Dear Editor,

Visualization of chromosome shapes and dynamics in a live cell is highly desirable and necessary in many areas of cell biology. For example, the copy number of a particular chromosome in cancer cells is often abnormal (e.g., more than two), and therefore probing chromosome copy numbers can aid cancer diagnosis. During interphase, each chromosome exists in its own territory in the nucleus, which can be imaged by fluorescence in situ hybridization (FISH) using sequence-specific probes of different colors [1, 2]. However, such chromosome painting has only been possible in fixed cells, and is not suitable for dynamic monitoring of live cells. Therefore, it would be valuable to visualize DNA replication of one chromosome during interphase, and follow chromosome dynamics in the M phase.

Recent development of clustered regularly interspaced short palindromic repeats/CRISPR-associated proteins (CRISPR/Cas) [3] has provided a powerful tool for live-cell imaging of genomic loci [4]. In particular, the nuclease defective Cas9 (dCas9) fused with enhanced green fluorescent protein (EGFP) is used to target a particular DNA sequence upstream of a protospacer adjacent motif (PAM) sequence. Such targeting is achieved through Watson-Crick base pairing of ~20-bp single-guide RNA (sgRNA) that is pre-complexed with the dCas9-EGFP protein. The targeted loci can thus be fluorescently labeled in live mammalian cells [5-8]. However, the labeling achieved by this method is usually restricted to the genomic loci that consist of repetitive sequences, and has not been attempted to track an entire chromosome in a live cell.

Here we report the specific labeling of a large number of loci in the genome, which makes it possible to paint an entire chromosome in a live cell. To do so, we designed a new strategy using a large number of sgRNAs targeting mainly the non-repetitive regions of the chromosome (Figure 1A and Supplementary information, Figure S1A). To design sgRNAs, we scanned the sequence of the entire chromosome 9 on human reference genome hg19. Each 19-23 bp genome sequence upstream of a PAM sequence NGG was taken as a candidate target region. Because the efficiency of sgRNA binding is dependent on its GC content [9, 10], sgRNAs with GC content of 45%-65% were selected (Supplementary information, Figure S1B). The sgRNAs that could also bind to other chromosomes were removed in order to assure labeling specificity and reduce fluorescent background. Additionally, when multiple targeting sequences overlapped in a chromosome region, only one of them was selected. Among all sgRNAs, only one protein-coding gene (DNAJB5) contains sgRNA-binding sites in the exon region (Supplementary information, Figure S1C), but its expression level was not affected by lentivirus infection or dCas9-EGFP/sgRNA targeting (Supplementary information, Figure S1D).

To obtain high densities of sgRNA that would give signals significantly above the intracellular fluorescent background, especially when the chromatin is in open state during G and S phases, we chose 15 clusters of target sites (c1-c15) on chromosome 9, each spanning 5 Kb and containing more than 30 targets. The clusters are placed at least 5 Mbp away from each other (Figure 1A). In total, we selected 1124 sgRNAs that are distributed on chromosome 9 (Supplementary information, Table S1). As shown in Supplementary information, Figure S1A, a cell line stably expressing dCas9-EGFP was constructed from one HeLa cell by lentiviral infection. In the meantime, the 1124 sgRNAs were packaged into lentiviruses and used to infect HeLa cells expressing dCas9-EGFP for three times. After labeling the chromosome with the 1124 sgRNAs, we imaged cells in the S phase using a Nikon structured illumination microscope (N-SIM) equipped with a 100× TIRF oil immersion objective (NA 1.49). Figure 1B shows that the projected z-stack EGFP images. We attribute the three bright EGFP signal regions in the nucleus to three copies of chromosome 9, which is consistent with the karyotype of the HeLa cells we used (Supplementary information, Figure S1E). We noted that the EGFP signal is stronger in the S phase, when DNA replication takes place, than in the G1 phase, indicating that dCas9-EGFP can bind rapidly to newly synthesized DNA. Taken together, the imaging results suggest the efficacy of our method in visualizing a desired chromo-
Painting a chromosome with CRISPR/Cas9

We found the imaging results for the chromosome territory in a live cell. Figure 1C shows a projected z-stack of N-SIM fluorescence images taken in the prophase during mitosis. The high resolution of structured illumination microscopy and the strong signal due to chromosome condensation, allowed for the visualization of the three pairs of sister chromatids (Figure 1C) after DNA replication.

