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Catalytically inactive Cas9 impairs DNA replication fork progression to induce focal genomic instability

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ABSTRACT

Catalytically inactive Cas9 (dCas9) has become an increasingly popular tool for targeted gene activation/inactivation, live-cell imaging, and base editing. While dCas9 was reported to induce base substitutions and indels, it has not been associated with structural variations. Here, we show that dCas9 impedes replication fork progression to destabilize tandem repeats in budding yeast. When targeted to the CUP1 array comprising ~16 repeat units, dCas9 induced its contraction in most cells, especially in the presence of nicotinamide. Replication intermediate analysis demonstrated replication fork stalling in the vicinity of dCas9-bound sites. Genetic analysis indicated that while destabilization is counteracted by the replisome progression complex components Ctf4 and Mrc1 and the accessory helicase Rrm3, it involves single-strand annealing by the recombination proteins Rad52 and Rad59. Although dCas9mediated replication fork stalling is a potential risk in conventional applications, it may serve as a novel tool for both mechanistic studies and manipulation of genomic instability.

INTRODUCTION

Cas9 is an RNA-guided endonuclease that cleaves double-stranded DNA at its target site, and the ease of designing single-guide RNA (sgRNA) has made it the most popular tool in genome editing. Catalytically inactive Cas9 (dCas9) bears mutations at two nuclease domains and has enabled a variety of applications. For instance, dCas9 can inhibit the progression of RNA polymerase to suppress transcription of the gene to which it binds (CRISPRi) (1). Moreover, dCas9 has been fused or complexed with fluorescent proteins, transcriptional activation/repression or

epigenetic modification domains, and adenosine/cytidine deaminases to enable live-cell imaging of genomic loci, targeted gene activation/inactivation, and base editing, respectively (2). These applications take advantage of the function of dCas9 as a programmable sequence-specific DNAbinding protein. Since dCas9 lacks nuclease activity, it was presumed to be non-mutagenic. However, it was also reported to promote mutagenesis at a frequency of $\sim 10^{-5}$ via R-loop formation (3). Most of these mutations were base substitutions attributable to spontaneous cytosine deamination of the non-target DNA strand of the dCas9-induced R-loop, whereas others included homopolymer instability and trans-lesion synthesis (TLS) (3). While dCas9 induces base substitutions and small indels, it has not been demonstrated, to our knowledge, to induce large structural variations (SVs). As recent studies reported Cas9 often induces unexpectedly large deletions around its target sites (4,5), the impact of dCas9-binding to genome DNA in vivo should be carefully examined in terms of SVs.

Eukaryotic genomes harbor many repetitive sequences in the form of tandem or interspersed repeats (6). They occasionally induce genomic instability leading to generation of SVs, including both pathogenic and adaptive copy number variations (CNVs). The most famous disease related to CNV of tandem repeats is triplet repeat disease, which is evoked by the expansion of arrays comprised of very short units, such as CAG, GAA, CGG, and CCG trinucleotides (7). In contrast, facioscapulohumeral muscular dystrophy is caused by the contraction of D4Z4 macrosatellite repeats comprised of a 3.3-kb unit harboring the DUX4 gene (8). CNV-mediated environmental adaptation has been well documented in budding yeast (9). When exposed to high concentrations of copper ions, yeast cells amplify the resistance gene CUP1 to rapidly generate adapted progenies. Most yeast strains contain tandemly iterated copies of CUP1 gene. Intriguingly, the presence of copper was shown to accelerate intra/sister chromatid recombination rate at CUP1 array (10). This enhanced recombination

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likely contributes to the generation of expanded CUP1 arrays and, hence, adapted progenies with enhanced copper resistance. Another example is rDNA comprising $\sim \! 150$ copies of tandemly iterated units, and its instability is involved in cellular senescence (11). Both examples notably involve replication fork stalling or collapse followed by its repair.

The replisome at the replication fork uses the Cdc45– Mcm2-7-GINS complex (CMG helicase) to unwind DNA for fork progression and DNA synthesis by replicative polymerases. The replicative CMG helicase associates with proteins in the replisome progression complex (RPC), which includes the checkpoint mediator Mrc1, the Tof1-Csm3 complex, the replisome adaptor protein Ctf4, the histone chaperone FACT and DNA topoisomerase I (12). Replication fork stalling occurs through the actions of these proteins upon encountering an obstacle. The accessory helicase Rrm3 removes DNA-bound proteins such as the origin recognition complex, the transcription regulator Rap1, and the replication fork-blocking protein Fob1, whereas the Tof1–Csm3 complex counteracts Rrm3 in the removal process (13,14). Depletion of these proteins induces the contraction or expansion of tandem repeats. For instance, $ctf4\Delta$ cells amplify the copy number of rDNA (15), and $mrc1\Delta$ or $rrm3\Delta$ cells destabilize both the CUP1 and rDNA arrays (14,16–18). Prolonged replication fork stalling results in its collapse, with or without DNA double strand breaks (DSBs). Cells have several pathways for coping with collapsed/broken forks, including homologous recombination (HR), non-homologous end joining (NHEJ), break-induced replication (BIR), TLS, template switching (TS) and single-strand annealing (SSA). During the repair process, repetitive sequences around the collapsed fork occasionally trigger the generation of SVs, including CNV of tandem repeat units. For the copper-accelerated CUP1 recombination described above, a model was proposed in which activated promoter activity-induced replication fork collapse is followed by BIR or fork restart using a homologous sequence on the sister chromatid (16).

Here, we report dCas9-induced CNV of tandem repeats. This finding led us to uncover that dCas9 impairs replication fork progression to induce focal genomic instability.

MATERIALS AND METHODS

Yeast strains

All yeast strains used in this study are derived from BY4741 ($MATa\,his3\,\Delta 1\,leu2\,\Delta 0\,met15\,\Delta 0\,ura3\,\Delta 0$) (19) (Supplementary Table S1). Standard culture media and genetic methods were used in this study (20). We deleted a gene of interest by transforming yeast cells with a DNA fragment composed of KanMX cassette sandwiched by the 5'- and 3'-flanking sequences of the open reading frame of the gene, which was amplified from the corresponding deletant strain in Yeast Deletion Clones MATa Complete Set (invitrogen) using PCR primers listed in Supplementary Table S2. To construct a strain bearing a URA3 insertion at the boundary of two neighboring CUPI repeat units, we transformed yeast cells with a DNA fragment composed of URA3 cassette sandwiched by the 3'- and 5'-end sequences of the repeat unit, which was obtained by PCR with

primers VIII214253::URA3-F and VIII214253::URA3-R listed in Supplementary Table S2. Transformants selected on agar plates of synthetic complete medium lacking uracil (SC-Ura) supplemented with 2% glucose were used for nanopore sequencing to determine the integration site of *URA3* in the *CUP1* array.

Plasmids

All plasmids used in this study are listed in Supplementary Table S3. All primers for plasmid construction were purchased from Sigma-Aldrich and Eurofins Genomics. Plasmids were constructed with the seamless cloning with HiFi DNA Assembly or Golden Gate Assembly obtained from New England Biolabs (NEB).

The integrative plasmid YIplac128-pCSE4-dCas9-tADH1 (*LEU2*) harbors a gene encoding *Streptococcus pyogenes* dCas9 fused with SV40 nuclear localization signal (NLS) as described previously (21) under the control of *CSE4* promoter. It was used for yeast transformation after NruI digestion to be integrated to the *CSE4* promoter on the genome.

The integrative plasmid YIplac128-pGAL1-dCas9-tADH1 (*LEU2*) harbors a gene encoding *S. pyogenes* dCas9 fused with SV40 NLS under the control of *GAL1* promoter. It was used for yeast transformation after AgeI digestion to be integrated to the *GAL1* promoter on the genome.

The integrative plasmid pFA6a-pACT1-yGEV-tADH1-HphMX (Hyg^R) harbors a gene encoding β -estradiol-responsive artificial transcription activator GEV (22) under the control of ACT1 promoter. It was used for yeast transformation after CpsCI digestion to be integrated to the ACT1 promoter on the genome. The GEV-coding sequence was codon-optimized for Saccharomyces cerevisiae.

The integrative plasmid pFA6a-pCUP2-yGEV-tADH1-HphMX (Hyg^R) harbors the codon-optimized GEV-coding gene under the control of CUP2 promoter. It was used for yeast transformation after MfeI digestion to be integrated to the CUP2 promoter on the genome.

Centromeric plasmids for sgRNA expression harbor sgRNA gene under the control of *SNR52* promoter or *GAL1* promoter. The sgRNA scaffold sequence contains a base-flip and an extension of a stem-loop for stable sgRNA expression (23). To cut off an unnecessary sequence from the 5′-terminal portion of the sgRNA-containing transcript, each sgRNA sequence is preceded by a hammer-head ribozyme (Supplementary Table S3). To define the 3′-terminus, each sgRNA sequence is followed by *SUP4* terminator on the *SNR52* promoter plasmid or by the HDV ribozyme on the *GAL1* promoter plasmid (Supplementary Table S3). For designing sgRNAs, CRISPRdirect (24) was used to select target sites in the yeast genome listed in Supplementary Table S4.

Gene editing

For constructing $rad52\Delta$ $rad59\Delta$ strains, we performed enAsCas12a-based gene editing. All gene-editing plasmids used in this study are listed in Supplementary Table S3. Each gene-editing centromeric plasmid (URA3, CEN) harbors a gene encoding enAsCas12a (25) fused with SV40

NLS and a gene coding CRISPR RNA (crRNA), both of which are under the control of GAL1 promoter. To improve the efficiency of gene editing, a 9-mer sequence (U₄AU₄) was attached to the 3'-end of crRNA (26). The crRNA is flanked by a hammerhead ribozyme and the HDV ribozyme at its 5'- and 3'-termini, respectively. For designing crRNAs, CRISPOR (27) was used to select target sites listed in Supplementary Table S4.

Yeast cells transformed with the gene-editing plasmid were spread on agar plates of SC-Ura supplemented with 2% galactose. After incubation at 30°C for 4–5 days, colonies were picked and streaked on a new plate. To examine successful gene-editing at the target site on the genome, we performed PCR to amplify a region spanning the target site. The PCR products were sequenced to reveal the size and position of indels around the target site (Supplementary Table S1).

To eliminate the gene-editing plasmid, the successfully gene-edited strains were grown in yeast extract/peptone/dextrose (YPD) liquid medium at 30°C overnight and streaked on a YPD agar plate to isolate single colonies. After incubation at 30°C for 2–3 days, each colony was streaked on agar plates of YPD medium and SC-Ura medium supplemented with 2% glucose to confirm the loss of the gene-editing plasmid.

Cell growth rate measurement

Cell growth rate was measured using the RTS-1 personal bioreactor (Biosan, Riga, Latvia). The properties were set as follow: the volume, 10 ml of SC medium supplemented with 2% glucose; the temperature, 25°C; the rotation speed, 1500 rpm; the measurement frequency, 10 times/min; and the reverse spin longitude, 1 s.

Cell culture for quantitative PCR (qPCR)

Yeast cells were grown at 25°C overnight in 5 ml of SC–Ura or SC-His medium supplemented with 2% glucose. On the following day, the OD_{600} of each sample was recorded and 10–50 μl of the culture was inoculated into 5 ml of the fresh medium containing 10 nM β-estradiol, supplemented with or without 5 mM nicotinamide (NAM). From the remaining overnight culture, genomic DNA was extracted with Gentra Puregene Yeast/Bact. Kit (QIAGEN) for qPCR. The same process was repeated every day. The division number per day was calculated from the change of OD_{600} .

qPCR

The concentration of genomic DNA was measured with Qubit dsDNA BR assay on Qubit 2.0 Fluorometer or Qubit Flex Fluorometer (Thermo Fisher Scientific). The DNA solution was diluted to a concentration of 0.5 ng/µl prior to qPCR. Each qPCR solution (20 μl) contained 2 μl of DNA (1 ng), 10 µl of TB Green Premix Ex Taq II (Tli RNaseH Plus) (Takara), 0.4 µl of ROX Reference Dye, 2 pmol each of the forward and reverse primers. The primers used for qPCR are listed in Supplementary Table S2. Each qPCR assay was performed in duplicate, using StepOnePlus or QuantStudio3 (Applied Biosystems) according to the manufacturer's instructions. Amplification condition was initial denaturation at 95°C for 30 s followed by 40 times iteration of a 3-step thermal cycle composed of 95°C for 10 s. 55°C for 30 s and 72°C for 5 s. All qPCR runs included 10fold serial dilutions to generate standard curves. The quantity of CUP1, ENA1 and URA3 was normalized to that of ACT1. The copy number of CUP1 and ENA1 in the standard curves was calibrated by the results of nanopore sequencing.

Genetic assay for the loss of *URA3* inserted into the *CUP1*

Yeast cells were grown at 25°C overnight in 5 ml of SC–His medium containing 2% glucose. On the following day, 15 μl of the culture was inoculated into 5 ml of fresh SC-His medium containing 2% glucose and 10 nM β-estradiol, supplemented with or without 5 mM NAM. The same process was repeated every day. After four days of cultivation, cells were appropriately diluted and spread onto SC glucose plates supplemented with 0.1% 5-FOA and YPAD plates to determine the frequency of 5-FOA-resistant clones.

