Research Article

A Structural Study of *Escherichia coli* Cells Using an *In Situ* Liquid Chamber TEM Technology

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Received 29 September 2014; Revised 8 January 2015; Accepted 13 January 2015

Academic Editor: Chih-Ching Huang

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Studying cell microstructures and their behaviors under living conditions has been a challenging subject in microbiology. In this work, *in situ* liquid chamber TEM was used to study structures of *Escherichia coli* cells in aqueous solutions at a nanometer-scale resolution. Most of the cells remained intact under electron beam irradiation, and nanoscale structures were observed during the TEM imaging. The analysis revealed structures of pili surrounding the *E. coli* cells; the movements of the pili in the liquid were also observed during the *in situ* tests. This technology also allowed the observation of features of the nucleoid in the *E. coli* cells. Overall, *in situ* TEM can be applied as a valuable tool to study real-time microscopic structures and processes in microbial cells residing in native aqueous solutions.

1. Introduction

High-resolution real-time observation of cells and cellular processes in their liquid environments is crucial for revealing cellular structures and functions. Electron microscopes (EMs) can easily achieve a nanometer level resolution [1]; therefore, EMs are efficient tools for observing cellular structures. However, traditional sampling procedures limit the application of EMs for real-time observation of liquid samples. Because of the high-vacuum operating conditions, extensive sample preparations (e.g., fixation, metal staining, plastic embedding and slicing [2], or freezing) are needed for biological TEM samples [3]. An in-depth understanding of cellular structures and intracellular processes requires real-time imaging of the entire biological object with high spatial resolutions in the native liquid environment. To allow real-time and fast observations of dynamic processes occurring in biological objects, the samples must be fully hydrated and nonfrozen. Environmental TEM uses a differential pump technology and therefore allows cellular studies under partially hydrated conditions [4]; however, this process is not conducted in pristine liquid environments [5, 6].

*In situ* liquid chamber TEM has been developed recently for nanostuctural systems; this enables the user to image and monitor biological samples in fully hydrated environments, thus providing real-time dynamic information [7]. In this method, two ultrathin electron-transparent window chips are used to construct a liquid cell chamber; the liquid sample is sealed between these chips. The *in situ* liquid cells can be used in standard TEM instruments [8]. A variety of samples and processes of nanomaterials have been studied with this technique, including nanomaterial reactions [9] and motion [10], nanocrystal depositions [11, 12], and self-assembly of nanolipoprotein discs [13]. Biological cells have also been imaged in liquid with nanometer resolution, such as human leukocytes [14], COS7 fibroblast cells [15], yeast cells [16], *K. pneumoniae* CG43S3 cells [7], and the bacteria *Deinococcus radiodurans* (*D. radiodurans*) [17]. *E. coli* has also been investigated previously [18]. Typically, a specimen kit equipped with *SiO₂* nanomembranes is applied in TEM...
imaging. This kit enables the observation of living *E. coli* [7]. *E. coli* has been reported to survive this TEM imaging process at 2.8 s under a 200 KeV electron beam [7]. Peckys et al. imaged *E. coli* [18] labeled with gold nanoparticles in liquid with scanning transmission electron microscopy (STEM). However, they only showed the overall morphology and viability of *E. coli* cells with limited resolution. The stability of the cells under the electron beam, the resolution, and the dynamic observation are key issues that need to be further addressed.

This study focuses on the fine structural analysis and dynamic observation of *E. coli* cells using *in situ* liquid chamber TEM. The integrity, stability, and viability of the cells under electron beams, which have not been examined in previous studies, are also examined.

### 2. Materials and Methods

*Escherichia coli* BL21(DE3) was grown in Luria-Bertani (LB) broths at 37°C for 12 h and then stored at 4°C before being used. The liquid chamber used for the *in situ* TEM was developed following a previously described procedure [9–11, 18]. In brief, silicon chips with 50 nm thick Si₃N₄ membranes (Ted Pella, Inc., CA, USA) were used as the electron transparent windows for the liquid chamber. The size of the windows was 0.5 × 0.5 mm. A JEOL JEM 2010 TEM (JEOL Ltd., Tokyo, Japan) was used for the *in situ* TEM and was operated under a 200 kV acceleration voltage. The beam intensity measured on the phosphorous screen is ∼50 pA/cm². The *E. coli* cells used for fluorescence microscopy were suspended in 0.85% NaCl after growth medium was removed. The sample was then mixed with 2x Bacterial Viability Kit stock solution with an equal volume. LIVE/DEAD BacLight Bacterial Viability Kit was from life technologies (Eugene, Oregon, USA). Briefly, a drop of 0.6 μL suspension was used in liquid chamber. The liquid chamber was imaged directly by fluorescence microscopy (Olympus IX51 inverted microscope, Tokyo, Japan) with exposure time: 67.08 ms. Green fluorescence image and corresponding red fluorescence image were merged together into one picture.

