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## Identification of 15 new bypassable essential genes of fission yeast

Aoi Takeda<sup>1</sup>, Shigeaki Saitoh<sup>2</sup>, Hiroyuki Ohkura<sup>3</sup>, Kenneth E. Sawin<sup>3</sup>, Gohta Goshima<sup>1,#</sup>

<sup>1</sup>Division of Biological Science, Graduate School of Science, Nagoya University, Furo-cho, Chikusa-ku, Nagoya 464-8602, Japan

<sup>2</sup>Division of Cell Biology, Institute of Life Science, Kurume University, Kurume, Fukuoka 830-0011, Japan

<sup>3</sup>Wellcome Centre for Cell Biology, School of Biological Sciences, University of Edinburgh, Michael Swann Building, Max Born Crescent, Edinburgh EH9 3BF, UK

### Abstract

Every organism has a different set of genes essential for its viability. This indicates that an organism can become tolerant to the loss of an essential gene under certain circumstances during evolution, via the manifestation of ‘masked’ alternative mechanisms. In our quest to systematically uncover masked mechanisms in eukaryotic cells, we developed an extragenic suppressor screening method using haploid spores deleted of an essential gene in the fission yeast *Schizosaccharomyces pombe*. We screened for the ‘bypass’ suppressors of lethality of 92 randomly selected genes that are essential for viability in standard laboratory culture conditions. Remarkably, extragenic mutations bypassed the essentiality of as many as 20 genes (22%), 15 of which have not been previously reported. Half of the bypass-suppressible genes were involved in mitochondria function; we also identified multiple genes regulating RNA processing. 18 suppressible genes were conserved in the budding yeast *Saccharomyces cerevisiae*, but 13 of them were non-essential in that species. These trends suggest that essentiality bypass is not a rare event and that each organism may be endowed with secondary or backup mechanisms that can substitute for primary mechanisms in various biological processes. Furthermore, the robustness of our simple spore-based methodology paves the way for genome-scale screening.

### Introduction

A recent genome-wide study using *S. cerevisiae* gave an insight into the ‘evolvability’ of essential cellular processes (Liu *et al.*, 2015), which can be also termed ‘bypass-of-essentiality’ (BOE) (Li *et al.*, 2019). The study surveyed the viability of every essential gene disruptant in *S. cerevisiae* (1,106 genes), and found that 9% of the gene disruptants proliferate and form colonies spontaneously (i.e. without artificial mutagenesis). Genome

<sup>#</sup>Correspondence should be addressed to: goshima@bio.nagoya-u.ac.jp; Phone: +81 52-788-6175.

#### Author contributions

G.G. conceived the project. G.G., S.S. H.O. and K.E.S. designed the research. A.T. and G.G. performed experiments. A.T., K.E.S. and G.G. analysed the data. S.S. and K.E.S. contributed resources. G.G. wrote the paper. S.S., H.O. and K.E.S. reviewed and edited the paper.

analysis showed that most of the proliferating strains had gained an extra chromosome (i.e. aneuploidy), which is typically an outcome of chromosome missegregation. This is a reasonable path to BOE in *S. cerevisiae*, because its haploid is tolerant to a chromosome gain for 13 of 16 chromosomes (Torres *et al.*, 2007). However, we speculated that there might be many more bypassable essential genes in yeast, as some non-bypassable essential gene disruptants might recover their viability by acquiring extragenic mutations, which are rarely introduced without mutagenesis.

Comprehensive identification of suppressor mutations would help to elucidate secondary or backup mechanisms that can substitute for primary mechanisms. Hitherto ‘masked’, these alternative mechanisms may act as the dominant pathways in specific cell types and/or diseased cells. To this end, we designed a BOE screening using the fission yeast *S. pombe*, which has a similar number (1,260) of essential genes to *S. cerevisiae* (Kim *et al.*, 2010). A notable difference from *S. cerevisiae* is that *S. pombe* has only 3 chromosomes, and the haploid yeast is inviable when an extra copy of either chromosome I or II (the two larger chromosomes) is inherited (Niwa and Yanagida, 1985). Thus, BOE via extra chromosome gain is likely an infrequent event in *S. pombe*. In the present study, we carried out BOE screening for randomly selected 92 essential genes in *S. pombe*, based on UV mutagenesis of spores in which essential genes were deleted.