Nanopore sequencing

Genomic DNA was extracted using Gentra Puregene Yeast/Bact. Kit (QIAGEN) and purified with 0.4× AM-Pure XP (Beckman Coulter) or Short Read Eliminator kit XL (Circulomics). To obtain high molecular weight DNA, we avoided vortexing and used mixing by gentle pipetting instead. DNA libraries were prepared using the ligation sequencing kit SQK-LSK109 (Oxford Nanopore Technologies) with or without barcoding. For barcoding, we used the native barcoding kit EXP-NBD104 or the rapid barcoding sequencing kit SQK-RBK004 (Oxford Nanopore Technologies) according to the manufacturer's instructions. We modified the protocol of the ligation sequencing kit as follows: DNA fragmentation, omitted; duration of the enzymatic repair steps at 20 and 65°C, both extended from 5 min to 30 min; and the duration of ligation step, extended from 10 to 30 min. The library was sequenced with the flowcell FLO-MIN106D R9.4.1 using the MinION sequencer (Oxford Nanopore Technologies). MinKNOW software was used to control the MinION device. The run time was set to 72 h. Base calling was performed using Albacore v2.3.1, Guppy v3.6.0, or Guppy v4.0.14. The assessment of sequencing data was performed using NanoPlot (28).

Dot plot analysis of nanopore sequencing reads

We used nanopore sequencing data in FASTA format to draw dot plots using YASS (29). We first selected reads spanning the entire array using 1-kb upstream and 1-kb downstream sequences of the CUP1 or ENA1 array as queries of minialign (https://github.com/ocxtal/minialign) and then used these reads as the first input sequence for YASS. As the second input, we used the reference sequence of interest (CUP1 repeat unit, ENA1 repeat unit, or URA3) or the selected read itself. By manually counting the number of diagonal lines appeared in each dot plot, we determined the CUP1 copy number, the ENA1 copy number and the location of *URA3* insertion in the *CUP1* array.

Computational counting of tandem repeat units in nanopore sequencing reads

To computationally count the copy number of tandem repeat units directly from each nanopore read, we developed a Fourier transform-based program termed DNA Sequence Detector, which works as follows.

Seq1 is a long DNA sequence sample to be examined (i.e. nanopore read), whereas Seq2 is a short and known DNA sequence (i.e. reference sequence of interest).

The nucleotide sequence of Seq1 is $s_0s_1...s_{n-1}$, and the nucleotide sequence of Seq2 is $r_0r_1...r_{m-1}$. Note that s_i and r_i are either A, T, G or C.

Then a matrix M is created as follows:

If s_i and $a_{i,j}$ are the same nucleotide, $a_{i,j}$ is replaced with '1'. If s_i and $a_{i,j}$ are not the same nucleotide, $a_{i,j}$ is replaced with '0'. Let the resulting matrix be M2.

$$M2 = \begin{bmatrix} & 0 & & & & & \\ & 1 & & & 0 & & \\ & 0 & & & 1 & & \\ & & 0 & & 1 & & \\ & & 1 & & \vdots & & \\ & & 1 & & 1 & & \\ & & 1 & & 0 & & \\ & & 0 & & & \end{bmatrix}$$

Next, each column in M2 is scanned. If '1' appears k times consecutively, these '1's are replaced with 'k'. Let the resulting matrix be M3.

$$M3 = \begin{bmatrix} 0 & & & & & \\ 1 & & & 0 & & \\ 0 & & & 12 & & \\ & & 0 & & 12 & & \\ & & 3 & & \vdots & & \\ & & 3 & & 12 & & \\ & & 3 & & 0 & & \\ & & 0 & & & 0 & & \end{bmatrix}$$

Next, all numbers below X (where X is a natural constant) are replaced with '0'. Let the resulting matrix be M4.

$$= [b_{i,j}], (0 \le i, j \le n-1)$$

 c_i is defined as follows:

$$c_i = \sum_{j=0}^{n-1} b_{i,j}, \ (0 \le i \le n-1)$$

Vector \vec{v} is defined as follows:

$$\vec{v} = \begin{bmatrix} c_0 & c_1 & \dots & c_{n-1} \end{bmatrix}$$

Next, the program searches a region in which values are dense and larger than a certain degree. Suppose that a region $R = [i \sim j]$ is found as such a region. This R is the region in which Seq2 is located.

Next, the program finds out the number of Seq2 present in R. The vector \vec{v} is smoothened to generate a vector $\vec{v}_{ave} = [d_0 \ d_1 \dots d_{n-1}]$. d_i is defined as follows:

$$d_{i} = \frac{\left(\sum_{j=i-L}^{i+L} c_{j}\right)}{2L+1} , (L \le i \le n-1-L)$$

 $d_i = c_i$, (i < L, i > n - 1 - L), L is a natural constant. By performing a discrete Fourier transform on a region R in \overline{v}_{ave} , the number of Seq2 in R is obtained.

Two-dimensional agarose gel electrophoresis (2D-AGE)

Yeast cells were grown at 25°C overnight in 5 ml of SC-Ura medium containing 2% glucose. Following the addition of 10 nM β-estradiol, the cells were cultivated for 2 h, diluted, and cultivated for 4 h. The genomic DNA was extracted with Gentra Puregene Yeast/Bact. Kit (QIAGEN) using a modified protocol, in which all vortexing steps were replaced by mixing with gentle pipetting to maintain the integrity of replication intermediates. The 2D-AGE followed by Southern blot hybridization was conducted as described previously (30,31) with some modifications. In brief, two micrograms of genomic DNA was fully digested with KpnI (Takara) or XcmI (NEB), precipitated with 1/10 volume of 3 M sodium acetate and one volume of isopropanol, washed with 70% ethanol, air dried, and finally dissolved in 30 µl of 10 mM HEPES-NaOH (pH 7.2). The first-dimension electrophoresis was performed on a 0.55% agarose gel (11 \times 14 cm) for 16 h at 22 V at room temperature. The second-dimension electrophoresis was performed on a 1.55% agarose gel (20 × 25 cm) containing ethidium bromide for 4 h at 260 mA at 4°C. The gel was sequentially soaked in depurination buffer, denaturing buffer, and neutralizing buffer, and the DNA was blotted to Hybond N+ membrane (Cytiva). Following UV-crosslinking, the blot was hybridized with a CUP1 probe at 55°C overnight. The probe was generated by PCR using the primers listed in Supplementary Table S4 followed by labeling with alkaline phosphatase using the labeling module of AlkPhos Direct Labelling and Detection System kit (Cytiva). Following appropriate washing of the blot at 60°C, chemiluminescent signals were generated using the CDP-Star Detection Reagent in the kit and detected with ImageQuant LAS4000 (Cytiva). Gel images were processed with ImageJ software (National Institutes of Health) for presentation images. The process involved rotating, cropping, and altering windowlevel settings. The spot of interest was selected as a circle for quantification, and the total internal intensity was divided by its area. The background was defined as the mean of area-normalized intensities of three randomly selected regions with no obvious signals.

Western blot

The expression of FLAG-tagged Rad52 was analyzed by western blotting. Proteins were extracted as described previously (32). Twenty micrograms of proteins (2 μ g/ μ l) were separated with sodium dodecyl sulfate-polyacrylamide gel electrophoresis using Any kD Mini-PROTEAN TGX Precast Gel (Bio-Rad). Transfer to membrane was performed with iBind Western System (Thermo Fisher Scientific) according to the manufacturer's protocol. Primary and secondary antibodies to detect Rad52-FLAG were FLAG M2 mouse monoclonal antibody (1:1000, Sigma-Aldrich) and goat anti-mouse IgG-HRP (1:2000, Santa Cruz Biotechnology), respectively. Primary and secondary antibodies to detect α-tubulin (loading control) were anti-alpha Tubulin antibody [YOL1/34] (1:2000, GeneTex) and goat Anti-Rat IgG H&L (HRP) (1:2000, Abcam), respectively. Following incubation with Clarity Western ECL Substrate (Bio-Rad), chemiluminescent signals were detected with ChemiDoc-Touch system (Bio-Rad). Gel images were processed with ImageJ software. The process involved cropping and altering window-level settings.

RESULTS

dCas9 induces copy number reduction of tandem repeat units

Cup1 is a metallothionein that buffers the concentration of intracellular copper in the budding yeast Saccharomyces cerevisiae (33,34). A \sim 2.0-kb unit including the CUP1 gene (CUP1 repeat unit) is tandemly iterated more than 10 times in the reference strain S288c to compose the CUP1 array on chromosome VIII (35). The CUP1 array in the parental strain used in this study was composed of ~16 repeat units (see below). The level of copper resistance linearly correlates with the CUP1 copy number (36–38). During experiments to target dCas9 to CUP1, we observed that the CUP1 copy number was decreased in a strain constitutively expressing CUP1-targeted dCas9 (Supplementary Figure S1A). The copy number was maintained in a control strain constitutively expressing dCas9 targeted to TEF1 on chromosome

XVI (39) (Supplementary Figure S1A). Moreover, the former strain, but not the latter, showed a sign of further decrease in the copy number during cultivation (Supplementary Figure S1B).

To further investigate this phenomenon, we constructed a strain in which dCas9 can be induced using β-estradiol without affecting cell growth (Figure 1A and Supplementary Figure S1C). This strain utilizes the artificial transcription factor GEV (Gal4 DNA-binding domain, estrogen receptor and VP16 transcription activation domain) (22), which translocates to the nuclei upon binding to β -estradiol and activates the GAL1 promoter to induce dCas9 expression. We grew the strain with daily dilution of the culture with fresh medium, extracted genomic DNA at various time points after B-estradiol addition, and measured the CUP1 copy number by qPCR. When TEF1-targeted dCas9 was induced, the copy number (~16 copies) did not show any significant change throughout the experiment (Figure 1B). In contrast, induction of CUP1-targeted dCas9 rapidly decreased the copy number in a time-dependent manner (Figure 1B). The extent of decrease varied from one sgRNA to another and was enhanced by the concurrent expression of three sgRNAs (CUP1a+b+c) (Figure 1B). The rate of decrease gradually slowed down, and the copy number appeared to reach a plateau in an extended culture (Supplementary Figure S1D).

As all three abovementioned CUP1 sgRNAs (CUP1a, b and c) bind to the same DNA strand, we designed an sgRNA that binds to the opposite strand (CUP1d) to test whether dCas9 reduces the CUP1 copy number in a strandspecific manner. CUP1d reduced the copy number with an efficiency largely comparable to that of CUP1a (Figure 1C). When combined, the two sgRNAs accelerated the copy number reduction (Figure 1C). These results suggested that dCas9 targeted to either DNA strand likely reduces the CUP1 copy number.

We next sought to determine whether the effect described above was specific to the CUP1 array. The yeast genome has several tandem repeats other than the CUP1 array, including the ENA1 array encoding P-type ATPase sodium pumps (40). The ENA1 array comprises a tandem array of three paralogous genes in the S288c reference genome sequence, namely ENA1, ENA2 and ENA5, but some strains harbor four or more paralogs (34,41). It was determined by nanopore sequencing that the strain used in this study had five paralogs (Figure 1D and Supplementary Figure S1E). We designed three sgRNAs for ENA1 to examine whether dCas9 targeting affects the copy number of ENA1 paralogs (Figure 1D). When dCas9 was targeted to ENA1 (ENA1a+b+c), the copy number decreased slowly (Figure 1E). This decrease was apparent in the presence of nicotinamide (NAM) (Figure 1E, see below).

Taken together, these results demonstrated that when targeted to tandem repeats, dCas9 reduces the copy number of repeat units in a sequence-specific manner.

NAM accelerates dCas9-induced copy number reduction of tandem repeat units

A previous study reported that NAM induces CUP1 CNV (16). NAM is an inhibitor of the NAD⁺-dependent his-

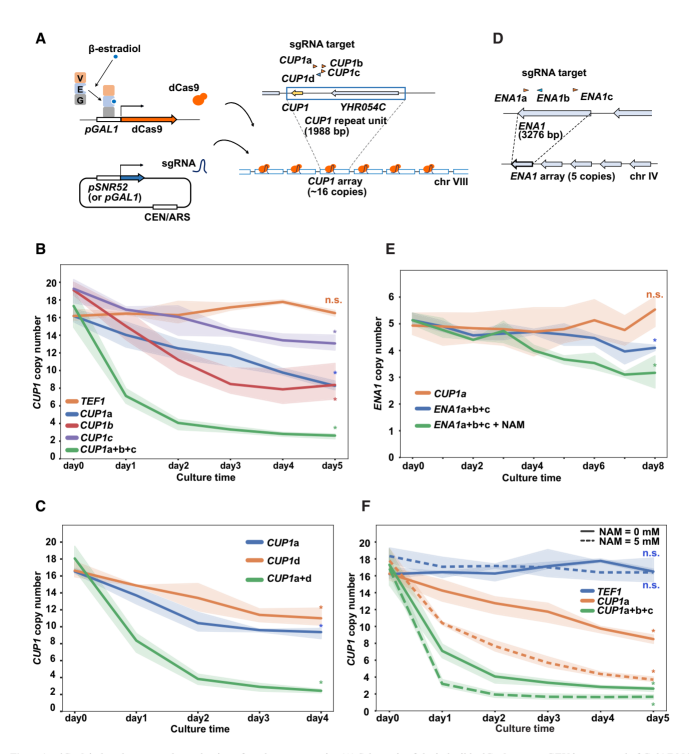


Figure 1. dCas9-induced copy number reduction of tandem repeat units. (A) Schematic of the inducible dCas9 system. GEV is composed of Gal4 DNA-binding domain (G), estrogen receptor (E) and VP16 transcription activation domain (V). GEV complexed with β-estradiol activates GAL1 promoter (pGAL1) to induce dCas9. sgRNAs are expressed from a centromeric plasmid under the control of SNR52 promoter (pSNR52) or pGAL1. Arrowheads above CUP1 repeat unit indicate four sgRNAs (orange, Watson strand; blue, Crick strand). (B) Time course of CUP1 copy number. The population-averaged copy number was determined with qPCR. Indicated sgRNAs were expressed using pSNR52. Expression of dCas9 was induced by the addition of 10 nM β-estradiol at day 0. Data are represented as mean ± standard deviation (i.e. line plots and shadows) (n = 3 or more biological replicates). Statistical significance of copy number alteration was examined between day 0 and day 5 using t-test (*t < 0.05). (C) Time course of t practices of t practices are included by t and sgRNAs. (E) Time course of t practices (*t < 0.05). (F) Effect of NAM on t on t day 4 using t-test (*t < 0.05). (F) Effect of NAM on t copy number. Similar to Figure 1B (t = 3 biological replicates). Statistical significance of copy number alteration was examined between day 0 and day 5 using t-test (*t < 0.05). (F) Effect of NAM on t on t copy number. Similar to Figure 1B (t = 3 biological replicates). Statistical significance of copy number alteration was examined between day 0 and day 5 using t-test (*t < 0.05).