### 3. Results and Discussion

To investigate the cellular fine structures of bacterial cells under aqueous conditions, *in situ* TEM was performed on an *E. coli* sample in the liquid chamber. Most of the cells appeared to be structurally robust against electron beam irradiation and preserved nanostructures during the TEM imaging. The *E. coli* cells in the *in situ* bright field TEM images (Figure I(a)) were oval-rod shaped.
Figure 2: In situ TEM images (a, c) and corresponding fluorescence microscopy images (b, d) of E. coli cells in the liquid chamber after high energy beaming. The fluorescence images were obtained after TEM exposing. (c) and (d) are magnifications from squares in (a) and (b). The red arrows in (c) pointed to two cells which emitted obvious green fluorescence in (d).

comparable to previous in situ liquid chamber TEM studies [7, 18]. Furthermore, as seen in Figures 1(b)–1(d), our E. coli images show more details than other reports in the literature, with several cellular structures detected outside of the oval-rod shaped bacterial cells and a few cells showing dim dark nucleoid centers, which are indicated by the two green arrows in Figure 1(b). The fine structures outside of the E. coli cells in the in situ TEM images may be attributable to pili, which are displayed as gray halos around some of the bacterial cells (Figure 1(c)). Almost half of the E. coli cells showed pili, but the other cells had no apparent pili structures (Figure 1(d)). Whether the pili were observed might be related to the biological activity and cell life cycle of the bacteria. This conforms with the property of pili. It is possible that the resolution of in situ TEM might not be high enough to detect additional intracellular details of the E. coli cells, such as a clear nucleoid. However, images have been reported with better than 1 nm and even atomic resolutions using in situ liquid chamber TEM, which indicates that getting enough resolution should not be an obstacle for the technology. In our in situ TEM images of E. coli cells, pili were observed clearly, which are known to have diameters less than 10 nm. Resolutions better than 10 nm have been obtained as measured from the TEM images, which is more than enough to image a clear nucleoid if it exists. Thus, the lack of details for E. coli is not due to the resolution limit of in situ TEM but may be because of the stage of the cells during the observation. The absence of low contrast in the dark centers may suggest that the nucleoids in the E. coli cells under the fully hydrated environment were not as condensed as those cells that were dried or processed to be used in other imaging techniques.

With nanometer resolutions and the capability of observing cells in their fully hydrated living environments, in situ TEM is a more powerful tool than conventional EMs. However, the length of time that a cell can withstand the electron beam irradiation without being subjected to structural damage or cell death is also of concern. Currently, studies have only successfully irradiated cells for several seconds using in situ TEM, which has largely limited its applications to in situ monitoring of dynamic changes in the cells. In this study, E. coli cells demonstrated good structural stability. The cells were observed under the electron beam for extended time, and the majority of the cells did not show beam-induced structural damage. The cell viability was examined by using LIVE/DEAD BacLight Bacterial Viability Kit, and the observations are shown in Figure 2. The cells exposed under TEM high energy electron beam (Figure 2(a)) still emitted green fluorescence (which indicated alive cells) (Figure 2(b)), instead of red fluorescence (which indicated dead cells). Figures 2(c) and 2(d) show magnified versions of the images. The red arrows in Figure 2(c) pointed to two cells which emitted obvious green fluorescence as shown in Figure 2(d). The other cells also emitted green fluorescence but could not be observed clearly, possibly due to a variation of imaging depth. These results further verified the survival of the E. coli cells through the analysis.

