Trapping and Manipulation of Single Cells in Crowded Environments

Qian Zhao1,2, Hao-Wei Wang1*, Pan-Pan Yu1, Shu-He Zhang3, Jin-Hua Zhou3, Yin-Mei Li1,4 and Lei Gong1*

1 Department of Optics and Optical Engineering, University of Science and Technology of China, Hefei, China, 2 Shandong Provincial Engineering and Technical Center of Light Manipulations and Shandong Provincial Key Laboratory of Optics and Photonic Device, School of Physics and Electronics, Shandong Normal University, Jinan, China, 3 Department of Biomedical Engineering, Anhui Medical University, Hefei, China, 4 Hefei National Laboratory for Physical Sciences at the Microscale, University of Science and Technology of China, Hefei, China

Optical tweezers provide a powerful tool to trap and manipulate living cells, which is expected to help people gain physiological insights at single-cell level. However, trapping and manipulating single cells under crowded environments, such as blood vessels and lymph nodes, is still a challenging task. To overcome this issue, an annular beam formed by the far-field Bessel beam is introduced to serve as an optical shield to isolate the target cells from being disturbed. With this scheme, we successfully trapped and manipulated single blood cells in a crowded environment. Furthermore, we demonstrated manipulation of two lymphocytes ejected from a lymph node independently with dual-trap optical tweezers, which paves the way for exploring cell interactions under living conditions. Such technique might be helpful in the study of how natural killer cells response to virus-infected cells or cancer cells.

Keywords: optical tweezers, crowded environment, lymphocytes, optical shield, optical force

INTRODUCTION

Since the pioneer work of Ashkin et al. (1986), optical tweezers have emerged as an essential tool for manipulating single cells and performing biomechanical characterizations at microscopic level (Chen et al., 2007; Zhong et al., 2013b; Xie et al., 2018). Distinct advantages of using tweezers for these characterizations include non-contact cell manipulation, Piconewton force accuracy (Williams, 2002; Falleroni et al., 2018), and amiability to liquid medium environments (Zhang and Liu, 2008). The wide application range of optical tweezers, such as transporting foreign materials into single cells (Hou et al., 2016; Johansen et al., 2016), delivering cells to specific locations (Xie et al., 2018), sorting cells in microfluidic systems (Wang et al., 2011; Landenberger et al., 2012), and quantifying intercellular interactions (Chen et al., 2007; Hongliang et al., 2008), have been proved previously. However, researches readily performed in vitro may not reflect the biological activities in vivo accurately because of the complexity of the in vivo environment. Recently, trapping, and manipulation of cells within living animals has been achieved using infrared optical tweezers (Zhong et al., 2013a; Johansen et al., 2016). Notably, a non-contact micro-operation has been managed to clear a blocked capillary in a living mice (Zhong et al., 2013a). Therefore, the ability of manipulating single cells in vivo is urgently demanded to verify our knowledge acquired from the in vitro studies.

Optical tweezers is capable of probing the physical properties of cells in living animals with microscale resolution (Paul et al., 2019), which might help people gain physiological insights...
at single-cell level. A recent work (Johansen et al., 2016) performed a unique way to stimulate immune response by optically trapping and manipulating injected bacteria inside a living zebrafish. Although there is no unconquerable obstacle for optical trapping inside living animals, isolating single cells under crowded environments remains a challenging task. When optically manipulating in vitro (Xie et al., 2002; Sinjab et al., 2018), samples are usually diluted to sparsely populated level so that the individual cells can be manipulated, and measured without disturbances from the ambient ones. However, living environments are usually congested with various classes of cells, and ambient cells enter the trap consequently and frequently. Thus, the tracking procedure is disturbed and the accuracy of the measurement is reduced. For example, the crowded environment inside the lymph node (Stoll et al., 2002; Willard-Mack, 2006) might be the most important obstacle to optical tweezers studies in this critical small gland related to infection and cancer development. In brief, the crowded environments inside living animals become a critical challenge for manipulating single cells in situ.