tone deacetylase family, which includes Sir2, Hst1, Hst2, Hst3 and Hst4. Accordingly, concurrent deletion of SIR2. HST3 and HST4 destabilized the CUP1 array (16). Conversely, deletion of RTT109, encoding the sole histone acetyltransferase responsible for histone H3 acetylated at Lys-56 (H3K56ac), suppressed the NAM-induced CNV (16). In our study, the effect of NAM on the *CUP1* copy number was not significant in the control strain with TEF1targeted dCas9 (Figure 1F). In contrast, NAM substantially accelerated copy number reduction in the presence of CUP1-targeted dCas9 (Figure 1F). NAM also accelerated the copy number reduction of ENA1 paralogs induced by ENA1-targeted dCas9 (Figure 1E). Consistent with the previous study (16), NAM failed to exert its effect on dCas9induced CUP1 CNV in the absence of Rtt109 (Supplementary Figure S1F). These results suggest that NAM enhances dCas9-induced destabilization of tandem repeats through the elevation of H3K56ac.

Binding of a single dCas9 molecule can destabilize the CUP1 arrav

We wondered whether binding of a single dCas9 molecule can affect the copy number of tandem repeat units. To address this issue, we deployed a classical genetic assay based on the loss of URA3 integrated into the CUP1 array. For this assay, we generated a strain carrying a URA3 cassette in the center of the CUP1 array comprising 16 repeat units (Figure 2A and Supplementary Figure S2A). Upon destabilization of the CUP1 array, a fraction of recombination events between the repeat units led to the loss of the URA3 cassette, conferring on cells resistance to 5fluoro-orotic acid (5-FOA). A four-day induction of CUP1targeted dCas9 reduced the average CUP1 copy number (Figure 2B). Following this, we spread the cells onto agar plates supplemented with or without 5-FOA. As expected, the CUP1-targeted strain contained more 5-FOA-resistant cells than the control TEF1-targeted strain (51.7% versus 0.1%, 374.9-fold) (Figure 2C).

Confirming the performance of the genetic assay with CUP1-targeted dCas9, we next tested whether URA3targeted dCas9 destabilizes the CUP1 array. We used four sgRNAs (URA3a, b, c and d) to generate four strains with URA3-targeted dCas9 and subjected them to both the qPCR and genetic assays (Figure 2A). The qPCR assay failed to detect any significant decrease in the average CUP1 copy number in the four strains, presumably because copy number reduction occurred only in a limited fraction of the cell population (Figure 2B). However, in the genetic assay, the URA3-targeted strains generated 5-FOA-resistant clones much more frequently than the control TEF1-targeted strain (URA3a, 0.6%; URA3b, 4.3%; URA3c, 2.6%; URA3d, 1.4%; TEF1, 0.1%) (Figure 2C). Point mutations of URA3 could also confer 5-FOA resistance, and dCas9 was reported to induce base substitutions and indels (3). However, qPCR using DNA isolated en masse from 5-FOA resistant colonies confirmed deletion of *URA3* cassette, indicating that the contribution of point mutations was negligible (Supplementary Figure S2B). We thus concluded that even a single molecule of dCas9 can destabilize the CUP1 array, albeit much less efficiently than multiple dCas9 molecules targeted to individual repeat units.

dCas9 both contracts and expands the CUP1 array

The qPCR assay using an aliquot of liquid culture revealed the population average copy number but did not demonstrate the cell-to-cell variation. To determine this variation. we isolated single colonies from the cells cultivated in a liquid medium supplemented with β-estradiol and determined the *CUP1* copy number of each clone by qPCR (Figure 3A). As expected from the decreased population average, most of the 38 clones examined had reduced copy numbers. However, three clones appeared to have higher copy numbers than the original strain (#36–#38, Figure 3A).

The observed increase in the CUP1 copy number does not necessarily indicate the expansion of CUP1 array, as the CUP1 repeat unit has been shown to exist as an extrachromosomal circular DNA (42). Furthermore, aneuploidy may include chromosome VIII bearing the CUP1 array. We thus performed long-read sequencing using the Min-ION nanopore sequencer to reveal the CUP1 array structure in the three clones (Figure 3A). We selected reads containing both 5'- and 3'-flanking regions of the CUP1 array (i.e. reads spanning the entire array), generated a dot plot between each read and the reference sequence of CUP1 repeat unit, and manually counted the number of units from each plot (Figure 3B). We also developed a Fourier transform-based algorithm to calculate the copy number directly from nanopore reads to validate the results of manual counting (Supplementary Figure S3A). Consequently, the clones #36, #37 and #38 were demonstrated to harbor CUP1 arrays composed of 22, 19 and 21 repeat units, respectively (Figure 3B).

We next examined the population structure of the CUP1 array by sequencing genomic DNA prepared from CUP1and TEF1-targeted strains at days 0 and 5 of induction. In the TEF1-targeted strain, the CUP1 copy number in the array did not change during the culture (16.0 and 15.7 copies on average at days 0 and 5, respectively, by both manual and computational counting) (Figure 3C and Supplementary Figure S3B). In contrast, in the CUP1-targeted strain, the copy number distribution was obviously different between days 0 and 5. The copy number at day 0 showed a relatively homogenous distribution within a range of 14– 16 copies (15.4 and 14.9 copies on average by manual and computational counting, respectively). This low level of heterogeneity was presumably attributable to leaky expression of dCas9, as it was observed only in the presence of CUP1 sgRNA. The copy number at day 5 exhibited a significantly heterogenous distribution (5.0 and 4.7 copies on average by manual and computational counting, respectively) (Figure 3C and Supplementary Figure S3B). While 16 reads contained only a single copy of the CUP1 repeat unit, two reads spanned arrays comprising 19 and 28 units (Supplementary Figure S3C). Note that longer arrays are underrepresented in population analysis by nanopore sequencing compared to that by qPCR, as a longer array will have fewer reads spanning the entire array (Supplementary Figure S3D). Nevertheless, nanopore sequencing unequivocally demonstrated the expansion of the CUP1 array. Interestingly, it

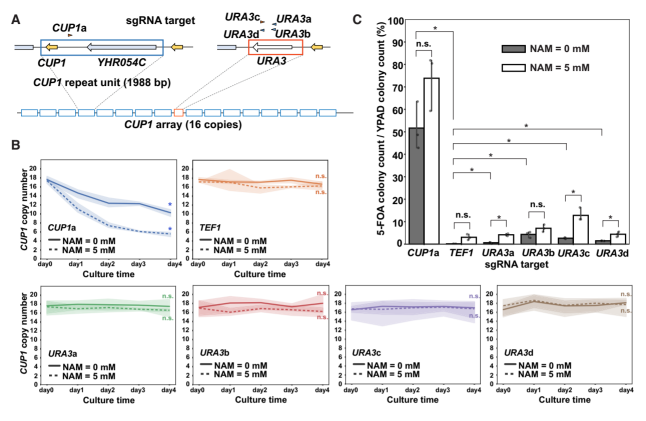


Figure 2. CUP1 array destabilization by single molecules of dCas9. (A) Schematic of URA3-bearing CUP1 array. Arrowheads indicate sgRNAs for CUP1 and URA3 (orange, Watson strand; blue, Crick strand). (B) Time course of CUP1 copy number in the strain with the URA3-bearing CUP1 array in the presence or absence of NAM. Similar to Figure 1B (n = 3 biological replicates). Statistical significance of copy number alteration was examined between day 0 and day 4 using t-test (*t > 0.05). (C) Frequency of t >

also revealed *CUP1* arrays with interstitial deletions (Supplementary Figure S3C), which led to the slight difference between the copy numbers estimated by manual and computational counting (Figure 3C and Supplementary Figure S3B).

Taken together, dCas9 contracts and expands the *CUP1* array in the majority and minority of cells, respectively, thereby inducing heterogeneity in the array structure.

dCas9 blocks replication fork progression in vivo

We hypothesized that dCas9-induced destabilization of the *CUP1* array stems from dCas9-mediated impairment of replication fork progression. Indeed, *CUP1*-targeted dCas9 failed to alter the copy number when the culture was saturated to terminate DNA replication (Figure 4A). To test this hypothesis directly, we conducted neutral-neutral 2D-AGE (43) and analyzed the status of DNA replication intermediates including the *CUP1* repeat unit by Southern blot hybridization (Figure 4B).

Prominent spots appeared on Y-arcs upon the induction of *CUP1*-targeted dCas9 (Figure 4C and D, blue arrows). These results indicated that dCas9 induced replication fork stalling in the *CUP1* unit. When KpnI-digested fragments were analyzed, the spot was observed approximately at the apex of Y-arc, indicating replication fork stalling around

the midpoint of the restriction fragment. Moreover, another spot appeared at the tip of X-spike, suggesting an accumulation of highly branched X-shaped molecules (Figure 4C, red arrow). It could be interpreted as two replisomes colliding around the midpoint of the fragment. When XcmI-digested fragments were analyzed, a spot was detected in the descending part of Y-arc upon induction of CUP1-targeted dCas9 (Figure 4D), suggesting replication fork stalling near an end of the fragment. Considering the dCas9-bound sites in the XcmI fragment, we speculate that the stalled replication fork likely originated from ARS813, located \sim 33-kb upstream of the *CUP1* array (Figure 4B). Although each CUP1 repeat unit has a weak replication origin (ARS810/811) (44,45), we failed to observe the bubblearc corresponding to the DNA replication bubble. This was presumably because ARS810/811 fires much less frequently than the closest neighboring replication origin for the CUP1 array. In summary, dCas9 impairs replication fork progression in the vicinity of its binding sites.

The RPC, accessory helicase and recombination proteins modulate dCas9-induced $\it CUP1$ CNV

A stalled replication fork may either resume progression or collapse. In the latter case, the cell exploits recombinational repair pathways to rescue the collapsed fork. We hypothe-

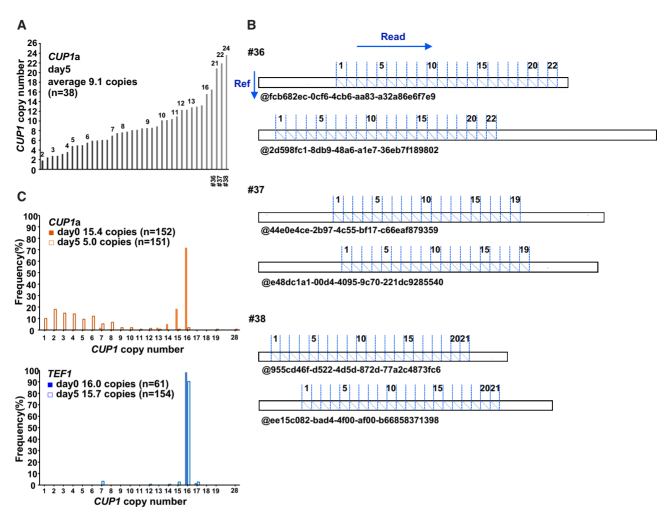


Figure 3. dCas9-induced expansion of CUP1 array. (A) CUP1 copy numbers in individual clones. Single colonies isolated at day 5 of dCas9 induction (Figure 1B) were used for qPCR. (B) CUP1 array structure revealed by nanopore sequencing. DNAs prepared from clones #36, #37 and #38 in Figure 3A were sequenced with MinION. Dot plots were generated between the reference sequence of CUP1 repeat unit (vertical axis) and nanopore sequencing reads (horizontal axis). Two representative reads are shown for each clone. (C) Population structure of CUP1 array. DNAs prepared from the cells with the indicated sgRNAs at days 0 and 5 of dCas9 induction (Figure 1B) were sequenced with MinION. CUP1 copy number determined from dot plots are

sized that dCas9-induced destabilization of tandem repeats is attributable to a repair process of replication forks stalled by dCas9. To obtain genetic evidence for and mechanistic insights into this process, we evaluated dCas9-induced reduction of the CUP1 copy number in a series of strains in which genes involved in replication fork stability and major DNA damage repair pathways were deleted (Figure 5A and Supplementary Figure S4A-C). We used qPCR to estimate the CUP1 copy number at days 0 and 2 of CUP1targeted dCas9 induction. To normalize the effects of differential growth rates among the strains, we evaluated the effects of gene deletions using a variation index (VI) defined as copy number change (%) per cell division (Supplementary Figure S4A, B). None of the 29 deletants examined abolished the dCas9-induced copy number alteration. The response to dCas9-mediated replication fork stalling may thus be redundant, with one pathway likely serving as a back-up for another. Nevertheless, certain strains showed significant change in VI (Figure 5A).

