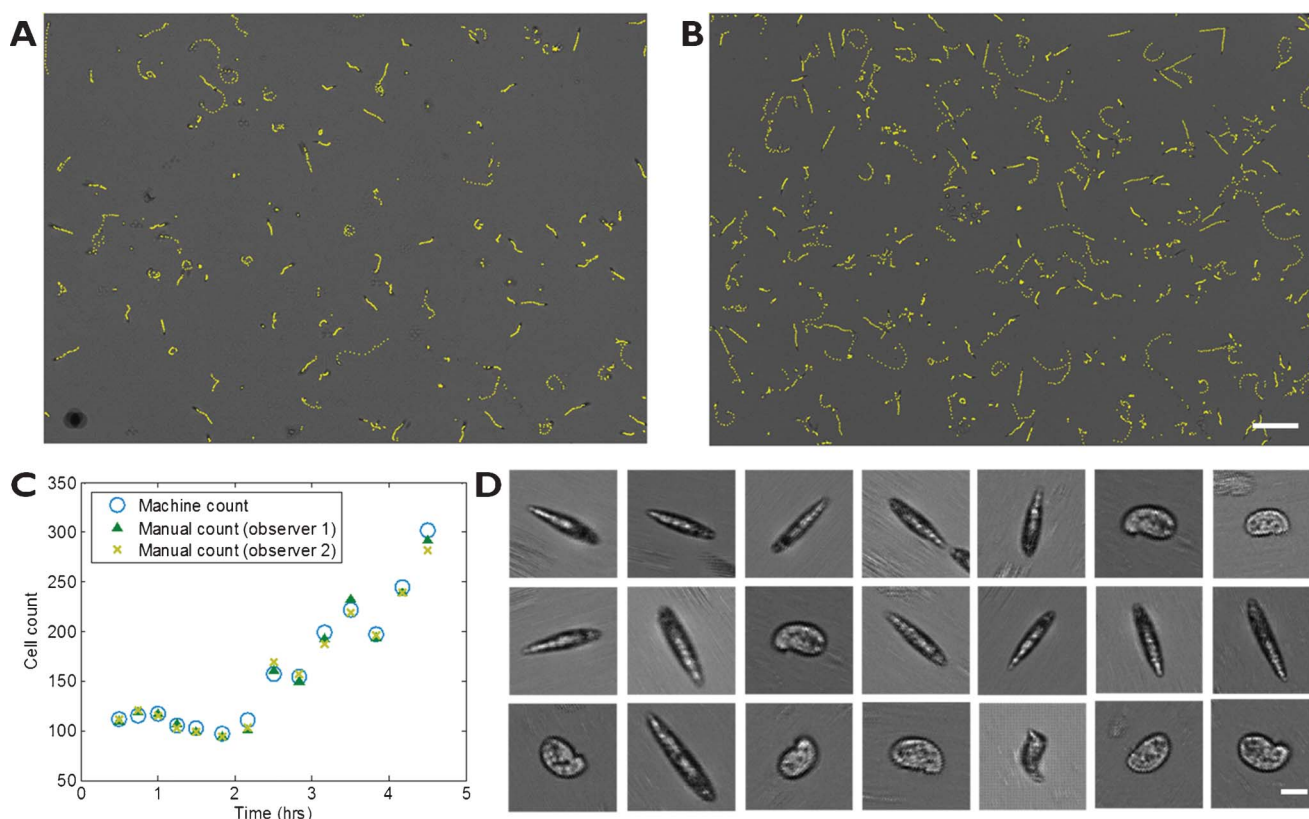


Fig. 4 Longitudinal imaging of *Euglena gracilis* on ePetri dish. Large FOV images of *Euglena gracilis* at (a) 1 h and (b) 4.5 h, respectively. Yellow line shows traces of each cell for 1 s. Scale bar $200\ \mu\text{m}$. (c) Growth in cell population counted with our cell counting software (blue) and manually counted (green, yellow). (d) High resolution images of *Euglena gracilis* reconstructed with SPMM method (selected). Scale bar indicates $20\ \mu\text{m}$.