Trapping with structured light beams is the frontier of optical tweezers. For example, the non-diffraction Bessel beam could overcome the limited depth of focus of Gaussian beam (Durnin et al., 1987). Bottle beams could help trap absorbing particles that cannot be stably confined by conventional optical tweezers (Gong et al., 2016). In this work, we trapped and manipulated single cells in the crowded environment with engineered trapping laser beam. Serving as an optical shield, an annular beam is introduced to avoid the disturbance from crowded environments during single cell manipulation. We adopt an optical configuration using a diffraction axicon and a converging lens to create the high efficiency annular beam. The established optical shield can be regulated by adjusting the distance between the axicon and the lens. Moreover, a theoretical analysis of the optical forces is presented. According to our analysis, the scattering force is stronger than the gradient force, and particles are propelled away from the center in the beam direction. As a proof of concept, we first trapped single blood cells within a crowded environment. Next, we applied the method to manipulate individual lymphocytes from a lymph node inside its normal physiological status. Our work is expected to benefit the research of the lymphocytes related immune response in the lymph nodes by keeping the natural physiological environment.

**MATERIALS AND METHODS**

**Principle of Generating Optical Shield**

To produce an optical shield, a hollow beam is constructed by an axicon and a lens as shown in Figure 1. When an aperture plane wave incidents on an axicon, a Bessel beam in the range $Z_{\text{max}} = a/(n - 1)\gamma$ can be created under the paraxial approximation, where $\gamma$ is the angle between the conical surface and the flat surface, $a$ is the semi-diameter of the aperture plane wave, $n$ is the index of the axicon. According to the focusing properties of aperture Bessel beam (Wei et al., 2005), as presented in Figure 1A, the hollow beam can be produced behind the lens when $z_0 < f < Z_{\text{max}}$, where $z_0$ is the distance between axicon and lens, $f$ is the focal length of the lens. We calculate the total force vector indicated along red arrows as a function of a particle's position over $x$-$y$ plane using ray optics model (Zhou et al., 2008; Shao et al., 2019). In the calculation, the particle (radius = 2.5 $\mu$m, $n_p = 1.50$) is immersed in water ($n_w = 1.33$), and other simulation parameters include $\lambda = 1064$ nm, $f = 260\lambda$, $z_0 = 0.6f$, and $\gamma = \pi/18$ were applied also. Details of trapping efficiency calculation are described in Supplementary Material. A bright annular intensity distribution is tightly focused on the focal plane of the lens ($z = 0$), as shown in Figure 1B. The total force vectors indicated by the red arrows are presented in $x$-$y$ plane as a function of particle's position. The length and the direction of arrows represent the magnitude and direction of the total force, respectively. Figures 1C,D show the transversal trapping efficiency and longitudinal trapping efficiency for the particle as a function of particle's displacement along $x$ axis, respectively. The blue dash line indicates the corresponding one-dimensional normalized intensity distribution. The distribution of the force arrows and the two trapping efficiency curves illustrate the manipulation capability of the annular beam. When the particle is initially located in the inner part of the annulus and subjected to a drag force outward in the radial direction, it will shift to the bright annular intensity where the transversal trapping efficiency and longitudinal trapping efficiency are balanced. Noted that the longitudinal optical force can be balanced by the gravity of the particle. Therefore, the hollow beam can clear a blank area in which the trapped individual cell can be shielded.

**Experimental Setup**

Our experimental setup is shown in Figure 2. A light beam from a 1064-nm continuous-wave (CW) laser was separated into two parts with a polarizer beam splitter (PBS1). The p-polarized beam passed through an axicon ($\gamma = 0.5^\circ$, Thorlabs) to produce a Bessel-like beam. Then, the Bessel-like beam was focused by lens L1 to create an annular beam, which acts as an optical shield. The s-portion was expanded by a beam expander and finally focused by an objective lens to create an optical trap. In addition, a 780-nm CW laser serves as the light source for another optical trap. The three beam paths were combined together by PBS2 and a dichroic mirror (DM1). Another dichroic mirror (DM2) reflects the combined beams toward the back focal plane of the objective. After the beams propagate through a tube lens, a 100 $\times$ oil immersion objective lens (NA = 1.4, Olympus) highly focus them to create desired optical traps. To move the positions of the two focused optical spots, scanning mirrors based on the high-speed Piezo tip/tilt platforms (S-330.30, Physik Instrumente) were adopted to control the beam incidence. The state of optical tweezers can be switched “on” and “off” independently with shutters. As a result, an annular intensity distribution, and two controllable optical traps formed at the focal plane of the objective. The laser power used for 1064-nm light is higher than that of 780-nm light in order to provide enough power to push cells away. The laser powers of the optical traps are ~120 mW (1064 nm), ~120 mW (1064 nm), and ~42 mW (780 nm). The whole optical tweezer setup was built on an Olympus IX71 inverted microscope, and real-time imaging of
the trapping process can be readily realized with the microscope and a CCD camera.