A significantly increased VI was observed in strains with CTF4, RRM3 and MRC1 deletions (ctf4 Δ , 5.57, P = 1.3E-06; $rrm3\Delta$, 5.35, P = 1.0E-05; $mrc1\Delta$, 5.08, P = 0.001; WT, 4.48) (Figure 5A). Although these genes have been implicated in tandem repeat stabilization (14–18), the CUP1 copy number was stably maintained in the $ctf4\Delta$, $rrm3\Delta$ and $mrc1\Delta$ strains with TEF1-targeted dCas9 (Supplementary Figure S4D). Note that Ctf4 and Mrc1 are components of the RPC and are known to affect fork stability (46–48). The accessory helicase Rrm3 removes obstacles in front of the replisome (17,49). Consistently, it appeared that dCas9induced replication fork stalling was enhanced in the $rrm3\Delta$ strain compared to the wild-type strain (Figure 5B and Supplementary Figure S4E-G). Compared to these three strains, the increment of VI in the $rad6\Delta$ strain was less significant (4.76, P = 0.014). Although the pol32 Δ strain was reported to accelerate CUP1 array contraction (16), it failed to exert a significant effect on the dCas9-induced CNV (4.69, P = 0.173).

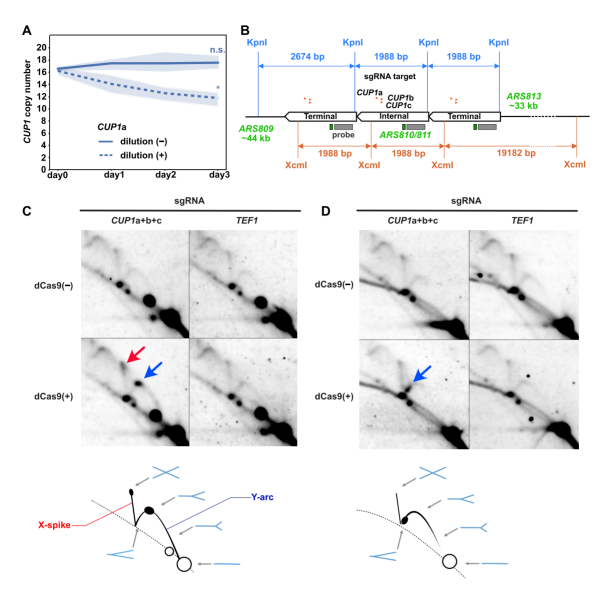


Figure 4. dCas9-mediated impairment of replication fork progression. (A) Effect of cell proliferation on dCas9-induced reduction of CUP1 copy number. Similar to Figure 1B, except that saturated cultures without daily dilution are included (n = 3 biological replicates). Statistical significance of copy number alteration was examined between day 0 and day 3 using t-test (*P < 0.05). (B) Design of 2D-AGE experiments. The CUPI array map indicates the positions of KpnI and XcmI restriction sites, sgRNAs, hybridization probe, ARSs and the lengths of bands detectable with the probe. Only a single copy of internal repeat units is depicted for clarity. (C) Representative 2D-AGE images of KpnI-digested DNA fragments. Blue and red arrows indicate the spots on the Y-arc (stalled replication fork) and the X-spike (highly branched X-shaped molecule), respectively. Schematics at the bottom indicate the interpretation of 2D-AGE patterns. Exposure time for signal detection was 4 h. (D) Representative 2D-AGE images of XcmI-digested DNA fragments. Blue arrow indicates the spot on the Y-arc. Schematics at the bottom indicate the interpretation of 2D-AGE patterns. Exposure time for signal detection was 4 h.

A significantly decreased VI was observed in strains with RAD52 and RAD59 deletions (rad 52Δ , 2.19, P = 6.62E– 09; $rad59\Delta$, 3.07, P = 7.31E-05). Notably, Rad52 plays critical roles in various recombinational repair pathways, and Rad59 is paralogous to and cooperates with Rad52. We thus used Cas12a-mediated gene editing (25) to disrupt RAD52 and RAD59 in the $rad59\Delta$ and $rad52\Delta$ strains, respectively. These double mutants failed to show a significantly decreased VI compared to the $rad52\Delta$ strain, suggesting that RAD52 is epistatic to RAD59 (Supplementary Figure S5A–D). Compared to the $rad52\Delta$ and $rad59\Delta$ strains, the decrease of VI was less significant in strains with EXO1, RTT109, MMS2 and TOF1 deletions (exo1 Δ , 3.39,

P = 0.010; $rtt109\Delta$, 3.67, P = 0.011; $mms2\Delta$, 3.72, P =0.002; $tof1\Delta$, 3.98, P = 0.030).

These results collectively indicated critical roles for the RPC components, accessory helicase, and recombination proteins in dCas9-induced CUP1 array contraction.

dCas9-induced CUP1 CNV involves SSA by Rad52

As RAD52 deletion had the largest impact on the VI (Figure 5A and Supplementary Figure S5A), we sought to determine how Rad52 contributes to dCas9-induced CUP1 array contraction. To this end, we took advantage of reported separation-of-function rad52 alleles. Rad52 is composed of

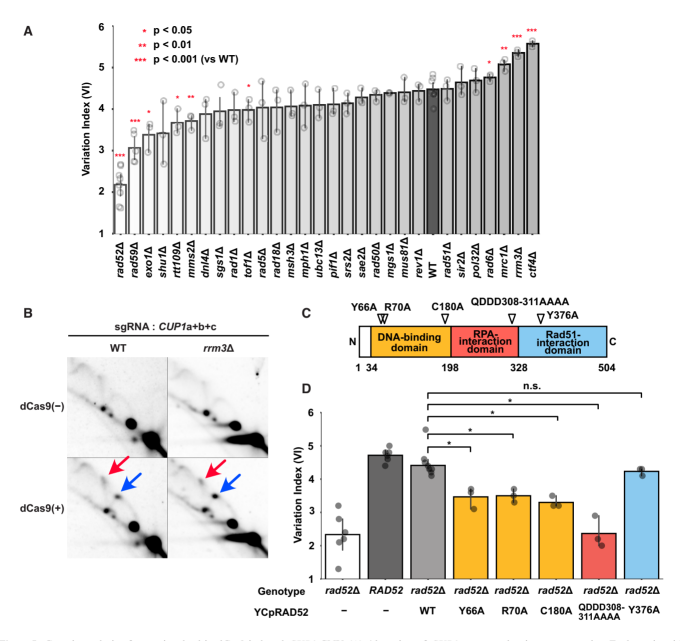


Figure 5. Genetic analysis of genes involved in dCas9-induced CUP1 CNV. (A) Alteration of CUP1 copy number in mutant strains. Each strain with three sgRNAs (CUP1a+b+c) was cultivated for two days in the presence of β-estradiol. Percentile change of CUP1 copy number estimated by qPCR was divided by the number of cell division estimated from absorbance to calculate VI. Data are represented as mean \pm standard deviation (n = 3 or more biological replicates). Statistical significance was calculated using t-test compared to the wild-type (WT) strain (*P < 0.05; **P < 0.01; ***P < 0.01). (B) Replication fork stalling in the wild-type and $rrm3\Delta$ strains. 2D-AGE patterns of KpnI-digested DNA are shown for the WT and $rrm3\Delta$ strains. Blue and red arrows indicate the spots on the Y-arc (stalled replication fork) and the X-spike (highly branched X-shaped molecule), respectively. The samples of the wild-type strain are identical to those in Figure 4C but exposure time for signal detection was 1 h. (C) Schematic of Rad52 domain structure with separation-of-function mutations. (D) Suppression of defective dCas9-induced CUP1 CNV in the rad52 Δ strain by wild-type and separation-of-function alleles. Data are represented as mean \pm standard deviation (n = 3 or more biological replicates). Statistical significance was examined between the strains harboring the WT and the separation-of-function alleles on YCpRAD52 using t-test (*P < 0.05).

three domains (Figure 5C). The N-terminal domain is evolutionarily conserved and mediates interactions with DNA, Rad52 (self-oligomerization), and Rad59. Class C mutants bearing mutations in this domain (rad52-Y66A, -R70A, -W84A, -R85A, -Y96A, -R156A, -T163A, -C180A and -F186A) are defective in SSA but proficient in HR (50–52). The central domain is required for binding to the singlestranded DNA (ssDNA)-binding protein RPA and formation of the Rad52 repair center. A mutant allele of this domain, rad52-QDDD308-311AAAA, encodes a protein that is defective in RPA-binding and mediator activity to exchange RPA for Rad51 on ssDNA, but is proficient in DNA binding, Rad51 binding, and SSA in vitro (53). The Rad52 C-terminal domain is also required for mediator activity. A mutant allele of this domain, rad52-Y376A, encodes a protein incapable of binding to Rad51 (54–56).

The wild-type and separation-of-function alleles were expressed in the $rad52\Delta$ strain under the control of the RAD52promoter on a centromeric plasmid vector. Immunoblot analysis confirmed comparable expression levels among the wild-type and mutant proteins except for Rad52-C180A (Supplementary Figure S5E). The *RAD52* allele suppressed defects in the $rad52\Delta$ strain, elevating the VI to a level comparable to that of the wild-type strain (Figure 5D and Supplementary Figure S5F-H). However, class C mutant alleles (rad52-Y66A, -R70A and -C180A) only partially suppressed the defect, indicating the involvement of the N-terminal domain and hence SSA (Figure 5D and Supplementary Figure S5F-H). Intriguingly, the rad52-QDDD308-311AAAA allele barely suppressed the defect (Figure 5D and Supplementary Figure S5F–H). In contrast, the rad52-Y376A suppressed the defect as efficiently as RAD52, demonstrating the dispensability of Rad51 binding and hence mediator activity (Figure 5D and Supplementary Figure S5F-H). Interestingly, depletion of Rad59, a C-terminally-truncated Rad52 paralog with SSA but not mediator activity, exerted a decelerating effect second only to that of Rad52 (Figure 5A). Furthermore, Rad51 depletion did not affect destabilization (Figure 5A). Thus, the data on the separation-offunction alleles were consistent with the deletant data.

Taken together, Rad52 likely contributes to dCas9-induced destabilization of tandem repeats through its involvement in the annealing of RPA-coated ssDNA, but not via its mediator activity.

DISCUSSION

dCas9 as a replication fork barrier

The highly plastic nature of the CUP1 array structure has been attracting attention for its role in environmental adaptation. A previous study reported critical roles for promoter activity and H3K56ac in CUP1 CNV (16). Here we showed that dCas9 induces CNV in the CUP1 and ENA1 arrays (i.e., array contraction and expansion), especially in the presence of NAM (Figures 1 and 3). Notably, dCas9 rapidly decreased the copy number even in the absence of transcriptional CUP1 induction (Figure 1). Although RTT109 deletion stabilizes the CUP1 array even in the presence of active transcription (16), CUP1-targeted dCas9 still destabilized the array in the $rtt109\Delta$ strain (Figure 5A). These results imply a high efficiency of dCas9 in inducing focal genomic instability. To our knowledge, this study is the first demonstration of dCas9-induced SVs. Intriguingly, NAM enhanced dCas9-induced CUP1 CNV in a H3K56ac-dependent manner (Supplementary Figure S1F). Since H3K56ac regulates replication-coupled nucleosome assembly (57), its hyperelevation likely alters chromatin status, thus affecting the stability of stalled replication fork and the accessibility of recombinational repair proteins and/or dCas9. These possibilities remain to be examined in future studies.

As both transcription and dCas9 induce *CUP1* CNV, dCas9 may serve as a mimic of RNA polymerase through its R-loop formation, with dCas9 generating an even longer R-loop than RNA polymerase. Notably, R-loop formation is a major threat to genome stability (58). We thus hypothesized that dCas9 and replisomes induce a conflict similar to that observed between transcription and replication. Consistent

with this scenario, we found that dCas9 impedes replication fork progression in vivo (Figure 4). Interestingly, a recently published report demonstrated the ability of dCas9 to block replisome progression in vitro (59).

Proteins tightly bound to DNA can impair replication fork progression, and some of them serve as physiological blocks (60). Examples of fork-blocking proteins include Tus binding to the replication terminator *Ter* of *Escherichia* coli, Fob1 binding to the replication fork barrier in rDNA of budding yeast, and Rtf1 binding to RTS1 in the mating locus of the fission yeast Schizosaccharomyces pombe (60). While these proteins function in an orientation-dependent manner, dCas9 appears to block the replication fork approaching from either side in vivo (Figure 1), consistent with the previous finding in vitro (59). The accumulation of highly branched X-shaped molecules at the tip of X-spike in 2D-AGE appeared to be consistent with replication fork stalling at both sides of dCas9-bound sites, although we should note that it may also reflect entanglements between sister chromatids (Figure 4C). It remains to be seen in future studies whether dCas9 is equally effective in blocking the replication fork approaching from either direction.

Aside from the professional fork-blockers, proteins such as LacI and TetR can impair replication fork progression when bound to highly iterated arrays of *lacO* and *tetO*, respectively (60). We investigated whether even a single molecule of dCas9 can serve as a sufficiently strong barrier to induce genomic instability. To address this issue, we repurposed the classical genetic assay using a *URA3*-bearing *CUP1* array to sensitive detection of focal genomic instability induced by a single molecule of *URA3*-targeted dCas9 (Figure 2). The results showed that the binding of even a single molecule of dCas9 can destabilize the array. If each dCas9 molecule independently destabilizes the *CUP1* array, then targeting of dCas9 to each repeat unit multiplies the destabilizing effects, thereby leading to rapid contraction of the array.