In addition to the overall stability and integrity under the in situ TEM condition, several E. coli cells ruptured. In Figure 3(a), a few cells in the right area of the picture...
that are circled by red dotted lines were found to be lighter, suggesting that they were broken cells. To the right of the figure, we detected several such broken cells forming a vertical line. In the magnified image in Figure 3(b), one cell was broken from the top end. Compared to the intact cells, the ruptured cell showed a size extension at the broken site, which might partially be because of the unfolding of the cell membrane and the extruding pressure from the cytoplasm. The nonruptured end appeared to be smaller, possibly related to the loss of cytoplasm and pressure inside the cell. In Figure 3(c), in addition to the partially broken cell in the upper part, a completely broken cell is displayed in the lower part of the image. This cell showed a dim cell membrane with a larger diameter than a complete cell, and the cellular region within the membrane boundary has a lighter shade than an unbroken cell. In Figure 3(d), in addition to the cell to the right that was broken at one end, one *E. coli* cell to the left was broken from the middle, forming a pore. Figure 3(e) shows a lytic broken cell forming membrane fragments, and as a result the outline of the oval shape of the cell was not clear. Several studies have shown that *E. coli* cells become lytic when the cell membrane is damaged [19], releasing intracellular components [20]. The phenomenon we observed could be cell lysis induced by damage to the membrane. In previous studies, pores have been suggested to form at the early stages. Additionally, the osmotic pressure across the membrane might also induce membrane rupture and cause more dramatic damage to a bacterial cell [21]. The initiation of this rupture was not clear and may not necessarily be caused by the imaging because these cells might have already broken before being examined.

Although rare, a dynamic process of cell damage has also been observed using *in situ* TEM. Within a minute, an oval-shaped *E. coli* was quickly damaged. The outer ring of the oval-shaped *E. coli* cell became deeper gray, getting darker and wider with time. The image of the *E. coli* cell became brighter with time (Figures 4(a)–4(d)). In the final image (Figure 4(f)), the contour of the *E. coli* was rod-like, smaller in size, and the brightness matched the background of the picture. In Figures 4(a)–4(e), approximately eight growing dark spots were also observed. The spots might originate from the dissolution of cellular debris. One reason for such coincidental damage might be the effect of radiation from beam-sample interactions. When high energy electrons pass
through an aqueous solution, the liquid system produces radicals and hydrated electrons, such as OH radicals [22, 23]. These radicals and hydrated electrons could damage the biomolecules of the cells. To analyze the effect of radicals on E. coli cells, 10 mg/mL glucose was added to the medium as radical scavengers. In the liquid chamber containing glucose, most of the E. coli cells kept their native structures without damage. This chamber showed lower red fluorescence than the liquid chamber without glucose (shown in Figure S1 in Supplementary Material available online at http://dx.doi.org/10.1155/2015/829302). This indicated less dead cells in liquid chamber with glucose.

In the higher resolution images, pili were distinguished in the halo around the cells. Figures 5(a) and 5(b) show video frames taken from the labeled region in Figure 1(a). Partially because of the higher magnification, we obtained better images and can clearly detect the pili surrounding the cells. These pili observed in the current work resemble those previously reported using conventional imaging techniques [24, 25]. Although the samples were in an isolated chamber, the liquid flow can still occasionally be observed in the sample. The liquid flow behavior was captured in the video. In Figure 5(c) (lower right part of the image), we detected a thicker liquid layer flowing into the cell, which got darkened and blurred. Figure 5(d) shows an image taken after the thick liquid covered the full imaging region, which produced an almost featureless picture because of the electron diffraction through the thick sample. Figures 5(e)–5(h) show magnified video frames taken from the labeled region in Figure 5(a); the arrow points to a pilus that changed shape with time. For the video of Figure 5 only, the imaging time was longer than 60 s, much longer than reported exposure time in the literature, and we did not see any structural damage to the cell. This makes a significant progress in dynamic observation of biosample using in situ liquid chamber TEM.

Generally, people believe there are big bubbles in the in situ liquid chamber [26, 27], filling the majority of the space, leaving possibly only an ultrathin layer of liquid on the two windows, which was indicated in Figure 6. Thus the cells are not necessarily surrounded by liquid only but are surrounded by liquid plus vapor, which explains the good image resolution. Even when surrounded by vapor only, the surrounding vapor pressure is much larger than in an environmental TEM; thus the cells are more hydrated. Thus, it is easily understood that the liquid layer thickness can change with time due to liquid flow, resulting in the image resolution change, supporting the observation in Figures 5(a)–5(d) [18]. In addition to the motion of the pili and the liquid-cell interactions observed, TEM allows the study of various events, such as flagellar motions, nanoparticle-cell interactions, and even cell divisions. Therefore, this tool could be beneficial in numerous important studies investigating cell structures and processes.