**Sample Preparation**

The samples used in the experiments are mice blood and an inguinal lymph node of mice. Mice were obtained from the Experimental Animal Center of University of Science and Technology of China. Mice blood was extracted from tail and diluted about 100 times with the isotonic Phosphate-buffered saline (PBS) buffer before injected into a glass chamber. Lymphocytes used in trapping experiment were ejected from the inguinal lymph node. An illustration cartoon (Supplementary Figure 1) was presented to illustrate procedure in detail. First of all, a lymph node was peeled off from mice and put into a chamber containing isotonic PBS. Then a cover glass was put on top of the lymph node to make it cling to the bottom of the chamber. Finally, we slightly pressed the cover glass so that the lymphocytes were ejected out of the lymph node via the efferent lymphatic. Since lymphocytes just left the lymph node, the biological activity and physiological status were well preserved (Miller et al., 2002; Stoll et al., 2002).

**RESULTS**

**Trapping a Single Blood Cell in a Crowd**

We first tested the ability of our scheme for trapping single cells within a crowded environment using mice blood cells. Figure 3 shows the process of manipulating individual blood cells under crowded environment. At the very beginning (t = 0 s), there is a crowd of cells in the field of view. When the hollow beam was switched on, cells around the beam center were pushed outward to the high-intensity ring because of optical forces. After 58 s, an optical shield with many cells distributing around the circumference was established. As a result, there exists a blank area on the center of optical shield. Then, an additional optical trap was used to capture and manipulate a single blood cell. A leukocyte (highlighted by the white circle) was trapped and moved to the center of the blank area via active manipulation, in which the optical trap is moved by tilting the scanning mirror, as shown in Figure 3b at the moment of 64 s. At the same time, another red blood cell highlighted by the white circle in Figures 3c-e, was trapped and moved by another optical trap toward the first trapped cell (Figure 3e).
Thus, two cells were isolated from the crowd and contacted with each other (Figure 3f).

**Manipulating Single Live Lymphocytes From a Lymph Node**

We applied the proposed scheme for trapping individual lymphocytes from an inguinal lymph node. Direct optical trapping of lymphocytes inside the lymph node is difficult because the wide-field microscope is not practicable through the thick capsule of the lymph node. Therefore, the lymphocytes were first pushed out of the lymph node via the efferent lymphatic, and selected lymphocytes were trapped and isolated from a crowd in the area around the efferent lymphatic. This implementation mimics the environment inside the lymph node because the physiological status of lymphocytes did not change as they just left the lymph node. Figure 4a shows a picture of a crowd of lymphocytes. Firstly, we switched on the optical shield, which gradually pushed the cells out forward and created an empty area. Then a selected lymphocyte (highlighted by a white circle in Figure 4c) was trapped and moved to the center of the shield with a Gaussian trap, as shown in Figures 4b–d. The relative motion of optical tweezers is achieved by tilting the incident angle of the laser beam. Additionally, the target cell and the optical shield can be moved to a new position via stage motion, as illustrated in Figures 4d–f, where the white cross indicates the reference point.

Furthermore, we demonstrated manipulation of two lymphocytes ejected from a lymph node independently with dual-trap optical tweezers. As shown in Figures 4g–i, two lymphocytes highlighted by white circles are trapped and pulled together at the center of the blank area in sequence, which perfectly mimic exploring cell interactions at cellular level, and such as the responses of natural killer cells to virus-infected cells or cancer cells.

**DISCUSSION**

In conclusion, we have proposed and implemented an optical shield scheme for manipulating individual cells in a crowded environment. An adjustable optical shield was created by using a diffraction axicon and a converging lens. Optical forces exerted on micro-particles by the optical shield were calculated to analyze its capability for optical clearing. Trapping and manipulating individual living lymphocytes from a lymph node was achieved...
with our proposed scheme in experiment. Furthermore, we have demonstrated independent manipulation of two lymphocytes with dual-trap optical tweezers, which could benefit studies of cell interactions in a crowded environment.