Whether the replisome and dCas9 collide on the genome and have a mutual contact remains unclear. As the replisome approaches dCas9 in vivo, the torsional stress between their binding sites likely increases and may finally prevent replisome progression. This could explain why 2D-AGE indicated replication fork stalling around the midpoint of KpnI fragment, although the dCas9-bound sites were slightly off-centered (Figure 4). A higher-resolution method for mapping the position of the stalled replication fork is required to address this issue in future studies.

Cellular responses to dCas9-mediated replication fork stalling

When the replication fork encounters with an obstacle on DNA, the stability of the former and the removal of the latter should be critical. Ctf4 was demonstrated to protect arrested replication forks against breakage to suppress genome rearrangements, including hyper-amplification of rDNA (47). Mrc1 interacts with Tof1–Csm3 to form the heterotrimeric fork protection complex. Deletion of CTF4 and MRC1 significantly accelerated dCas9-induced reduction of the CUP1 copy number (Figure 5A). In the $ctf4\Delta$ and $mrc1\Delta$ strains, dCas9-mediated replication fork stalling

appeared to be diminished compared to the wild-type strain (Supplementary Figure S4E–G), presumably reflecting the breakage of destabilized replication forks. The accessory helicase Rrm3 is responsible for the removal of obstacles in front of the replisome. In the $rrm3\Delta$ strain, the VI was significantly increased and dCas9-mediated replication fork stalling appeared to be enhanced (Figure 5A, B and Supplementary Figure S5G). Consistently, depletion of Tof1, which counteracts Rrm3, resulted in a modest but significant decrease of the VI (Figure 5A). These data collectively underscored the importance of replisome protection and dCas9 removal in tandem repeat stability.

Despite the activities for fork protection and obstacle removal, stalling is occasionally prolonged to result in fork collapse. Cells have various mechanisms to cope with collapsed forks, which likely induce CUP1 CNV. Our genetic analysis indicated that RAD52 and its paralog RAD59 have the largest and second-largest contributions to CUP1 CNV, respectively (Figure 5A). Genetic analysis using separationof-function alleles indicated that Rad52 destabilizes tandem repeats via its SSA activity, but not its mediator activity to exchange RPA for Rad51 (Figure 5D). Although Rad52 and Rad59 mediate SSA, Rad52 but not Rad59 can perform this function in the presence of RPA (61). Based on the result of rad52 allele encoding a protein defective in RPA binding, we assumed that Rad52 mediates the annealing of RPA-coated ssDNA. Note that Rad52 was dispensable to restore rDNA copy number in the absence, but not the presence, of histone chaperone Asf1 (62). It would be intriguing to examine whether the requirement of Rad52 for dCas9induced CUP1 CNV is mitigated in the absence of Asf1.

Conventional SSA occurs after DSB and subsequent end resection. However, qPCR failed to provide evidence for DSB around dCas9-bound sites. Similarly, time-lapse imaging failed to reveal significant difference in Rfa1 focus formation, indicative of DSB, between the strains with CUP1targeted dCas9 and with no sgRNA. Thus, we have so far not obtained clear evidence for dCas9-induced DSB. The decease of VI in the $exol \Delta$ strain defective in end resection was less significant compared to the $rad52\Delta$ and $rad59\Delta$ strains (Figure 5A). It is conceivable that dCas9 induces destabilization without forming prominent DSBs, such as those generated by replication fork breakage. In this context, a new mechanism termed inter-fork strand annealing (IFSA) has attracted our attention. IFSA explains the interrepeat recombination induced by Rtf1/RTS1 system in fission yeast, involves Rad52 and Exo1 but not Rad51, and occurs without replication fork breakage (63). An IFSAlike mechanism may operate in budding yeast to mediate dCas9-induced CNV of tandem repeat units. Alternatively, tandem repeat structure may help SSA-mediated DSB repair to proceed too quickly to be detected with conventional approaches. Interstitial deletions occasionally found by nanopore sequencing may indicate at least a limited involvement of DSB in copy number alterations (Supplementary Figure S3C).

Less prominent but significant effects on destabilization were observed in the $mms2\Delta$ and $rad6\Delta$ strains (Figure 5A). Both MMS2 and RAD6 encode components of the errorfree TS pathway (64). However, the $mms2\Delta$ and $rad6\Delta$ strains exerted mutually opposite effects. Moreover, depletion of the other components of this pathway (Rad18,

Ubc13 and Rad5) failed to have significant effects on VI. Since all these proteins are involved in ubiquitination of proliferating cell nuclear antigen (PCNA), it is intriguing to examine the ubiquitination-defective PCNA mutant (Pol30-K146R). In anyway, we assumed that TS has little if any contribution to dCas9-induced tandem repeat destabilization. Similarly, HR, NHEJ, BIR and TLS did not appear to play major roles because no significant change of VI was observed in the $rad51\Delta$, $dnl4\Delta$, $pol32\Delta$ and $rev1\Delta$ strains, respectively (Figure 5A).

Taken together, it remains to be seen in future studies how Rad52 and Rad59 mediate the cellular response to dCas9mediated stalling of replication fork. It is also intriguing to examine the response in other species, including mammals, in which the preference in the choice of recombinational repair pathways may be different from that in the budding yeast.

Potential risk and application of dCas9-mediated replication fork stalling

This work has identified a potential risk of dCas9, distinct from the previously reported mutagenicity of the R-loop (3). For instance, for the sake of sensitivity, live-cell imaging studies often target dCas9 fused or complexed with fluorescent proteins to tandem repeats. Extended cultivation of such cells may result in contraction of the targeted tandem repeats, leading to not only compromised sensitivity but also an unexpected outcome. Even at a single-copy target site, dCas9 can impede replication fork progression and may thus induce SVs. This is especially true when recombinogenic genomic features are present around the target site, as was in the case for the URA3 cassette integrated in the CUP1 array (Figure 2). In this context, it is intriguing to note that the results of our genetic analysis (Figure 5) suggests a potential utility of Rad52 inhibitors (65,66) in reducing the risk of dCas9-induced focal genomic instability, albeit at the expense of general defects in various types of recombination.

Conversely, our findings imply that dCas9 provides a versatile tool for impeding replication fork progression at the genomic site of interest in vivo. Indeed, controlled replication fork stalling can accelerate mechanistic studies on genome stability. For this purpose, the Tus/*Ter*-system has been successfully used in both yeast and mammalian cells (67,68). However, this system requires its users to integrate Ter sequences into the regions of interest. In contrast, dCas9 is readily targetable to virtually any genomic regions by simply designing appropriate sgRNAs. Moreover, since dCas9mediated replication fork stalling works without modifying the genomic sequence, it would enable recapitulation of natural SV generation, thus providing a novel approach for modeling evolution and pathogenesis. Initial amplification of a single-copy gene likely involves mechanisms such as re-replication-induced gene amplification (69) and origindependent inverted repeat amplification (70). In both mechanisms, the borders of amplified regions are defined by the positions of replication fork collapse. We therefore expect that dCas9-mediated replication fork stalling provides a versatile tool to manipulate SVs including gene duplication, a critical driver of evolution.

DATA AVAILABILITY

The source code of DNA Seq Detector used in this study is available at GitHub (https://github.com/poccopen/DNA_Sequence_Detector). Nanopore sequencing data used in this study were deposited in DRA under accession number DRA010708.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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Conflict of interest statement. None declared.

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Supporting Information

Catalytically inactive Cas9 impairs DNA replication fork progression to induce focal genomic instability

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SUPPLEMENTARY FIGURE LEGENDS

Supplementary Figure S1. dCas9-induced copy number reduction of tandem repeat units.

- (A) CUP1 copy number of strains with constitutive expression of CUP1- and TEF1-targeted dCas9. Both strains use SNR52 and CSE4 promoters to constitutively express sgRNA and dCas9, respectively. Data are represented as the mean of two biological replicates.
- (B) Time course of CUP1 copy number of the strains used in Supplementary Figure S1A.
- (C) Effect of dCas9 induction on cell growth. Growth in the presence of 0, 10, and 100 nM β -estradiol were examined for the strains using *CUP2* or *ACT1* promoter for GEV expression (n = 1). Doubling time was determined from the growth curve drawn based on optical density measured every 10 min.
- (D) Time course of *CUP1* copy number. Similar to Figure 1B, except that culture was extended to day 9 (n = 1).
- (E) Dot plots of a nanopore sequencing read spanning the entire *ENA1* array. Top, self dot plot for the read. Bottom, dot plot between the *ENA1* reference sequence of *ENA1* (vertical axis) and the read (horizontal axis).
- (F) Effects of NAM on dCas9-induced *CUP1* copy number reduction in the WT and $rtt109\Delta$ strains. Similar to Figure 1B (n = 3 or more biological replicates).

Supplementary Figure S2. Destabilization of *URA3*-bearing *CUP1* array by *URA3*-targeted dCas9.

- (A) Dot plots between a nanopore sequencing read spanning the *URA3*-bearing *CUP1* array (horizontal axis) and the reference sequence of *CUP1* repeat unit (RU) or *URA3* cassette (vertical axis).
- (B) qPCR of CUP1 and URA3 in DNA isolated en masse from 5-FOA-resistant colonies.

Supplementary Figure S3. dCas9-induced expansion of CUP1 array.

- (A) Computational counting of CUP1 repeat units in nanopore reads. Boxed are the tandemly iterated regions identified by DNA Sequence Detector software. Dot plots between the reads and the reference sequence of CUP1 repeat unit are shown on the top. Note that the software correctly identifies the tandemly iterated regions from a nanopore read.
- (B) Population structure of the *CUP1* array. Similar to Figure 3C, except that copy numbers are computationally counted with DNA Sequence Detector.
- (C) Various structures of *CUP1* array revealed by nanopore sequencing. Dot plots between the *CUP1* reference sequence (vertical axis) and individual reads (horizontal axis) are shown. Arrows indicate the positions of interstitial deletions.
- (D) Histogram of nanopore read lengths after log transformation. Green vertical lines indicate the length of the wild-type *CUP1* array composed of 16 repeat units (~32 kb).

Supplementary Figure S4. Genetic analysis of genes involved in dCas9-induced CUP1 CNV.

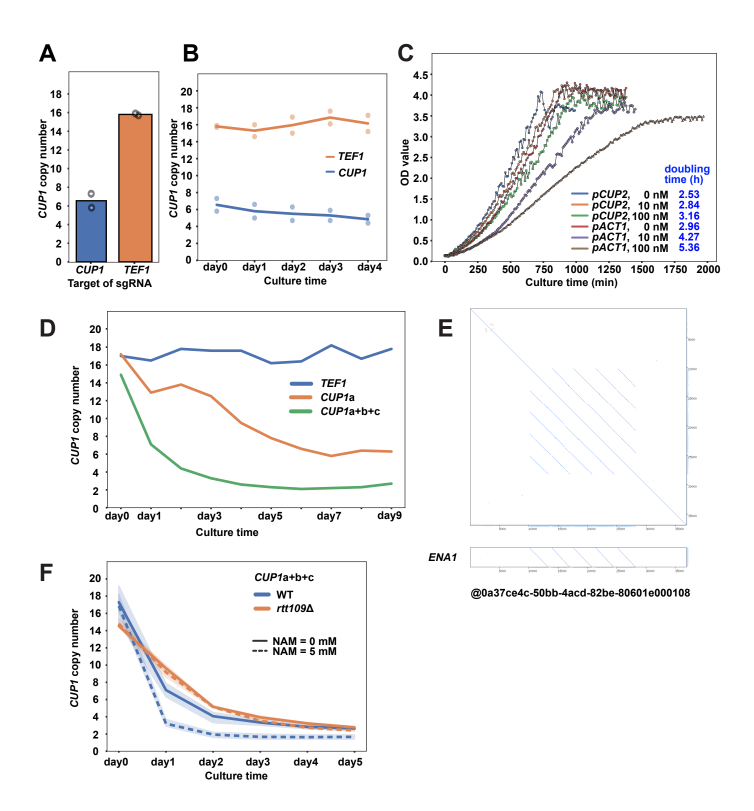
- (A) Percentile decrease of *CUP1* copy number in the strains used in Figure 5A.
- (B) Number of cell division in the strains used in Figure 5A.
- (C) Initial CUP1 copy number at day 0 in the strains used in Figure 5A.
- (D) CUP1 copy number change in the WT, $ctf4\Delta$, $rrm3\Delta$, and $mrc1\Delta$ strains with TEF1-targeted dCas9 (n = 2 or more biological replicates).
- (E) 2D-AGE images of KpnI-digested DNA fragments of the WT, $rrm3\Delta$, $ctf4\Delta$, and $mrc1\Delta$ strains. Panels labeled as WT and $rrm3\Delta$ #1 are identical to those in Figure 5B. Panels labeled as $rrm3\Delta$ #2 show biological duplicate data on the $rrm3\Delta$ strain. Exposure time for signal detection was 1 h.
- (F) Strategy for quantitative evaluation of replication fork stalling. 2D-AGE images identical to those in Figure 5B are shown as example. Two images were obtained by exposing the same membrane for 1 h and 1 min. The Stall Spot appeared on the Y-arc in the presence of dCas9 (i.e., stalled replication fork) was quantified from the 1-h exposure image as described in Methods. Similarly, the 1N Spot including the linear ~2-kb fragment was quantified from the 1-min exposure image. We divided the Stall Spot intensity by the 1N Spot intensity on the same membrane, thereby normalizing the amount of DNA prior to comparison among gels.
- (G) Quantitative comparison of replication fork stalling among the WT, *rrm3*Δ, *ctf4*Δ, and *mrc1*Δ strains. The Stall Spot was quantified from the gels shown in Supplementary Figure S4E according to the method described in Supplementary Figure S4F. Top panel indicates the simple ratio of Stall Spot against 1N spot, whereas the bottom panel indicates the background-subtracted ratio. The values in each mutant was normalized to that of WT (n =2 or more biological replicates). Note that replication fork stalling was enhanced in *rrm3*Δ but diminished in *ctf4*Δ and *mrc1*Δ strains.