## Materials and methods

### Yeast strains and media

A diploid named G29 was used as the host ( $h^+$  *his2 leu1 ura4-D18 ade6-216* /  $h^-$  *leu1 ura4-D18 ade6-210 rpl42.sP56Q*), where *rpl42.sP56Q* allele was used as a counter-selection marker against cycloheximide (Roguev *et al.*, 2007). Conventional genetic experiments followed (Moreno *et al.*, 1991). Yeast was grown on complete YE5S medium (YE supplemented with leucine, uracil, adenine, lysine, and histidine) or the synthetic PMG or EMM medium at 32°C (plate) or 30°C (liquid). Sporulation was induced on the SPA plate or in the EMMG liquid medium (i.e. PMG containing 1 g/L sodium glutamate instead of 3.75 g/L).

### Gene disruption

Essential genes were selected based on information found in the Pombase database (Wood *et al.*, 2012). 92 genes on chromosome II were randomly selected. Conventional one-step replacement was conducted using ~500-bp homologous sequences (5'UTR and 3'UTR of the gene to be deleted) (Krawchuk and Wahls, 1999). A tandem G418-resistance (kanMX) / *ura4+* cassette was used as the selection marker (however, *ura4+* marker was not actually used for selection). For most genes, we directly generated a linear construct (5'UTR-G418-*ura4+*-3'UTR) by two rounds of PCR using two sets of primers (i.e. nested PCR). In some cases, the PCR fragment was cloned into a vector using an Infusion kit (Takara), and the linear construct was amplified with T7/T3 primer set from the plasmid template. The linear DNA was transformed into the G29 diploid strain using the standard lithium acetate/PEG-mediated method, and disruption of the target gene was confirmed by colony PCR using KOD-Fx-Neo or KOF-ONE kit (Toyobo). When the endogenous gene and G418-*ura4+*

cassette had a similar length, we used a longer version of G418-*ura4+* cassette to distinguish disrupted and endogenous alleles by length. *hul5* was deleted with the hygromycin-resistance cassette. PCR primers for gene disruptions and their confirmation are listed in Table S1.

### Spore isolation

Exponentially growing heterozygous diploid cells in YE5S (+10 µg/ml G418) were harvested and transferred to EMMG medium ( $1 \times 10^6$  cells/ml). After shaking at 200 rpm and 30°C for 36 h, cells were harvested, washed twice with sterile water, and resuspended in 0.5% glusulase solution. The solution was shaken at 80 rpm at room temperature overnight to digest non-sporulated cells. The spores were harvested and treated with 30% ethanol for 30 min (80 rpm, room temperature) to further remove diploid cell contamination. The purified spores were resuspended in sterile water and stored at 4°C.

### Spore quality check

Prior to UV mutagenesis, the viability and purity of spores were determined by plating onto normal YE5S plate and YE5S supplemented with G418 (100 µg/ml) and cycloheximide (100 µg/ml), respectively. No haploid spores were expected to grow on the G418/cycloheximide plate, since an essential gene had been replaced with G418. However, colonies were always formed at  $>1 \times 10^{-6}$  frequencies. Cells in these colonies were diploids, which we interpreted to be derived from diploid spores generated at low frequency during meiosis; diploid spores would also be resistant to glusulase or ethanol. In cases in which the putative diploid contamination frequency was  $< 5 \times 10^{-4}$ , we moved on to UV mutagenesis and screening. In cases in which the contamination frequency was  $\geq 5 \times 10^{-4}$ , we discarded the sample and repeated the spore isolation process. The reason for differences in the prevalence of putative diploid spores is unknown.