Longitudinal imaging of *Euglena gracilis*

To demonstrate longitudinal imaging and motion analysis capabilities, we cultured *Euglena gracilis* protist in our ePetri system. Long term culture monitoring and image-based motion analysis with ePetri system can benefit many biological assays and scientific studies using *Euglena gracilis*. Due to their sensitivity to the changes in the chemical environment, *Euglena gracilis* is used for various microbiological assays, including vitamin B12 assay²³ and aquatic toxicity assays.^{7,13,24} Their phototactic and gravitactic movement has been intensively studied and utilized as sensors.^{10,25} Their shape also changes under different illumination conditions.⁶

Euglena cells were cultured in the ePetri platform and continuously imaged for 10 h. We used two image acquisition modes; large field-of-view imaging with low frame rate (16 fps) for continuous monitoring and smaller field-of-view scanning with a higher frame rate (180 fps) for SPMM high resolution image reconstruction. From the large FOV image, statistical analysis on cell counting, tracking (speed and trajectory), size and shape distribution (aspect ratio of the cells) can be obtained for >200 cells. For a closer look into each cell, we performed SPMM imaging by dividing the entire imaging area into smaller windows (400 × 200 pixels) to obtain sequences with higher temporal resolution.

Fig. 4a shows growth of *Euglena* cells cultured in an ePetri system. 5 μL of *Euglena* medium containing 140 cells were dispensed into an ePetri chip and cultured at room temperature with a green LED illumination (0.2 W m⁻²). Two snapshots of the culture at 1 h and 5 h show distinct increases in the cell population in the culture. High-resolution images using the SPMM method reveal details in the shape, size and opacity variations in each microorganisms (Fig. 4d). We observed *Euglena gracilis* in various stages of growth including young *Euglena* cells with elongated shapes, old rounded cells, cells undergoing mitosis and non-motile palmella stage cells that are covered with mucilage. However, since the SPMM method relies on the movement of the target cells, images of immotile cells such as mitotic, palmella stage and lysed cells cannot be reconstructed with enhanced resolution.

Image analysis for cell counting was performed by first identifying each cell *via* thresholding the raw gray scale images (*Euglena* cells or the boundary of *Euglena* cells appear darker than the background), converting into binary images and then counting the number of areas within a predefined size range to filter out debris and noises. From the counting results, the growth of the cells showed a clear increase with the number of cells doubling in approximately 5 h. To assure the accuracy of our cell counting software, our results were compared with manual counting results of 2 observers. The results showed the average percent difference of 3.0% between the manual and the machine count, where percent difference between the observers is 3.1%. (See table S1†). Subsequent analysis for cell tracking and shape analysis was performed with simple image processing techniques. We traced each cells for 16 frames (1 s) to obtain the speed of the cells, which is an important parameter for the status of the culture. The mean velocity of the cells remained within 10% variance during the 5 h culture, indicating that the culture environment in ePetri dish was adequate.

Next we performed differential experiment with two sets of *Euglena* cultures. We used spring water as medium for culture A

and the *Euglena* culture medium for culture B. The two cultures were kept in ePetri for >5 h under same temperature and illumination conditions. We used our analysis software to obtain the distribution of the motility and the cell shapes (Fig. 5). The result shows that the cells cultured in fresh water have more population of elongated cell types with high aspect ratio and lower motility, whereas larger population of *Euglena gracilis* cultured in the medium showed faster motion and rounded shapes.

Conclusions

We demonstrated the imaging of motile microorganisms in an ePetri platform by using the inherent motion of the cells for the pixel super-resolution reconstruction. Rototranslatory movement of *Euglena gracilis* allows capturing of sequential shadow