Supplementary Figure S5. Roles for Rad52 in dCas9-induced CUP1 CNV.

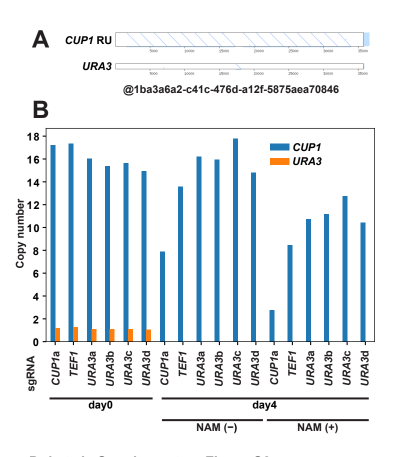
- (A) Alteration of *CUP1* copy number in the strains lacking both Rad52 and Rad59. The *rad52*Δ *rad59*Δ strains #1, #2, #3, and #4 (SY7-1, -2, -3, and -4) were generated from the *rad52*Δ strain by Cas12a-mediated gene editing with crRNAs *RAD59*g1, g6, g6, and g8, respectively (Supplementary Table S1 and S4). The *rad59*Δ *rad52*Δ strains #1 and #2 (SY7-6 and -7) were generated from the *rad59*Δ strain by Cas12a-mediated gene editing with crRNA *RAD52*g2 (Supplementary Table S1 and S4). The size and position of deletion in each strain is indicated in Supplementary Table S1. Data are represented as mean ± standard deviation (n = 3 or more biological replicates). Statistical significance was examined between the *rad52*Δ strain and the other strains using t-test (*p<0.05).
- (B) Percentile decrease of CUP1 copy number in the strains used in Supplementary Figure S5A.
- (C) Number of cell division in the strains used in Supplementary Figure S5A.
- (D) Initial CUP1 copy number at day 0 in the strains used in Supplementary Figure S5A.
- (E) Western blot analysis of Rad52 proteins expressed from a centromeric plasmid in the rad52Δ strain. Note that Rad52 proteins were C-terminally FLAG-tagged for the detection with anti-FLAG antibody.

Tubulin- α was used as a loading control for each sample.

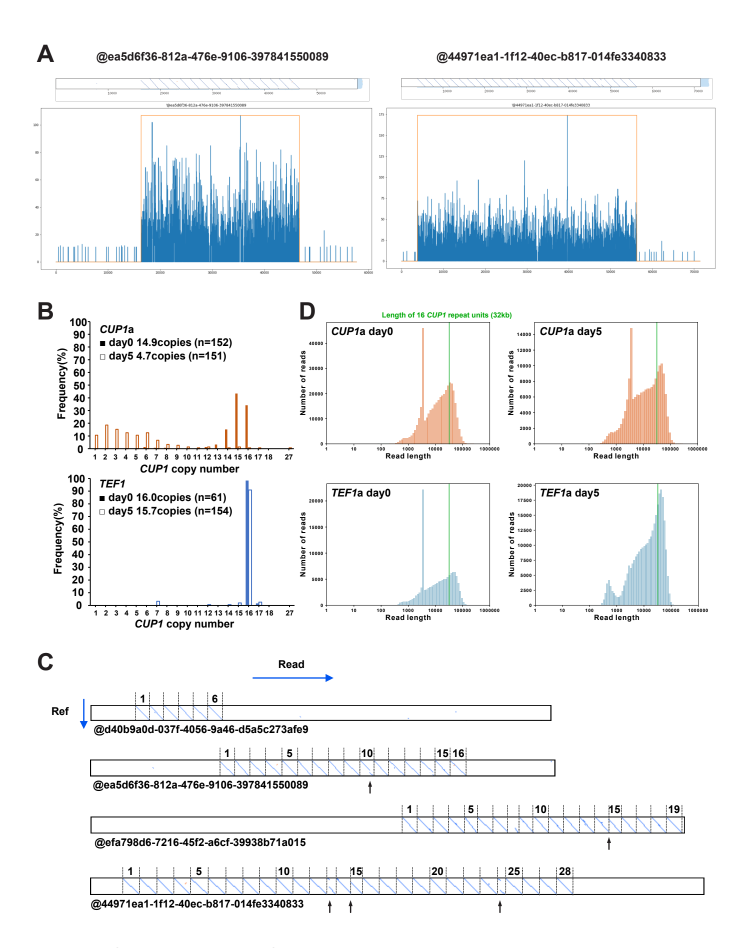
- (F) Percentile decrease of *CUP1* copy number in the strains used in Figure 5D.
- (G) Number of cell division in the strains used in Figure 5D.
- (H) Initial CUP1 copy number at day 0 in the strains used in Figure 5D.



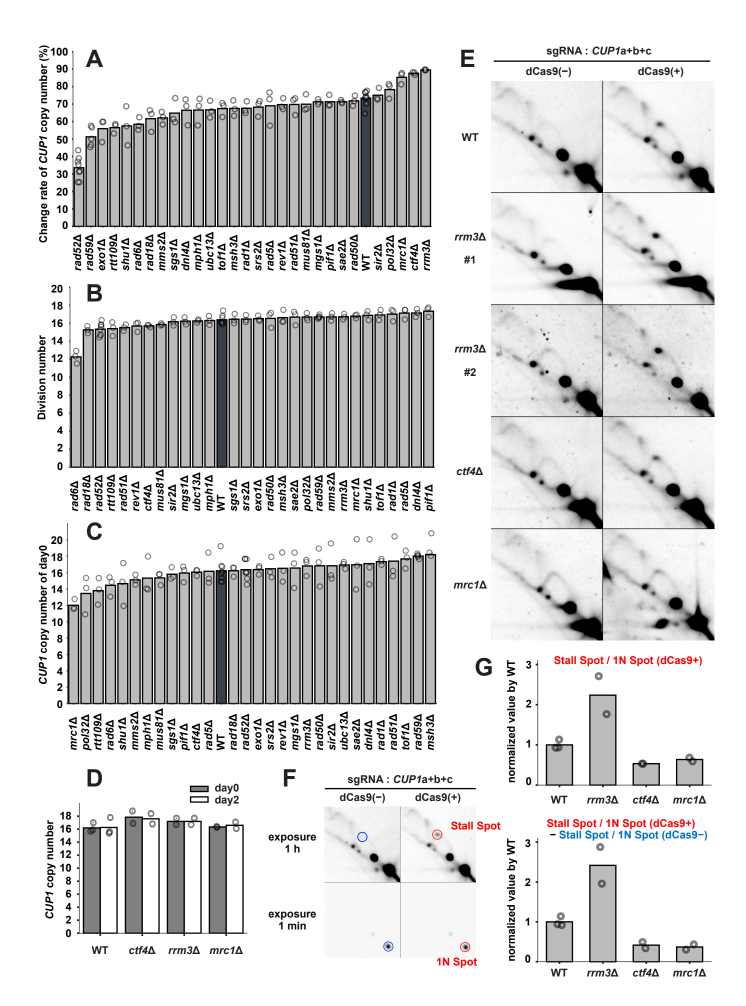
Doi et al., Supplementary Figure S1



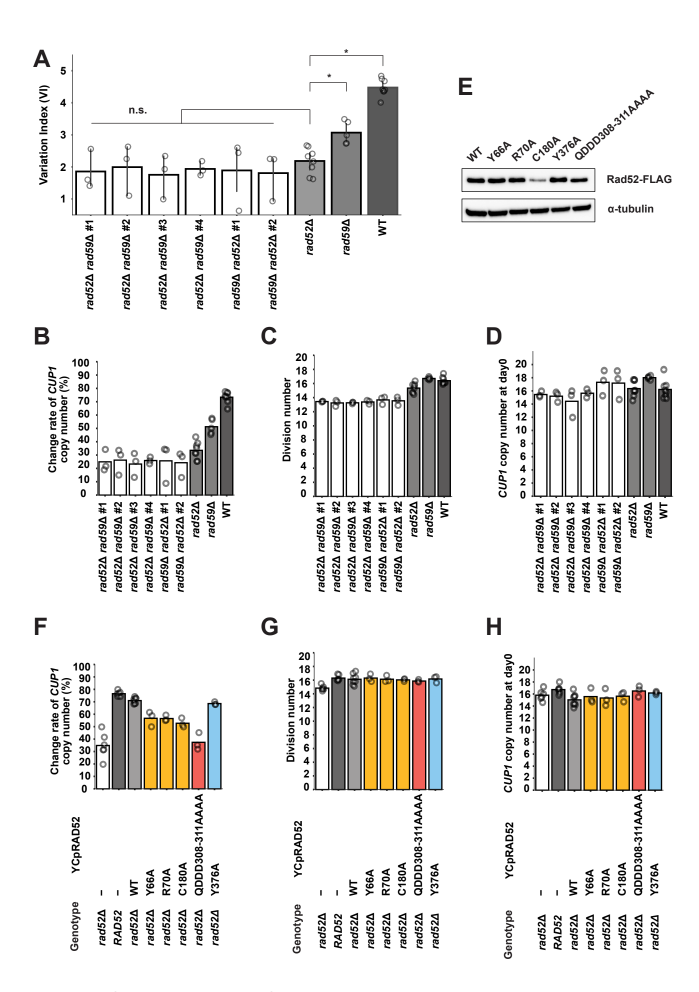
Doi et al., Supplementary Figure S2



Doi et al., Supplementary Figure S3



Doi et al., Supplementary Figure S4



Doi et al., Supplementary Figure S5

Supplementary Table S1. Yeast strains used in this study.