### BOE screening with UV mutagenesis

$1 \times 10^7$  spores were plated onto a YE5S plate containing G418 (100 µg/ml) and cycloheximide (100 µg/ml), followed by UV irradiation ( $90 \times 100 \mu\text{J}/\text{cm}^2$ : UV Crosslinker, CL-1000, 254 nm, 100 V, 8 W [UVP/*Analytik Jena*] or Stratilinker UV crosslinker Model 1800 [Stratagene]). Under these conditions, spore viability was approximately 1%. Cycloheximide allows counter-selection against *rpl42+* gene; in the presence of cycloheximide, haploids possessing the *rpl42.sP56Q* allele can grow but not parental heterozygous diploids (Roguev et al., 2007). Plates were incubated at 32°C for 7 d. In most cases, we detected colonies. To check if each colony represents BOE or diploid contamination, we replica-plated onto EMM minus adenine and SPA plates. After checking spore formation by iodine treatment on SPA, we selected the Ade- and non-spore-forming colonies as candidate BOE haploids; colonies that did not match this criterion were likely diploids and disregarded. The candidate colonies were subjected to colony PCR, with which the disruption of the target gene was reconfirmed (see Fig. 2B). For *top3*, we performed mutagenesis in a *rad13* (DNA repair-deficient) background, in order to decrease UV power ( $1,500 \mu\text{J}/\text{cm}^2$ , 5% viability) and avoid cytotoxicity. However, since we obtained expected BOE results for *cut7* in the presence of *rad13+*, we did not introduce *rad13* for any other genes.

## Whole-genome sequencing and sequence analysis

To identify suppressor mutations, bulk segregant analysis was performed. Survivor strains were crossed with a wild-type strain, and spores were plated on G418-containing YE5S plates. After 7 d, ~1,000 colonies were collected and DNA was extracted with Dr. GenTLE (Takara). Genomic DNA (1 µg) was sequenced by BGI or Novogene (1 Gb), and the reads were mapped to the reference genome (Schizosaccharomyces\_pombe.ASM294v2.genebank.gb) using CLC Genomics Workbench. Unique and homogenous Indels and SNPs identified for each strain were manually inspected using *Integrative Genomics Viewer* (IGV).

## Results & Discussion

Fig. 1 illustrates the scheme of our BOE screening. A heterozygous diploid in which one copy of an essential gene was replaced with a drug (G418)-resistant marker was sporulated. The spores were plated on G418-containing plates and simultaneously mutagenized by UV irradiation. If a haploid colony is obtained on this plate, it has likely acquired a suppressor mutation(s), indicating that the essentiality of the gene has been bypassed.

We first applied this method to two gene disruptants, *cut7* (kinesin-5) and *top3* (type I topoisomerase), the lethality of which is known to be suppressed by the loss of function of Pkl1 (kinesin-14) and Rqh1 (recQ helicase), respectively (Goodwin *et al.*, 1999; Olmsted *et al.*, 2014; Syrovatkina and Tran, 2015). For *cut7*, we obtained a total of 8 haploid colonies in the first experiment and 30 more in a later experiment, in which 5-fold more spores were mutagenized (Fig. 2A, B). We analysed 26 colonies by target sequencing of *pk11* and *msd1* genes (Msd1 is a positive regulator of Pkl1 (Yukawa *et al.*, 2015)), whole-genome sequencing, and/or genetic linkage test (*pk11* locus is close to the *rpl42* locus, at which a mutation was introduced to confer cycloheximide resistance in our strain). The combined results suggested that suppressor mutations reside in *pk11* or *msd1* for 19 or 7 strains, respectively (Fig. 2C, E). Mutagenesis of *top3* yielded 3 haploid strains (Fig. 2D), and direct sequencing of the *rqh1* gene identified a mutation in all cases (Fig. 2E). Thus, our screening successfully elucidated known BOE relationships.

We then expanded the screening to 92 essential genes located on chromosome II. For 20 of these, we obtained 1~17 haploid colonies, which corresponds to 22% (Fig. 3). This frequency is much higher than that obtained in the previous mutagenesis-free screening in *S. cerevisiae* (Liu *et al.*, 2015).