4. Conclusions

In conclusion, in situ liquid chamber TEM was used to image E. coli cells in liquid environments. The in situ TEM results clearly displayed contours and fine cellular structures (such as pili) of the E. coli cells. Notably, the nucleoids in the E. coli cells in the fully hydrated environment appeared to be in noncondensed forms. Additionally, during our in situ TEM experiments, most of the E. coli cells were robust under the electron beam, without showing structural damage over time. The liquid chamber TEM allowed several fine structure and motion observations that were impossible with traditional
Figure 5: Video frames taken from the labeled region in Figure 1(a): (a) 0 s, (b) 2 s, (c) 40 s, and (d) 60 s. (e)–(h) Magnified video frames taken from the labeled region in Figure 5(a): (e) 0 s, (f) 10 s, (g) 20 s, and (h) 30 s. The arrow in (e) points to a pilus that changed shape with time. The scale bar in (h) is 0.2 μm.

Figure 6: Schematic of the in situ liquid chamber. Liquid with *E. coli* cells was filled in chamber formed by two Si₃N₄ membranes. A bubble existed in the chamber, which was benefit to high-resolution images of biosamples.

TEM or SEM analyses. Liquid chamber TEM is capable of in situ monitoring of significant biological processes, including cell rupture and cellular structural movement. A variety of cell damage types were also observed. Notably, liquid flow behavior was observed. Additionally, the motion of the pili was recorded for the first time using the liquid chamber TEM technology. Overall, the in situ liquid chamber TEM is a facile and efficient tool for *E. coli* analysis. This method provides reliable studies of the cellular structures and behaviors in their native growing environments.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

Acknowledgments

This work was supported by the Major Programs of Ministry of Education of China and National Natural Science Foundation of China (21303050 and 31471659), the China Postdoctoral Science Foundation (2013M540334), the Science and Technology Commission of Shanghai Municipality Project (11nm0507000), the Shanghai Leading Academic Discipline Project (B502), and the SRF for ROCS, SEM.
References

[1] R. F. Egerton, “Electron energy-loss spectroscopy in the TEM,” Reports on Progress in Physics, vol. 72, no. 1, Article ID 016502, 2009.

[2] D. R. Keene, S. F. Tufa, G. P. Lunstrum, P. Holden, and W. A. Horton, “Confocal/TEM overlay microscopy: a simple method for correlating confocal and electron microscopy of cells expressing GFP/YFP fusion proteins,” Microscopy and Microanalysis, vol. 14, no. 4, pp. 342–348, 2008.

[3] O. Medalia, I. Weber, A. S. Frangakis, D. Nicastro, G. Gerisch, and W. Baumeister, “Macromolecular architecture in eukaryotic cells visualized by cryoelectron tomography,” Science, vol. 298, no. 5596, pp. 1209–1213, 2002.

[4] A. Rigort, F. J. B. Bäuerlein, A. Leis et al., “Micromachining tools and correlative approaches for cellular cryo-electron tomography,” Journal of Structural Biology, vol. 172, no. 2, pp. 169–179, 2010.

[5] A. Hoenger and J. R. McIntosh, “Probing the macromolecular organization of cells by electron tomography,” Current Opinion in Cell Biology, vol. 21, no. 1, pp. 89–96, 2009.

[6] A. Leis, B. Rockel, L. Andrees, and W. Baumeister, “Visualizing cells at the nanoscale,” Trends in Biochemical Sciences, vol. 34, no. 2, pp. 60–70, 2009.

[7] K.-L. Liu, C.-C. Wu, Y.-J. Huang et al., “Novel microchip for in situ TEM imaging of living organisms and bio-reactions in aqueous conditions,” Lab on a Chip, vol. 8, no. II, pp. 1915–1921, 2008.

[8] M. J. Williamson, R. M. Tromp, P. M. Vereecken, R. Hull, and F. M. Ross, “Dynamic microscopy of nanoscale cluster growth at the solid-liquid interface,” Nature Materials, vol. 2, no. 8, pp. 532–536, 2003.

[9] X. Chen, L. H. Zhou, P. Wang, C. J. Zhao, and X. L. Miao, “A study of nano materials and their reactions in liquid using in situ wet cell TEM technology,” Chinese Journal of Chemistry, vol. 30, no. 12, pp. 2839–2843, 2012.

[10] X. Chen and J. Wen, “In situ wet-cell TEM observation of gold nanoparticle motion in an aqueous solution,” Nanoscale Research Letters, vol. 7, Article ID 598, 2012.

[11] Y. Liu, X. Chen, K. W. Noh, and S. J. Dillon, “Electron beam induced deposition of silicon nanostructures from a liquid phase precursor,” Nanotechnology, vol. 23, no. 38, Article ID 385302, 2012.