Lymph nodes are critical small glands closely related to infection and cancer development for mammals (Willard-Mack, 2006). Trapping and manipulation of individual cells inside the lymph nodes is expected to help the study of how natural killer cells response to virus-infected cells or cancer cells. In our current experiment, trapping lymphocytes was actually performed in the saline solution, and trapping depth through the lymph node capsule is limited to about 100 μm due to the short working distance of the objective. A water immersion objective with longer working distance should be helpful to improve the trapping depth (Zhong et al., 2013b). However, the wavefront distortion caused by the thick capsule and limited imaging depth of the microscope are still obstacles of optical trapping inside the lymph node. Stripping the capsular surface of the lymph node or wavefront correction could be helpful in deep imaging and trapping. Because the inability of wide-field microscope in distinguishing T-cells and B-cells, fluorescence labeling technologies is demanded to recognize different kinds of cells, and which may promote the screening of immunological interactions (Stoll et al., 2002). In particular, our proposed trapping scheme can be combined with some other techniques such as Raman spectroscopy (Sinjab et al., 2018), position tracking (McAlinden et al., 2014), and imaging (Berg and Holler, 2016) for advanced studies of individual live cells, where the optical shield will enable measurements without disturbances.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this manuscript will be made available by the authors, without undue reservation, to any qualified researcher.

REFERENCES

Ashkin, A., Dziedzic, J., Bjorkholm, J., and Chu, S. (1986). Observation of a single-beam gradient force optical trap for dielectric particles. Opt. Lett. 11, 288–290. doi: 10.1364/ol.22.000816
Berg, M. J., and Holler, S. (2016). Simultaneous holographic imaging and light-scattering pattern measurement of individual microparticles. Opt. Lett. 41, 3363–3366. doi: 10.1364/OL.41.003363
Chen, H., Ge, K., Li, Y., Wu, J., Gu, Y., Wei, H., et al. (2007). Application of optical tweezers in the research of molecular interaction between lymphocyte function associated antigen-1 and its monoclonal antibody. Cell Mol. Immunol. 4, 221–225.
Durnin, J., Miceli, J., Jr., and Eberly, J. (1987). Diffraction-free beams. Phys. Rev. Lett. 58, 1499–1501.
Fallon, R., Torre, V., and Cojoc, D. (2018). Cell mechanotransduction with piconewton forces applied by optical tweezers. Front. Cell. Neurosci. 12:130. doi: 10.3389/fncel.2018.00130
Gong, L., Liu, W., Zhao, Q., Ren, Y., Qiu, X., Zhong, M., et al. (2016). Controllable light capsules employing modified Bessel-Gauss beams. Sci. Rep. 6:29001. doi: 10.1038/srep29001
Hongliang, R., Lihui, Z., and Yinmei, L. (2008). Measurement of interaction potential between colloidal particles using dual optical tweezers. Chin. J. Lasers 35, 151–155.

ETHICS STATEMENT

The animal study was reviewed and approved by The Ethical Committee of the University of Science and Technology of China. Written informed consent was obtained from the owners for the participation of their animals in this study.

AUTHOR CONTRIBUTIONS

LG, QZ, and H-WW conceived the idea. QZ performed the experiment, analyzed the data. P-PY contributed to constructing the experimental setup. QZ, H-WW, and LG discussed the results. QZ and LG wrote and edited the manuscript. S-HZ and J-HZ contributed to the force calculation. Y-ML supervised the study. All the authors reviewed and revised the manuscript.

FUNDING

National Science Foundation of China (NSFC; 61535011, 11704369, 11974333, and 91950104). The National Key Research and Development Program of China (2017YFA0304800).