Strain name	Lab strain number	Genotype	Strain background	Source	Figure
BY4741	YIT5001	MATa his341 leu2 Δ0 met/5Δ0 ure3Δ0	BY4741	(Brachmann et al., 1998)	riguie
D25-8	YIT6523	pCSE4-dCas9-iADH1:LEU2 [pSNR52-HHribo-sqRNAFE(TEF1)-tSUP4-CU]	BY4741	This study	S1A, S1B
D25-12	YIT6527	pCSE4-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(CUP1a)-tSUP4-CU]	BY4741	This study	S1A, S1B
D54-1	YIT6865	pACT1-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2	BY4741	This study	S1C
D54-2	YIT6941	pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2	BY4741	This study	S1C
D68-8	YIT6972	pCUP2-yGEV-tADH1:HphiMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(TEF1)-tSUP4-CU]	BY4741	This study	1B, 1F, S1D, 3C, S3B, S3D, 4C, 4D
D68-9	YIT7087	pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(CUP1a)-tSUP4-CU]	BY4741	This study	1B, 1E, 1F, S1D, S1E, 3A, 3B, 3C, S3A, S3B, S3C, S3D, 4A
D68-15	YIT7702	pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(CUP1b)-ISUP4-CU]	BY4741	This study	1B
D68-16 D68-7	YIT7703 YIT6971	pCUP2-yGEV-tADH1-HphMX pGAL1-tiCas9-tADH1:LEU2 [pSNr52-HHribo-sgRNAFE(CUP1c)-tSUP4-CU] pCUP2-yGEV-tADH1-HphMX pGAL1-tiCas9-tADH1:LEU2 [pSNr52-HHribo-sgRNAFEs-tSUP4-CU](CUP1s+b+c)]	BY4741 BY4741	This study This study	1B
D68-12	YIT7219	pour-zyce-vount injimin poni taless-muni izeu (ponica-ininosgininez-sour-e-o(our istue)) pour-zyce-vount injimin poni taless-muni izeu (ponica-ininosgininez-sour-e-o(our istue)) pour-zyce-vount injimin poni taless-muni izeu (ponica-ininosgininez-sour-e-o(our istue))	BY4741	This study This study	1B, 1F, S1D, S1F, 4C, 4D, S4A, S4B, S4C, S4D, S4F, S4G, 5A, 5B, 5D, S5A, S5B, S5C, S5D, S5F, S5G, S5H
D68-13	YIT7220	pCUP2-yGEV-4Dh11:HpMX pGA1.1-dGSS-4DH11:EU2 [pGA1.1-HHnito-sgRNAFE(CUP1d)-4DH1-CU]	BY4741	This study	10
D68-14	YIT7268	pCUP2-yGEV-4ADH::hphMX pGAL1-4Cas9-4ADH::LEU2 [pGAL1-Hhribo-sgRNAFEs-1ADH1-CU(CUP1s+d)]	BY4741	This study	1C
D82-13	YIT7375	pCUP2-yGEV-4ADH1:HghMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU[ENA1a-b+c)]	BY4741	This study	1C
D84-0D	YIT7576	CUP1::URA3 pCUP2-yGEV-iADH1:HphMX pGAL1-dCas9-iADH1:LEU2	BY4741	This study	S2A
D84-9	YIT8148	CUP1::URA3 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(CUP1a)-tSUP4-CH]	BY4741	This study	2A, 2C, S2B
D84-10	YIT8149	CUP1::URA3 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 (pSNR52-HHribo-sgRNAFE(TEF1)-tSUP4-CH]	BY4741	This study	2B, 2C, S2B
D139-1	YIT8867	CUP1::URA3 pCUP2-yGEV-tADH1:HphMX pGAL1-tlCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(URA3a)-tSUP4-CH]	BY4741	This study	2B, 2C, S2B
D139-2	YIT8868	CUP1::URA3 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(URA3b)-tSUP4-CH]	BY4741	This study	2B, 2C, S2B
D139-3	YIT8869	CUP1::URA3 pCUP2-yGEV-4ADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(URA3e)-tSUP4-CH]	BY4741	This study	2B, 2C, S2B
D139-4 C72-1	YIT8870 YIT7202	CUP1::URA3 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(URA3d)+SUP4-CH]	BY4741 BY4741	This study	2B, 2C, S2B
C72-1 C72-2	YIT7202 YIT7203	rad5∆::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 (pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)] rtt109∆::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 (pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c))	BY4741 BY4741	This study This study	S4A, S4B, S4C, 5A S1F, S4A, S4B, S4C, 5A
C72-2 C72-3	YIT7204	ITTUBLINAND PULVE-YIGEV-VADUHT-INDIANN FORLT-T-GUSSIN-PUNTILEUZ (IDNINSZ-HITTIDO-SIGNUNES-SI-SUN-EU-(LUVIT-SI-SHO-I)) TIMBEL TENNING FOUPE-YIGEV-VADUHT-INDIANN FORLT-T-GUSSIN-PUNTILEUZ (IDNINSZ-HITTIDO-SIGNUNES-SI-SUN-EU-(LUVIT-SI-SHO-I)) TIMBEL TENNING FOUPE-YIGEV-VADUHT-INDIANN FORLT-T-GUSSIN-PUNTILEUZ (IDNINSZ-HITTIDO-SIGNUNES-SI-SUN-EU-(LUVIT-SI-SHO-I))	BY4741	This study This study	S1F, S4A, S4B, S4C, SA S4A, S4B, S4C, SA
C72-4	YIT7205	Iminas a manimos pour ayout vinoum rispinino pour monassimos manimos pour a manimos pour a pour a manimos pour pour manimos pour ma	BY4741	This study	S4A, S4B, S4C, 5A
C72-5	YIT7215	rad18_KanMX6 pCUP2-yGEV-4ADH.14phMX pGAL 1-Class-4ADH1:LEU2 [pSNR52-HHribo-sgRNAFE-t-SUP4-CU[CUP1a-b-c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-6	YIT7216	srs2 &::KanMX6 pCUP2-yGEV-IADH1:HphMX pGAL1-dCas9-IADH1::EU2 [pSNR52-HHribo-sgRNAFEs-ISUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A. S4B. S4C. 5A
C72-7	YIT7217	ubc13 b::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribb-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-8	YIT7218	rad6Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-9	YIT7230	rad51 b.:KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-10	YIT7231	sir2 L::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-11	YIT7232	mgs1Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-12	YIT7233	mph 1 D.: KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs+SUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-13	YIT7264	rm3Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU[CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, S4E, S4G, 5A, 5B
C72-14 C72-15	YIT7265 YIT7266	sgs1 \(\text{SanMX6}\) pCUP2-yGEV-ADH1:HphMX\) pGAL1-dCas9-ADH1:LEU2\(\text{pSNR52-HHribo-sgRNAFEs-ISUP4-CU(CUP1a+b+c)}\) shu1\(\text{\text{SanMX6}\) pCUP2-yGEV-ADH1:HphMX\) pGAL1-dCas9-ADH1:LEU2\(\text{pSNR52-HHribo-sgRNAFEs-ISUP4-CU(CUP1a+b+c)}\)	BY4741 BY4741	This study This study	S4A, S4B, S4C, 5A S4A, S4B, S4C, 5A
C72-15	YIT7267	SIGLI EL REMINA DI DUTZ-YGEV-BUTH I INJURAN DAGI. I-06-855-BUTH I ELEV (DISTINSE-THIRDES)GIVENDO SIGNI PER SISUPE-CU(CU) 1449-E)] INTERNAS DUTZ-YGEV-BUTH I INJURAN DAGI. I-16-855-BUTH I ELEV (DISTINSE-THIRDES)GIVENDO SIGNI PER SISUPE-CU(CU) 1449-E)]	BY4741	This study This study	S4A, S4B, S4C, S4E, 5A
C72-17	YIT7329	rad52 A: KanMX6 pCUP2-yGEV-tADH1-HpMX7 pGAL1-dCas9-tADH1:EU2 [pSNR52-HHribo-sqRnAFE-tSUP4-CU(CUP1a-bb-c)]	BY4741	This study	S4A, S4B, S4C, 4F, 5A, 5D, S5A, S5B, S5C, S5D, S5F, S5G, S5H
C72-18	YIT7330	rad50\text{\text{2.5}}'\text{\text{LMNX6}} pCUP2-yGEV-tADH1:HphMx pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sqRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A. S4B. S4C. 5A
C72-19	YIT7331	sae2 Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-20	YIT7332	rev1∆::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-21	YIT7479	dnl4 \(\text{\Lambda} \) L:KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-22	YIT7480	rad1 \(L::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)] \)	BY4741	This study	S4A, S4B, S4C, 5A
C72-23	YIT7481	msh3Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-24	YIT7652	pif1 Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A
C72-25 C72-27	YIT7653	rad27_A_KanMX6_pCUP2-yGEV-tADH1:HphMX_pGAL1-dCas9-tADH1:LEU2_[pSNR32+HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741 BY4741	This study This study	S4A, S4B, S4C, 5A
C72-28	YIT8656 YIT8657	rad59∆::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNr52-HHribo-sgRNAFEs-tSUP4-CU(CUP1s+b+c)] mrc1∆::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNr52-HHribo-sgRNAFEs-tSUP4-CU(CUP1s+b+c)]	BY4741	This study	S4A, S4B, S4C, 5A, S5A, S5B, S5C, S5D S4A, S4B, S4C, S4E, S4G, 5A
C72-29	YIT8658	IND I EXEMBNO POR PSET VEND I THOMAS PARE TROOS PROFILED INTO SERVICE IN HOUSE AND LESSOFT POOL PROFILED IN THE PROFILED IN TH	BY4741	This study	S4A, S4B, S4C, 5A
C72-30	YIT8659	sml1 \(\triangle \)-KanMX\(6\) pCUP2-yGEV-4DH1:HphMX pGAL1-0Cas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A. S4B. S4C. 5A
C72-31	YIT8699	mus81 \(\textit{\alpha}\)	BY4741	This study	S4A, S4B, S4C, 5A
C72-34	YIT8700	ctf4Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)]	BY4741	This study	S4A, S4B, S4C, S4E, S4G, 5A
D143-13	YIT9203	rm3∆::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE[TEF1]-tSUP4-CU]	BY4741	This study	S4D
D143-28	YIT9204	mrc1Δ::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(TEF1)-tSUP4-CU]	BY4741	This study	S4D
D143-34	YIT9205	ct/4\Delta:KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFE(TEF1)-tSUP4-CU]	BY4741	This study	S4D
D93-17	YIT7799	rad52 D.:KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(WT)-tRAD52]	BY4741	This study	5D, S5F, S5G, S5H
D96-1 D96-2	YIT7907 YIT7908	rad52 \(\text{AsmMX6}\) pCUP2-yGEV-4ADH1-HphMX\(\text{pGAL1-dCas9-VADH1:LEU2\(\text{pSNR52-HHribo-sgRNAFEs-\sUP4-CU(CUP1s+b+c)-pRAD52-RAD52\(\text{Y66A}\)-RAD52\(\text{Y66A}\)-RAD52\(\text{TabMX}\) pGAL1-dCas9-VADH1:LEU2\(pSNR52-HHribo-sgRNAFEs-\sUP4-CU(CUP1s+b+c)-pRAD52-RAD52\(\text{RAD52\(\text	BY4741 BY4741	This study	5D, SSF, S5G, S5H 5D, SSF, S5G, S5H
D96-2 D96-3	YIT7909	Idazia I. nalimiko jourz-yisez-vinuri. Inpiniko podu. I-leusas-vinuri. Letuz (piximoz-miniur-signivin-ta-si-sur-vin-tu)(upi interher)-pixiusz-vinuzieri, vinurusz-jiximoz- datisa I.: Kankiko (pUPz-yisez-vinuti). Kanki podu. I-leusas-vinuti i. Letuz (piximoz-miniur-ta-si-sur-vinutus) (piximoz-vinutusz) (piximoz-vinutu	BY4741	This study This study	5D, S5F, S5G, S5H
D96-7	YIT8036	lauce Insuland pour angle mental in pamin pour inclusion pour incl	BY4741	This study	5D, S5F, S5G, S5H
D96-8	YIT8544	laduz Inamino por 2-yez-vicum i njimin porirucasa-indini iLLUz posiniza-i il illuzagivini Estabi-vicul izadiz Cultura Cultura il illuzagivini Estabi-vicul illuzagivini E	BY4741	This study	5D. S5F. S5G. S5H
D132-1	YIT8692	rad52 b.:KanMX6 pCUP2-yGEV-IADH1:HphMX pGAL1-dCas9-IADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU[CUP1a+b+c]-pRAD52-RAD52[WT]-FLAG-IRAD52]	BY4741	This study	S5E
D132-2	YIT8693	rad52 b.::KanMX6 pCUP2-yGEV-tADH1:HphmX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(Y66A)-FLAG-tRAD52]	BY4741	This study	S5E
D132-3	YIT8694	rad52 L::KanMX6 pCUP2-yGEV-IADH1:HpMX pGAL1-dCas9-IADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(R70A)-FLAG-IRAD52]	BY4741	This study	S5E
D132-4	YIT8695	rad52 L::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(C180A)-FLAG-tRAD52]	BY4741	This study	S5E
D132-5	YIT8696	rad52 L::KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(Y376A)-FLAG-RAD52]	BY4741	This study	S5E
D132-6	YIT8697	radS2 D.:KanMX6 pCUP2-yGEV-tADH1:HphMX pGAL1-dCas9-tADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(QDDD308-311AAAA)-FLAG-tRAD52]	BY4741	This study	S5E
SY7-1	YIT9156	rad52Δ::KanMX6 rad59Δ(nt57-60) pCUP2-yGEV-IADH1:HphMX pGAL1-dCas9-IADH1:LEU2 [pSNR52-HHribo-sgRNAFEs-tSUP4-CU[CUP1a+b+c]]	BY4741	This study	S5A, S5B, S5C, S5D
SY7-2 SY7-3	YIT9157 YIT9158	rad52.1-KanMX6 rad5941_rit33-64] pCUP2-yGEV-4ADH+1-HpMXR pGAL-1-dCas9+ADH+1.EUZ [pSNR52.+HHrbo-spRNAFEs+ISUP4-CU[/CUP1a-b-c)] rad52.1-KanMX6 rad594(rit33-64) pCUP2-yGEV-4ADH+1-HpMX pGAL-1-dCas9+ADH+1.EUZ [pSNR52.+HHrbo-spRNAFEs+ISUP4-CU[/CUP1a-b-c)]	BY4741 BY4741	This study This study	S5A, S5B, S5C, S5D
SY7-4	YIT9158 YIT9159	/BBD22::ABMIXO FBD934(M3-9-9) PUUP-ZYGEV-RUHTI-II:PPMIX PD41E-10-889-PUHTI-ELU [D5NRO2-HIRIDO-SQRIVEN-ES-I-UPV-LU[CUPT-B-PO-C]] ### ### #### #### ##################	BY4741 BY4741	This study	S5A, S5B, S5C, S5D S5A, S5B, S5C, S5D
SY7-6	YIT9161	labusat.nammoi audusatinis-svoi puori-sijostinomini injumini puori-silostinomini injumini inj	BY4741	This study	S5A, S5B, S5C, S5D
SY7-7	YIT9162	rad594::KanMX6 rad524(nt50-54) pCUP2-yGEV-IADH1-IIphINX pCAL1-i-dCas9-IADH1-LEU2 [pSNR32-HHribo-sgRNAFEs-tSUP4-CU/(CUP1++b+c)]	BY4741	This study	S5A, S5B, S5C, S5D

Supplementary Table S2. Primers used in this study.