[12] H. Zheng, S. A. Claridge, A. M. Minor, A. P. Alivisatos, and U. Dahmen, “Nanocrystal diffusion in a liquid thin film observed by in situ transmission electron microscopy,” Nano Letters, vol. 9, no. 6, pp. 2460–2465, 2009.

[13] J. E. Evans, K. L. Jungjohann, P. C. K. Wong et al., “Visualizing macromolecular complexes with in situ liquid scanning transmission electron microscopy,” Micron, vol. 43, no. 11, pp. 1085–1090, 2012.

[14] D. F. Parsons, “Structure of wet specimens in electron microscopy,” Science, vol. 186, no. 4162, pp. 407–414, 1974.

[15] N. de Jonge, D. B. Peckys, G. J. Kremers, and D. W. Piston, “Electron microscopy of whole cells in liquid with nanometer resolution,” Proceedings of the National Academy of Sciences of the United States of America, vol. 106, no. 7, pp. 2159–2164, 2009.

[16] D. B. Peckys and N. de Jonge, “Liquid scanning transmission electron microscopy applied to study live cells,” Microscopy and Microanalysis, vol. 17, supplement 2, pp. 204–205, 2011.

[17] T.-W. Huang, S.-Y. Liu, Y.-J. Chuang et al., “Self-aligned wet-cell for hydrated microbiology observation in TEM,” Lab on a Chip, vol. 12, no. 2, pp. 340–347, 2012.

[18] D. B. Peckys, G. M. Veith, D. C. Joy, and N. de Jonge, “Nanoscale imaging of whole cells using a liquid enclosure and a scanning transmission electron microscope,” PLoS ONE, vol. 4, no. 12, Article ID e8214, 2009.

[19] X.-F. Li, X.-Q. Feng, S. Yang, G.-Q. Fu, T.-P. Wang, and Z.-X. Su, “Chitosan kills Escherichia coli through damage to be of cell membrane mechanism,” Carbohydrate Polymers, vol. 79, no. 3, pp. 493–499, 2010.

[20] K. Xing, X. G. Chen, C. S. Liu, D. S. Cha, and H. J. Park, “Effect of oleoyl-chitosan nanoparticles as a novel antibacterial dispersion system on viability, membrane permeability and cell morphology of Escherichia coli and Staphylococcus aureus,” Carbohydrate Polymers, vol. 76, no. 1, pp. 17–22, 2009.

[21] M. Hartmann, M. Berditsch, J. Hawecker, M. F. Ardakani, D. GERTHSEN, and A. S. ULRICH, “Damage of the bacterial cell envelope by antimicrobial peptides gramicidin S and PGLa as revealed by transmission and scanning electron microscopy,” Antimicrobial Agents and Chemotherapy, vol. 54, no. 8, pp. 3132–3142, 2010.

[22] P. Fenter, S. S. Lee, Z. Zhang, and N. C. Sturchio, “In situ imaging of orthoclase–aqueous solution interfaces with X-ray reflection interface microscopy,” Journal of Applied Physics, vol. 110, no. 10, Article ID 102211, 2011.

[23] B. C. Garrett, D. A. Dixon, D. M. CAMAIONI et al., “Role of water in electron-initiated processes and radical chemistry: Issues and scientific advances,” Chemical Reviews, vol. 105, no. 1, pp. 355–390, 2005.

[24] E. Hahn, P. Wild, U. Hermanns et al., “Exploring the 3D molecular architecture of Escherichia coli type I pili,” Journal of Molecular Biology, vol. 323, no. 5, pp. 845–857, 2002.

[25] E. Miller, T. Garcia, S. Hultgren, and A. F. Oberhauser, “The mechanical properties of E. coli type I pili measured by atomic force microscopy techniques,” Biophysical Journal, vol. 91, no. 10, pp. 3848–3856, 2006.

[26] G. Zhu, Y. Jiang, W. Huang, H. Zhang, F. Lin, and C. Jin, “Atomic resolution liquid-cell transmission electron microscopy investigations of the dynamics of nanoparticles in ultrathin liquids,” Chemical Communications, vol. 49, no. 93, pp. 10944–10946, 2013.

[27] X. Chen, K. W. Noh, J. G. Wen, and S. J. Dillon, “In situ electrochemical wet cell transmission electron microscopy characterization of solid-liquid interactions between Ni and aqueous NiCl$_2$, Acta Materialia, vol. 60, no. 1, pp. 192–198, 2012.