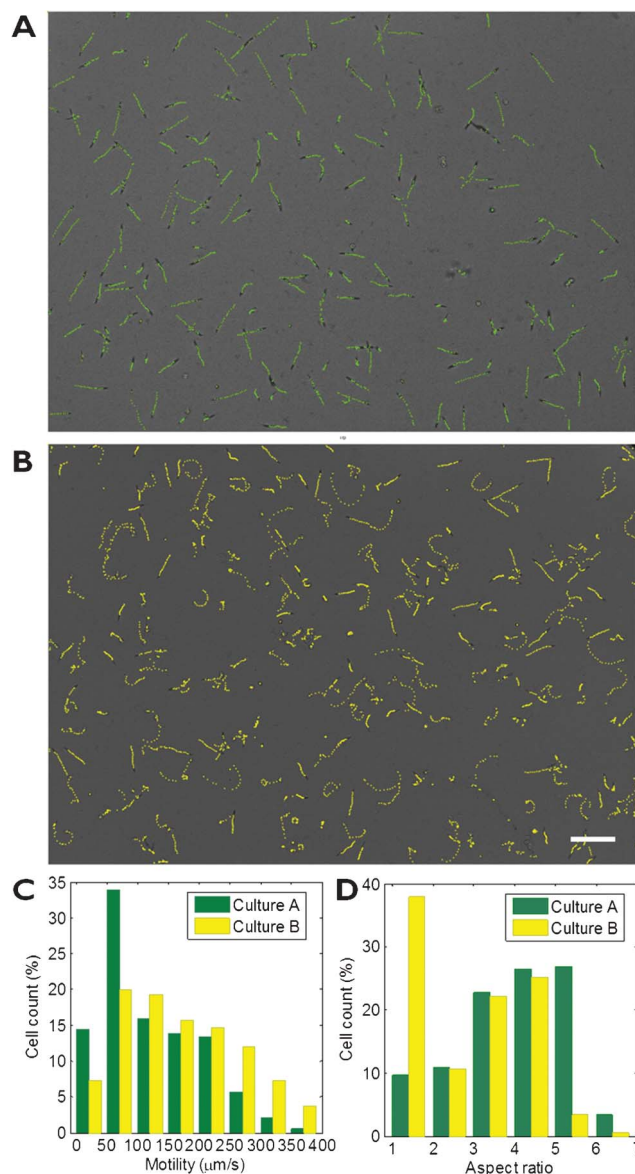


Fig. 5 Motion tracking and shape analysis using ePetri. (a–b) *Euglenas* cultured in spring water (a) and in *Euglena* medium. Scale bar indicates 200 μm. (b) Showing difference in their motion and the cell sizes. (c) Motility distribution of the cells. (d) Aspect ratio (major axis/minor axis) distribution of the cells.

images with sub-pixel shifts, which allows for the imaging of individual cells with enhanced resolution compared to the pixel size of the sensors. Using the SPMM method, we achieved high resolution imaging over 6 mm × 4 mm area without using any optical elements. In addition, we show long-term culture of *Euglena gracilis* in a SPMM ePetri platform, and demonstrate image-based analysis for automatic cell counting, motion analysis and shape analysis.

This work, particularly focused on a motile microorganism, shows the versatility of our SPMM ePetri system in the choice of the sample and the type of information collectible through our system. The key advantage of our SPMM ePetri platform is that it reduces the labour required and the cost of culture experiments by miniaturizing and automating the monitoring system. The user can simply dispense the specimen on an ePetri chip and monitor the growth through our software with the minimum access to the culture. It can also improve the quality of experiments a by improving the temporal resolution of the monitoring and allowing for streamlined and parallel process.

Experimental

Preparation of ePetri chip

We used MT9P031 (2.2 µm pixel, Aptina) image sensors. We removed the colour filter and the microlens layer by treating the sensor under oxygen plasma for 10 min (120 W). The PDMS wall is prepared by mixing 1 : 10 with base and curing agent, then spin coated on a 3 in. silicon wafer followed by baking at 80 °C for 1 h. The surfaces of ePetri chips are treated with poly-L-lysine (0.01%, Sigma-Aldrich) for 15 min and washed with distilled water three times.

Euglena gracilis culture

Euglena gracilis (Carolina scientific) were cultured in a custom made *Euglena* medium (spring water, 40 L⁻¹ wheat grains, 35 L⁻¹ rice grains and 5 cm³ L⁻¹ dried skimmed milk, recipe provided by Carolina scientific) under natural sunlight illumination. All experimentation was performed at room temperature where the *Euglena gracilis* are healthy and their reproduction is well promoted.