Primer name	Sequence	Description
RAD5-in-5	CTCATTCTGGACCTTTCTGC	
		RAD5 deletion
RAD5-in-3	TTGATCCCGACTGATGA	RAD5 deletion
RTT109-in-5	CCCGTTAAGTCGTAAACGTC	RTT109 deletion
RTT109-in-3	CTCTTTAGCAAATTGTGGCC	RTT109 deletion
MMS2-in-5	ACCCACCACTATTGCTCATT	MMS2 deletion
MMS2-in-3	GCAACAGTCTTTCTGTCGTC	MMS2 deletion
POL32-in-5	CGGCGCTAAGCTTAATTTTT	POL32 deletion
POL32-in-3	AACTAAATGGGATGACGCTG	POL32 deletion
RAD18-in-5	ATCAATTGCAGAAGAGGCAG	RAD18 deletion
RAD18-in-3	TCTTTGCCTAATCCAGTGGC	RAD18 deletion
SRS2-in-5	TCTAGCACTCCTTCCAACCA	SRS2 deletion
SRS2-in-3	CTTTGACCCAACGCCTTCTA	SRS2 deletion
UBC13-in-5	TGATTAGATTCTGCGGGAGC	UBC13 deletion
UBC13-in-3	TAGTGTAGGACGGTCGATCC	UBC13 deletion
RAD6-in-5	CCTGGATATGGTACCGATGT	RAD6 deletion
RAD6-in-3	CGGGTATCGGCAGTTATAAC	RAD6 deletion
RAD51-in-5	ATCACCGTCTCACCATATCC	RAD51 deletion
RAD51-in-3	AGGAAGTAGTCATCGGGAAG	RAD51 deletion
SIR2-in-5	TCTAGCACTCCTTCCAACCA	SIR2 deletion
SIR2-in-3	CTTTGACCCAACGCCTTCTA	SIR2 deletion
MGS1-in-5	AGAGCAAAACGGATTTGACA	MGS1 deletion
MGS1-in-3	GGTCGAGGAAAAGGAAGAAC	MGS1 deletion
MPH1-in-5	GCACTCAATTTGCAAGGTTG	MPH1 deletion
MPH1-in-3	GTGCTTCATGACAGAAAGGT	
		MPH1 deletion
RRM3-in-5	CCGGTCTATCTTCCCTTACC	RRM3 deletion
RRM3-in-3	AGGATCTGATTTCCCTCACT	RRM3 deletion
SGS1-in-5	TATAGTTCAGCCGTGCGTTT	SGS1 deletion
SGS1-in-3	TAACTGAGCAATGTGCACAC	SGS1 deletion
SHU1-in-5	CTCTCGGATTCACTTACCGA	SHU1 deletion
SHU1-in-3	TCTATGTAAGCATGCGTTGC	SHU1 deletion
TOF1-in-5	CGTTTGCTTTAGGAAGCTGT	TOF1 deletion
TOF1-in-3	CATGCGTCTTCAAGCAATGA	TOF1 deletion
RAD52-in-5	TATCATCTGCCTGCCCTGTA	RAD52 deletion
RAD52-in-3	CCGAGTTGCCATATTTGTAT	RAD52 deletion
RAD50-in-5	TAACTGCAGTAGAACCCACC	RAD50 deletion
RAD50-in-3	GAAAAGAACAACTAGCTGACG	RAD50 deletion
SAE2-in-5	GGGCCCATAAGTCTCTGATAA	SAE2 deletion
SAE2-in-3	CGAAAATAACGTCGACGTTC	SAE2 deletion
REV1-in-5	CAGACCAAGACGGAAAAAAG	REV1 deletion
REV1-in-3	CTTTCGGAATGGTGGACAAC	REV1 deletion
DNL4-in-5	AGATTCGCCGAGTATAGATCA	DNL4 deletion
DNL4-in-3	GCGATGAGGTGAAAGACAAA	DNL4 deletion
RAD1-in-5	TGAAGTGTTCTCTGTTTGCC	RAD1 deletion
RAD1-in-3	AGATTCCACCAAATACGGTT	RAD1 deletion
MSH3-in-5	CAATAGTGTTTCCCCGACTC	
		MSH3 deletion
MSH3-in-3	TGTACAAGGCCAAGGCATAA	MSH3 deletion
PIF1-in-5	TGCTGTTCAGCTCATCTCTT	PIF1 deletion
PIF1-in-3	GTGAGTTAGTCTCCTTTGGC	PIF1 deletion
RAD27-in-5	ACTATTGGCTTAGCTATGGC	RAD27 deletion
RAD27-in-3	TACGATGGTTCCGATATGCC	RAD27 deletion
	TATCGCGTTGGACACTAATG	RAD59 deletion
RAD59-in-5		
RAD59-in-3	CTTCGTTACCTTGGAATGGT	RAD59 deletion
MRC1-in-5	CACACACACTTTCGTTTAC	MRC1 deletion
MRC1-in-3	CCTACTCTTTTGCTCCA	MRC1 deletion
EXO1-in-5	AGGCGGGAAGTACAATAACT	EXO1 deletion
EXO1-in-3	CGCGCTCGCAAAACATATAA	EXO1 deletion
MMS2-in-5	ACCCACCACTATTGCTCATT	
		MMS2 deletion
MMS2-in-3	GCAACAGTCTTTCTGTCGTC	MMS2 deletion
SML1-in-5	CATTTTGGGAAGTCATCCGT	SML1 deletion
SML1-in-3	TTTCTCATTTGTCTCGGGCC	SML1 deletion
MUS81-in-5	ATTGGGCACCTTTATGTCC	MUS81 deletion
MUS81-in-3	GGTATCCAGAAGTCAAAGGT	MUS81 deletion
CTF4-in-5	AGGTGAAATCGTAGTTTGCC	CTF4 deletion
CTF4-in-3	GTACAGGACCAGGTGTTACA	CTF4 deletion
VIII214253::URA3-F	GCCTAACTTACAACTCTGTATAGAATCCCCAGATTTCGAATAAAAAAAA	URA3 cassette integration into CUP1 array
VIII214253::URA3-R	CAATAGAGGCAGGTATCGGAGATAGGTTTTCAGCAGCGGGTACCATGAATGGGTAATAACTGATATAATTAAATTGAAGC	URA3 cassette integration into CUP1 array
ACT1-F	CGCTGCTCAATCTTCTCAA	ACT1 quantification by qPCR
ACT1-R	GTAGTTTGGTCAATACCGGC	ACT1 quantification by qPCR
CUP1-F	CACGTATAGCGCCCAAATGT	CUP1 quantification by qPCR
CUP1-R	ATTGACAAGGATTTGGAATCTGA	CUP1 quantification by qPCR
ENA1-F	TCACTACTGACCGGTGAATC	ENA1 quantification by qPCR
ENA1-R	GACAATACCCTTGGCTCTTC	ENA1 quantification by qPCR
		L quantinoation by qr Oil
LIDAS E	0040044004474070040	UDAO
URA3-F	CCACCAAGGAATTACTGGAG	URA3 quantification by qPCR
URA3-F URA3-R	CCACCAAGGAATTACTGGAG TTGTACTTGGCGGATAATGC	URA3 quantification by qPCR URA3 quantification by qPCR
URA3-R	TTGTACTTGGCGGATAATGC	URA3 quantification by qPCR
URA3-R CUP1-probe-F1	TTGTACTTGGCGGATAATGC GGAAACGCAAGCAAAATTGA	URA3 quantification by qPCR CUP1 repeat unit probe

Supplementary Table S3. Plasmids used in this study

Number	Name	Plasmid type	Marker gene
4-30	Ylplac128-pCSE4-dCas9-tADH1	Integrative	LEU2
8-75	Ylplac128-pGAL1-dCas9-tADH1	Integrative	LEU2
9-38	pFA6a-pACT1-yGEV-tADH1-HphMX	Integrative	HygR
9-55	pFA6a-pCUP2-yGEV-tADH1-HphMX	Integrative	HygR
4-78	pSNR52-HHribo-sgRNAFE(TEF1)-tSUP4-CU	Centromeric	URA3
3-52	pSNR52-HHribo-sgRNAFE(CUP1a)-tSUP4-CU	Centromeric	URA3
3-49	pSNR52-HHribo-sgRNAFE(CUP1b)-tSUP4-CU	Centromeric	URA3
3-51	pSNR52-HHribo-sgRNAFE(CUP1c)-tSUP4-CU	Centromeric	URA3
4-41	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)	Centromeric	URA3
9-71	pGAL1-HHribo-sgRNAFE(CUP1a)-HDV-tADH1-CU	Centromeric	URA3
10-15	pGAL1-HHribo-sgRNAFE(CUP1d)-HDV-tADH1-CU	Centromeric	URA3
10-39	pGAL1-HHribo-sgRNAFE(CUP1a+d)-HDV-tADH1-CU	Centromeric	URA3
11-37	pSNR52-HHribo-sgRNAFEs(ENA1a+b+c)-tSUP4-CU	Centromeric	URA3
8-20	pSNR52-HHribo-sgRNAFE(TEF1)-tSUP4-CH	Centromeric	HIS3
12-79	pSNR52-HHribo-sgRNAFE(CUP1a)-tSUP4-CH	Centromeric	HIS3
15-76	pSNR52-HHribo-sgRNAFE(URA3a)-tSUP4-CH	Centromeric	HIS3
15-77	pSNR52-HHribo-sgRNAFE(URA3b)-tSUP4-CH	Centromeric	HIS3
15-78	pSNR52-HHribo-sgRNAFE(URA3c)-tSUP4-CH	Centromeric	HIS3
15-79	pSNR52-HHribo-sgRNAFE(URA3d)-tSUP4-CH	Centromeric	HIS3
12-51	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(WT)-tRAD52	Centromeric	URA3
13-11	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(Y66A)-tRAD52	Centromeric	URA3
13-12	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(R70A)-tRAD52	Centromeric	URA3
13-13	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(C180A)-tRAD52	Centromeric	URA3
13-65	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(Y376A)-tRAD52	Centromeric	URA3
14-63	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(QDDD308-311AAAA)-tRAD52	Centromeric	URA3
15-6	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(WT)-FLAG-tRAD52	Centromeric	URA3
15-7	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(Y66A)-FLAG-tRAD52	Centromeric	URA3
15-8	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(R70A)-FLAG-tRAD52	Centromeric	URA3
15-9	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(C180A)-FLAG-tRAD52	Centromeric	URA3
15-10	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(Y376A)-FLAG-tRAD52	Centromeric	URA3
15-11	pSNR52-HHribo-sgRNAFEs-tSUP4-CU(CUP1a+b+c)-pRAD52-RAD52(QDDD308-311AAAA)-FLAG-tRAD52	Centromeric	URA3
16-48	pGAL1-yenAsCas12a-tADH1-pGAL1-tRNA(Gly)-HindIII-AsCpf1gRNA(RAD59g1)-U4AU4-HDV-tCYC1-CU	Centromeric	URA3
16-53	pGAL1-yenAsCas12a-tADH1-pGAL1-tRNA(Gly)-HindIII-AsCpf1gRNA(RAD59g6)-U4AU4-HDV-tCYC1-CU	Centromeric	URA3
16-55	pGAL1-yenAsCas12a-tADH1-pGAL1-tRNA(Gly)-HindIII-AsCpf1gRNA(RAD59g8)-U4AU4-HDV-tCYC1-CU	Centromeric	URA3
16-57	pGAL1-yenAsCas12a-tADH1-pGAL1-tRNA(Gly)-HindIII-AsCpf1gRNA(RAD52g2)-U4AU4-HDV-tCYC1-CU	Centromeric	URA3

Supplementary Table S4. sgRNAs and crRNAs used in this study

sgRNA/crRNA	Protospacer	PAM	Chromosome	Strand	Start	End	Standard Name	Systematic Name	Start	End	Standard name	Systematic Name	Start	End	Standard name	Systematic Name	Source
CUP1a	TCTGACAATCCATATTGCGTTGG	TGG	chrVIII	+	214865	214887	CUP1-1	YHR053C	212867	212889) CUP1-2	YHR055C					This study
CUP1b	TATCAGCCTGAAATAAAGGGTGG	TGG	chrVIII	+	215024	215046	CUP1-1	YHR053C	213026	213026	CUP1-2	YHR055C					This study
CUP1 c	AGATATCAGCCTGAAATAAAGGG	GGG	chrVIII	+	215021	215043	S CUP1-1	YHR053C	213023	213045	CUP1-2	YHR055C					This study
CUP1 d	CTGTCAAGAGATTCTTTTGCTGG	TGG	chrVIII	-	214954	214976	CUP1-1	YHR053C	212956	212978	3 CUP1-2	YHR055C					This study
TEF1	TTGATATTTAAGTTAATAAACGG	CGG	chrXVI	+	700460	700482	TEF1	YPR080W									(Gilbert et al, 2013)
ENA1a	CGGAATATAGACGACGGGGAAGG	AGG	chrIV	+	531553	531575	ENA1	YDR040C	531553	531575	ENA2	YDR039C	527668	3 527690	ENA5	YDR038C	This study
ENA1b	TGATCGAAGCTTTACACCGTAGG	AGG	chrIV	-	532339	532361	ENA1	YDR040C	532339	532361	ENA2	YDR039C	528454	528476	ENA5	YDR038C	This study
ENA1c	ACCACGACTTACTAGGATCGCGG	CGG	chrIV	+	533817	533839	ENA1	YDR040C	533817	533839	ENA2	YDR039C	529932	2 529954	ENA5	YDR038C	This study
URA3a	CATTACGAATGCACACGGTG	TGG	chrV	+	116517	116536	URA3	YEL021W									This study
URA3b	GCACACGGTGTGGTGGGCCC	AGG	chrV	+	116527	116546	URA3	YEL021W									This study
URA3c	TTGTCATCTAAACCCACACC	GGG	chrV	-	116773	116792	URA3	YEL021W									This study
URA3d	TTGATTATGACACCCGGTGT	GGG	chrV	+	116758	116777	URA3	YEL021W									This study
RAD59g1	TATCCAAGCCCGGTGCTGTGCCG	TTTA	chrlV	+	344878	344900	RAD59	YDL059C									This study
RAD59g6	GAGCATATCGTATGATTCGACTA	GTTC	chrlV	-	344905	344927	RAD59	YDL059C									This study
RAD59g8	GTCGAATCATACGATATGCTCGA	TGTA	chrlV	+	344907	344929	RAD59	YDL059C									This study
RAD52 g2	GTCTGTATGTCCTCGGAATGGTT	TTTG	chrXIII	+	213863	213885	RAD52	YML032C									This study