The 20 suppressible genes possess divergent known biological functions. 10 genes (50%) were related to mitochondrial function (Fig. 4A). This may be partly explained by the fact that, in the regular medium containing >2% glucose, cell proliferation does not depend much on mitochondrial respiration (Takeda *et al.*, 2015). 6 genes were associated with RNA processing and ribosome functions; the basis of these trends are unclear. Overall, 90% of the genes had clear orthologues in *S. cerevisiae* and *H. sapiens*, indicating that BOE is not limited to unconserved genes (Fig. 4B, C). However, the orthologues of 70% genes were reported to be non-essential in *S. cerevisiae* (Fig. 4D). An obvious next step would be to

identify suppressor mutations of each survivor to understand how an essential mechanism can be bypassed.

In the course of this project, a conceptually identical study was published (Li et al., 2019). In this study using *S. pombe* haploid, BOE was screened by 3 methods: chemical mutagenesis (termed C-BOE), transposon-based mutagenesis (T-BOE), and gene library overexpression (OP-BOE). While C-BOE is the most similar approach to ours, the methodology employed is different. Li et al (2019) did not use spores; instead, the essential gene disruptant was kept viable by the transformation of a plasmid that contains the deleted gene: if a colony that had lost the plasmid was recovered, it was interpreted to indicate BOE. Li et al. (2019) obtained survivors for 27% of the genes in one or more BOE assays, which is a similar frequency to ours.

Coincidentally, in the two studies, 29 common essential genes were screened. Upon comparison, 21 genes were not bypassable in both studies, whereas 5 genes were common BOE hits. 2 and 1 hits were uniquely found in their and our studies, respectively (Fig. 3). Thus, the comparison indicates that both screens have a good agreement in bypassability, but also suggests that a single screen cannot identify all possible BOE.

Li et al (2019) further identified suppressor genes responsible for BOE. For example, they found that the mutation/overexpression of 6 components of the 19S proteasome compensates for mitochondrial dysfunction, suggesting a link between proteasome alteration and mtDNA dispensability. To test if our screen identified the same set of extragenic suppressor genes as Li et al. (2019), we determined the whole-genome sequences of 3 BOE strains for *mrp18* (mitochondrial ribosome protein), which was a C-BOE hit in Li et al (2019). Interestingly, we identified mutations in *atp1* (F1-F0 ATP synthase alpha subunit (Falson *et al.*, 1991)) and *rpt3* (19S proteasome base subcomplex ATPase subunit (Kitagawa *et al.*, 2014)), which are very similar to what were found in Li et al. (2019) (*atp3*; F1-F0 ATP synthase gamma subunit: *mts4*; 19S proteasome regulatory subunit (Wilkinson *et al.*, 1997)). In addition, our screen uniquely identified a mutation in *hul5* (HECT-type ubiquitin-protein ligase E3 (Fang *et al.*, 2011)); we subsequently obtained the double *mrp18/hul5* deletion strain. Hul5 might function upstream of the proteasome. *pir2* (RNA silencing factor (Sugiyama *et al.*, 2016)) was another common hit, and Li et al. (2019) reported a single gene mutation in *dis3* (exosome 3'-5' exoribonuclease subunit (Murakami *et al.*, 2007)). However, we could not find *dis3* mutations in any of the 3 BOE strains we obtained, indicating that other genes had acquired suppressive mutations.

In summary, we have established an alternative sensitive—and perhaps less labour-intensive—methodology for mutagenesis-based BOE screening in fission yeast, and expanded the list of genes whose essentiality is bypassable. Our methodology allows for a straightforward scale-up of the screen, from which we expect to reveal masked cellular mechanisms.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