ACKNOWLEDGMENTS

We would like to acknowledge Ziqiang Wang and Yifan Liu for helpful discussions.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fbioe.2020.00422/full#supplementary-material

Hou, X., DeSantis, M. C., Tian, C., and Cheng, W. (2016). Optical manipulation of a single human virus for study of viral-cell interactions. Proc. SPIE Int. Soc. Opt. Eng. 9922:992212. doi: 10.1117/12.2239051
Johansen, P. L., Fenaroli, F., Evensen, L., Griffiths, G., and Koster, G. (2016). Optical micromanipulation of nanoparticles and cells inside living zebrafish. Nat. Commun. 7:10974. doi: 10.1038/ncomms10974
Landenberger, B., Hofemann, H., Wadle, S., and Rohrbach, A. (2012). Microfluidic sorting of arbitrary cells with dynamic optical tweezers. Lab. Chip 12, 3177–3183. doi: 10.1039/c2lc21099a
McAlinden, N., Glass, D. G., Millington, O. R., and Wright, A. J. (2014). Accurate position tracking of optically trapped live cells. Biomed. Opt. Express 5, 1026–1037. doi: 10.1364/BOE.5.001026
Miller, M. J., Wei, S. H., Parker, I., and Cahalan, M. D. (2002). Two-photon imaging of lymphocyte motility and antigen response in intact lymph node. Science 296, 1869–1873. doi: 10.1126/science.1070051
Paul, C. D., Bishop, K., Devine, A., Paine, E. L., Staunton, J. R., Thomas, S. M., et al. (2019). Tissue architectural cues drive organ targeting of tumor cells in zebrafish. Cell Syst. 9, 187–206. doi: 10.1016/j.cels.2019.07.005
Shao, M., Zhang, S., Zhou, J., and Ren, Y.-X. (2019). Calculation of optical forces for arbitrary light beams using the Fourier ray method. Opt. Express 27, 27459–27476. doi: 10.1364/OE.27.027459
Sinjab, F., Awuah, D., Gibson, G., Padgett, M., Ghaemmaghami, A. M., and Nottenberg, I. (2018). Holographic optical trapping Raman micro-spectroscopy
for non-invasive measurement and manipulation of live cells. *Opt. Express* 26, 25211–25225. doi: 10.1364/OE.26.025211

Stoll, S., Delon, J., Brotz, T. M., and Germain, R. N. (2002). Dynamic imaging of T cell-dendritic cell interactions in lymph nodes. *Science* 296, 1873–1876. doi: 10.1126/science.1071065

Wang, X., Chen, S., Kong, M., Wang, Z., Costa, K. D., Li, R. A., et al. (2011). Enhanced cell sorting and manipulation with combined optical tweezer and microfluidic chip technologies. *Lab. Chip* 11, 3656–3662. doi: 10.1039/c1lc20633b

Wei, M.-D., Shiao, W.-L., and Lin, Y.-T. (2005). Adjustable generation of bottle and hollow beams using an axicon. *Opt. Commun.* 248, 7–14. doi: 10.1364/OL.39.000630

Willard-Mack, C. L. (2006). Normal structure, function, and histology of lymph nodes. *Toxicol. Pathol.* 34, 409–424. doi: 10.1080/01926230600867727

Williams, M.C. (2002). *Optical Tweezers: Measuring Piconewton Forces*. Biophysics Textbook. Available Online: http://www.biophysics.org/btol

Xie, C., Dinno, M. A., and Li, Y. (2002). Near-infrared Raman spectroscopy of single optically trapped biological cells. *Opt. Lett.* 27, 249–251. doi: 10.1364/ol.27.000249

Xie, M., Shakoor, A., and Wu, C. (2018). Manipulation of biological cells using a robot-aided optical tweezers system. *Micromachines* 9:245. doi: 10.3390/mi9050245

Zhang, H., and Liu, K.-K. (2008). Optical tweezers for single cells. *J. R. Soc Interface* 5, 671–690. doi: 10.1098/rsif.2008.0052

Zhong, M.-C., Gong, L., Zhou, J. H., Wang, Z. Q., and Li, Y. M. (2013a). Optical trapping of red blood cells in living animals with a water immersion objective. *Opt. Lett.* 38, 5134–5137. doi: 10.1364/OL.38.005134

Zhong, M.-C., Wei, X.-B., Zhou, J.-H., Wang, Z.-Q., and Li, Y.-M. (2013b). Trapping red blood cells in living animals using optical tweezers. *Nat. Commun.* 4:1768. doi: 10.1038/ncomms2786

Zhou, J.-H., Ren, H.-L., Cai, J., and Li, Y.-M. (2008). Ray-tracing methodology: application of spatial analytic geometry in the ray-optic model of optical tweezers. *Appl. Opt.* 47, 6307–6314. doi: 10.1364/ao.47.006307

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2020 Zhao, Wang, Yu, Zhang, Zhou, Li and Gong. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.