Acknowledgements

We acknowledge funding support from the National Institute of Health (NIH) Agency Award R01AI096226-01.

References

- 1 C. Ascoli, M. Barbi, C. Frediani and A. Murč, *Biophys. J.*, 1978, **24**, 585.
- 2 T. Höpfner, A. Bluma, G. Rudolph, P. Lindner and T. Scheper, *Bioprocess Biosyst. Eng.*, 2010, **33**, 247.
- 3 C. M. Waterman-Storer, A. Desai, J. Chloe Bulinski and E. D. Salmon, *Curr. Biol.*, 1998, **8**, 1227.
- 4 Y. Aizu and T. Asakura, *Opt. Laser Technol.*, 1991, **23**, 205.
- 5 M. Tim, *J. Immunol. Methods*, 1983, **65**, 55.
- 6 D.-P. Häder and K. Vogel, *J. Math. Biol.*, 1991, **30**, 63.
- 7 H. Tahedl and D.-P. Häder, *Ecotoxicol. Environ. Saf.*, 2001, **48**, 161.
- 8 I. Levin-Reisman, O. Gefen, O. Fridman, I. Ronin, D. Shwa, H. Sheftel and N. Q. Balaban, *Nat. Methods*, 2010, **7**, 737.
- 9 G. Wei, P. Cosman, C. C. Berry, F. Zhaoyang and W. R. Schafer, *IEEE Trans. Biomed. Eng.*, 2004, **51**, 1811.
- 10 P. R. Richter, M. Schuster, H. Wagner, M. Lebert and D. P. Häder, *J. Plant Physiol.*, 2002, **159**, 181.
- 11 L. E. Waggoner, G. T. Zhou, R. W. Schafer and W. R. Schafer, *Neuron*, 1998, **21**, 203.
- 12 T.-W. Su, A. Erlinger, D. Tseng and A. Ozcan, *Anal. Chem.*, 2010, **82**, 8307.
- 13 H. Ahmed and D.-P. Häder, *J. Appl. Phycol.*, 2010, **22**, 785.
- 14 D. N. Breslauer, R. N. Maamari, N. A. Switz, W. A. Lam and D. A. Fletcher, *PLoS One*, 2009, **4**, e6320.
- 15 X. Cui, L. M. Lee, X. Heng, W. Zhong, P. W. Sternberg, D. Psaltis and C. Yang, *Proc. Natl. Acad. Sci. U. S. A.*, 2008, **105**, 10670.
- 16 G. Zheng, S. A. Lee, S. Yang and C. Yang, *Lab Chip*, 2010, **10**, 3125.
- 17 G. Zheng, S. A. Lee, Y. Antebi, M. B. Elowitz and C. Yang, *Proc. Natl. Acad. Sci. U. S. A.*, 2011, **108**, 16889.
- 18 S. Farsiu, M. Robinson, M. Elad and P. Milanfar, *IEEE Trans. Image Process.*, 2004, **13**, 1327.
- 19 P. Sung Cheol, P. Min Kyu and K. Moon Gi, *IEEE Signal Process. Mag.*, 2003, **20**, 21.
- 20 C. Ascoli, M. Barbi, C. Frediani and D. Petracchi, *Biophys. J.*, 1978, **24**, 601.
- 21 M. Cramer and J. Myers, *Arch. Microbiol.*, 1952, **17**, 384.
- 22 S. A. Lee, R. Leitao, G. Zheng, S. Yang, A. Rodriguez and C. Yang, *PLoS One*, 2011, **6**, e26127.
- 23 G. Ross, *J. Clin. Pathol.*, 1952, **5**, 250.
- 24 K. Aoyama, K. Iwahori and N. Miyata, *Mutat. Res., Genet. Toxicol. Environ. Mutagen.*, 2003, **538**, 155.
- 25 K. Ozasa, J. Lee, S. Song, M. Hara and M. Maeda, *Lab Chip*, 2011, **11**, 1933.