We next aimed to verify the dCas9-EGFP signals with FISH using probes targeting repetitive sequences so that a single FISH probe could access many sites. We used another two sgRNA sequences, C9-1 and C9-2, which were previously described and named [7], to replace the three non-repetitive clusters c13-c15. C9-1 binds to a region with pericentromeric repeats on chromosome 9 and C9-2 loci labeled by Cy3-tagged FISH probes (red). Scale bar, 5 μm. (F) Dynamics of the three copies of chromosome 9 in S phase in a live HeLa cell. Z maximum intensity projection of 10.5 μm. Scale bar, 5 μm. See also Supplementary information, Movie S1. (G) Distribution of sgRNA sequences from a single clonal sample detected by DNA sequencing after PCR amplification using the common primers for the sgRNA sequences. Single clone 1 (S1) was selected from a cell pool that has been painted with 802 sgRNAs. Majority of the sgRNA sequences have been incorporated into a cell single clone. The horizontal dashed line indicates the level with 10 sgRNA sequences per cluster, which is necessary for effective chromosome painting. (H) Distribution of numbers of reads for different sgRNA sequences, showing efficient incorporation and even distribution of the pooled 802 sgRNA library in the cell population.

After labeling the chromosome with the 802 sgRNAs, we found the imaging results for the chromosome territory and three pairs of sister chromatids are very similar to those labeled with c1-c15 containing 1 124 sgRNAs (Supplementary information, Figure S2D and E). The labeled chromosome 9 in the M phase can be easily seen even with a wide-field fluorescence microscopy. Figure 1D shows the cells in the prophase, metaphase, anaphase and telophase during mitosis. The cells were also stained with a DNA-specific dye Hoechst 33342, which allows co-localization of chromosome 9 with the rest of chromosomes. The three pairs of sister chromatids in the prophase and metaphase, though not clearly resolved due to the reduced resolution, separate in the anaphase and telophase. As a final control experiment, Figure 1E shows co-localization of labeled chromosome 9 of the EGFP signal with the Cy3-tagged C9-1 and C9-2 FISH probes. The fact that the EGFP areas were larger than the diffraction-limited FISH areas proves again the successful labeling of chromosome 9.

Having demonstrated stable labeling of chromosome 9, we next applied our method to dynamic monitoring using a DeltaVision Imaging System (Applied Precision/GE) equipped with a 100×/1.4 NA oil immersion objective, which allows less photobleaching, longer time for data acquisition, and confocality based on deconvolution. Figure 1F and Supplementary information, Movie S1 show chromosome dynamics at a single chromosome level for a cell in the S phase. Supplementary information, Figure S2F and Movie S2 show the dynamics of chromosome 9 during a period of 2 h from the late S phase to M phase. In addition to the three chromosome 9 spots, the nucleoli exhibit unintended EGFP signal, likely due to nonspecific binding of dCas9-EGFP protein with small RNAs in the nucleoli [8]. During the data collection period, EGFP signal from the nucleoli faded away due to the disappearance of the nucleoli in the M phase, while chromosome 9 fluorescent signal became stronger due to chromosome condensation. Supplementary information, Movie S3 shows another cell in the M phase for a period of ~3 h. After cell division, the two daughter cells maintained strong fluorescence signals from the replicated chromosomes, indicating that chromosome labeling is kept in the daughter cells. In this movie, significant conformational fluctuation of the condensed chromosomes was observed.
Although hundreds of sgRNAs were delivered by lentivirus into the cells, no obvious effects on cell proliferation were observed (Supplementary information, Figure S2G).

To evaluate how many sgRNAs were actually introduced to each cell, we carried out DNA sequencing of clonally amplified cells after PCR amplification with the common primers for sgRNA sequences [11]. Figure 1G shows the distribution of the detected 510 non-repetitive sgRNA sequences in chromosome 9 in one clonally amplified population. Single clones with higher labeling efficiency (S1) had more sgRNAs sequences incorporated than single clones with weaker fluorescence (S34) (Supplementary information, Figure S2H). Figure 1H shows the distribution of read numbers for all sgRNA sequences in the population of cells, showing that majority of the sgRNAs has been sufficiently incorporated into the cell population.

To address the question of how many sgRNAs are sufficient for painting a chromosome, we reduced the number of non-repetitive sgRNAs to 485. We found no significant deterioration of the image quality when compared with the images produced with 802 sgRNAs (Supplementary information, Figure S2I), when we imaged chromosome 9 at different phases of the M phase. Based on this, we conclude that more than 20 sgRNAs in each cluster and at least 300 types of total sgRNAs are needed in one cell for effective chromosome 9 painting. As the required number of the sgRNAs should be dependent on the length of the chromosome, we anticipate that using our design strategy ~800 sgRNAs should be enough to label the longest human chromosome 1, which is about twice as long as chromosome 9, while ~100 sgRNAs could be enough for the shortest chromosome. Of course, for painting other chromosomes, additional optimization and validation may be required.

In summary, by introducing hundreds of specific and non-repetitive sgRNAs in a human cell, we are able to paint an entire chromosome in a live cell for fluorescent imaging. We have visualized the spatial arrangements of homologous chromosomes and sister chromatids and tracked the movement of a particular chromosome in dividing cells. Our method will facilitate studies of functional organization of chromosomes, interactions among different chromosome regions, and long-term chromosomal dynamics in live mammalian cells.

Acknowledgments

We thank Dr Yujie Sun for providing the dCas9-EGFP expressing plasmid and Dr Ping Wei from the Center of Quantitative Biology for the Tet-on 3G plasmids. We thank Shaoshuai Meng for his early involvement in this project. We appreciate the assistance of Bo He, Junya Zhao and Zujia Wang from Central South University in karyotype analysis. We are grateful to Xiaochen Li, Xuemei Hao and Chunyan Shan in Peking University imaging user facility for their help of acquiring cell division movies and N-SIM images. We thank Dr Patty Purcell for editing of the manuscript. Meanwhile we appreciate the assistant of Huan Hu for the guidance of FISH protocol. This work is supported by grants from the National Natural Science Foundation of China, NSFC 21390412 and NSFC21327808 for XSY, and NSFC31430025, NSFC31170126, and NSFC81471909 for WW, and in part by a grant from National Basic Research Program of China, 2015AA020407 for LH.

Yuexin Zhou1,2,3,*, Ping Wang1,2,*, Feng Tian3,4,5,*, Ge Gao3,5, Lei Huang1,2, Wensheng Wei1,2,3,4, X Sunney Xie1,2,6

1 Biodynamics Optical Imaging Center (BIOPIC), School of Life Sciences, Peking University, Beijing 100871, China; 2 Beijing Advanced Innovation Center for Genomics (ICG), Peking University, Beijing 100871, China; 3 State Key Laboratory of Protein and Plant Gene Research, School of Life Sciences, Peking University, Beijing 100871, China; 4 Peking-Tsinghua Center for Life Sciences (CLG), Academy for Advanced Interdisciplinary Studies, Peking University, Beijing 100871, China; 5 Center for Bioinformatics, School of Life Sciences, Peking University, Beijing 100871, China; 6 Department of Chemistry and Chemical Biology, Harvard University, Cambridge, MA 01238, USA

Correspondence: Lei Huang, Wensheng Wei, X Sunney Xie

E-mail: leihuangmed@gmail.com
E-mail: wswei@pku.edu.cn
E-mail: xie@chemistry.harvard.edu

*These three authors contributed equally to this work.

References

1. Cremer T, Cremer C. Nat Rev Genet 2001; 2:292-301.
2. Croft JA, Bridger JM, Boyle S, et al. J Cell Biol 1999; 145:1119-1131.
3. Jinek M, Chylinski K, Fonfara I, et al. Science 2012; 337:816-821.
4. Chen B, Gilbert LA, Cimini BA, et al. Cell 2013; 155:1479-1491.
5. Shao S, Zhang W, Hu H, et al. Nucleic Acids Res 2016; 44:e86.
6. Ma H, Tu LC, Naseri A, et al. Nat Biotechnol 2016; 34:528-530.
7. Ma H, Naseri A, Reyes-Gutierrez P, et al. Proc Natl Acad Sci USA 2015; 112:3002-3007.
8. Chen B, Hu J, Almeida R, et al. Nucleic Acids Res 2016; 44:e75.
9. Gilbert LA, Horlbeck MA, Adamson B, et al. Cell 2014; 159:647-661.
10. Wang T, Wei JJ, Sabatinii DM, et al. Science 2014; 343:80-84.
11. Zhou Y, Zhu S, Cai C, et al. Nature 2014; 509:487-491.

(Supplementary information is linked to the online version of the paper on the Cell Research website.)

This work is licensed under a Creative Commons Attribution 4.0 Unported License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/

© The Author(s) 2016