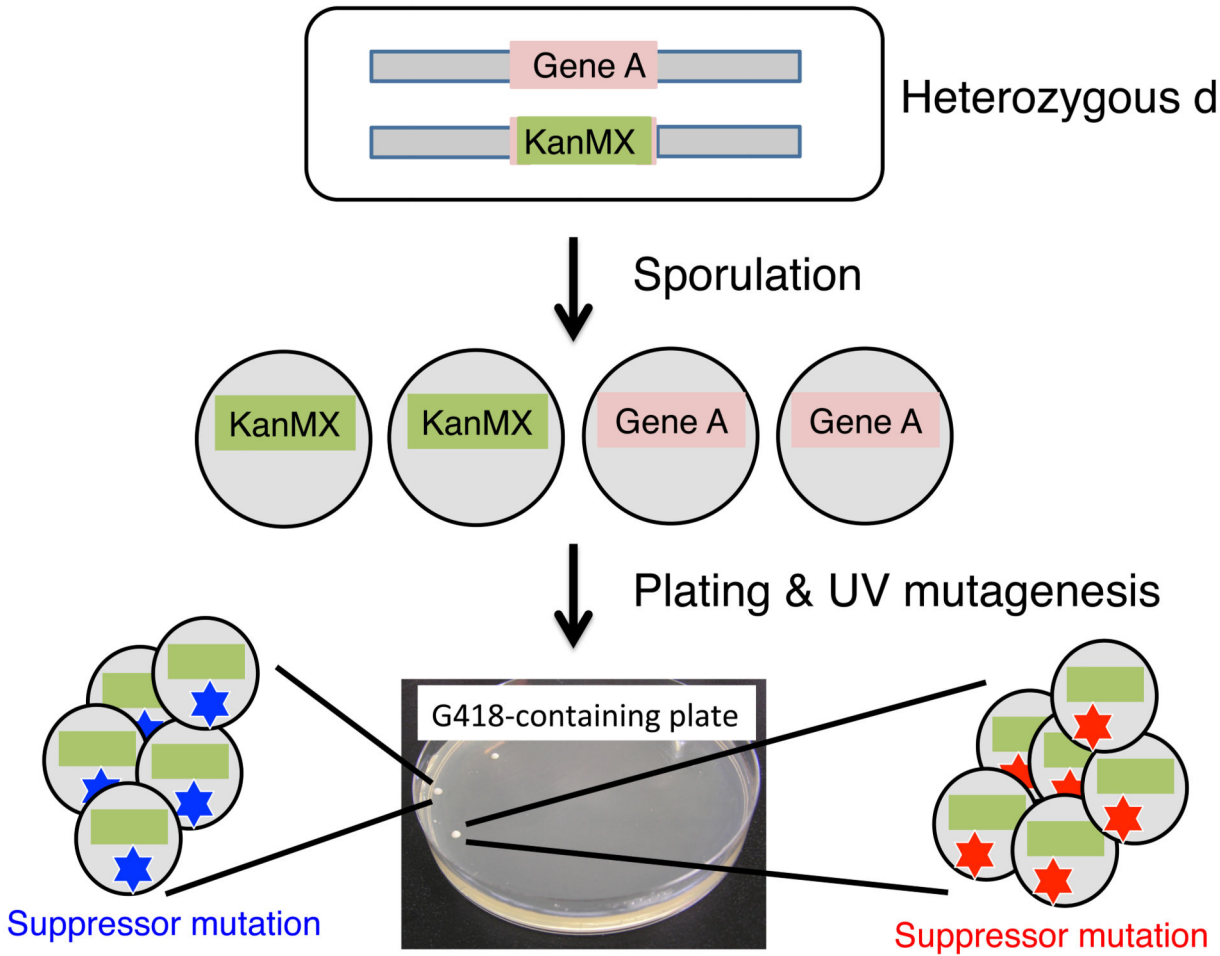
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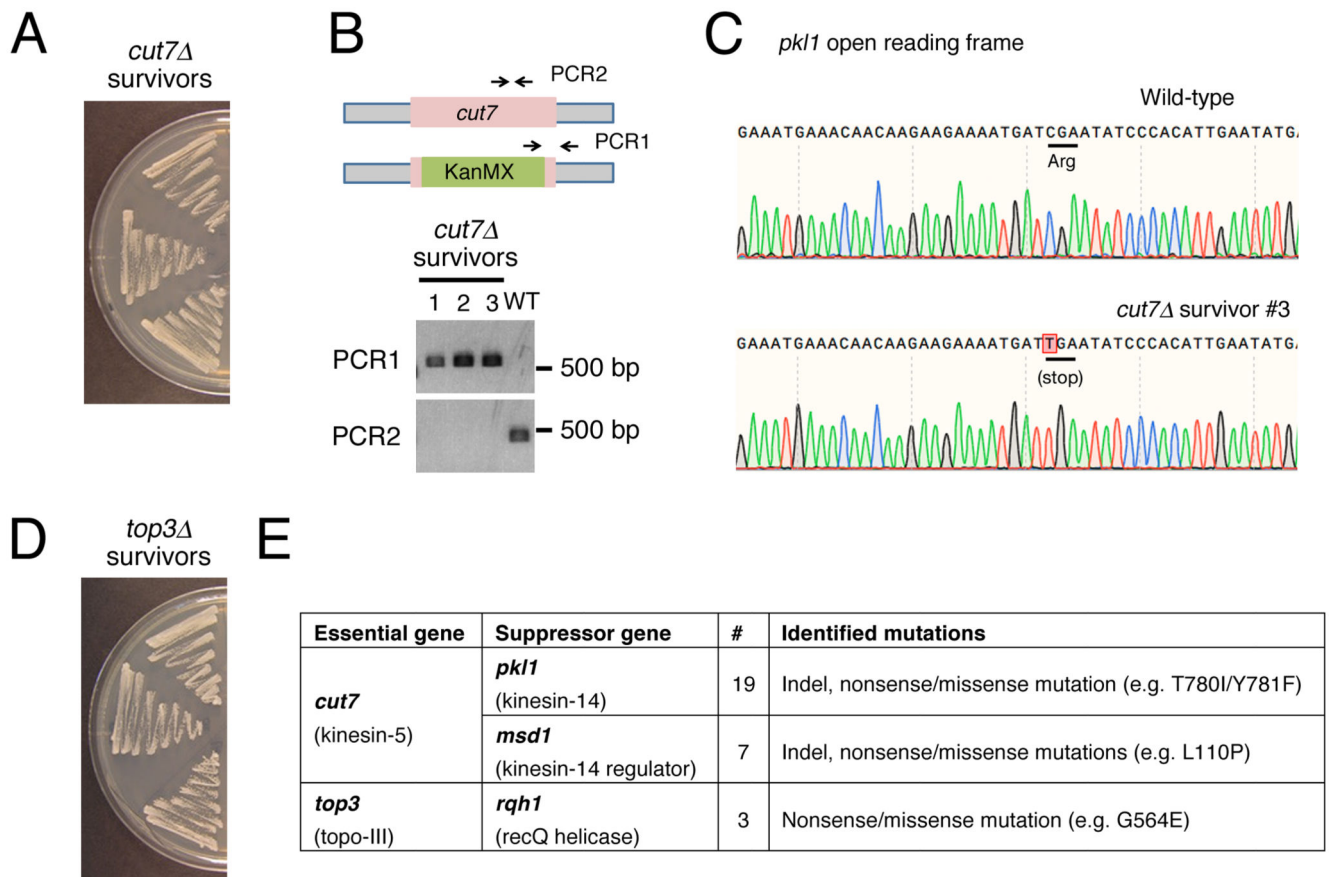
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**Figure 1. Flowchart of BOE screening using haploid spores of fission yeast**

A copy of an essential gene (named 'A' in this figure) is replaced by the G418-resistant cassette (KanMX) in the diploid strain. This heterozygous diploid is viable since another copy of gene A remains intact. The diploid is sporulated in the nitrogen-limited medium. Spores with or without gene A are obtained at 1:1 ratio. The spores are spread onto G418-containing plate and then immediately irradiated with UV for mutagenesis. Only a spore with a suppressor mutation can grow and form a colony on the medium. The lack of gene A is confirmed by colony PCR. Note that, in the actual screening, we implemented the *rpl42*-cycloheximide system to more specifically select the haploid on the G418-containing plate (see Materials and Methods for details).





**Figure 2. Successful identification of known BOE**

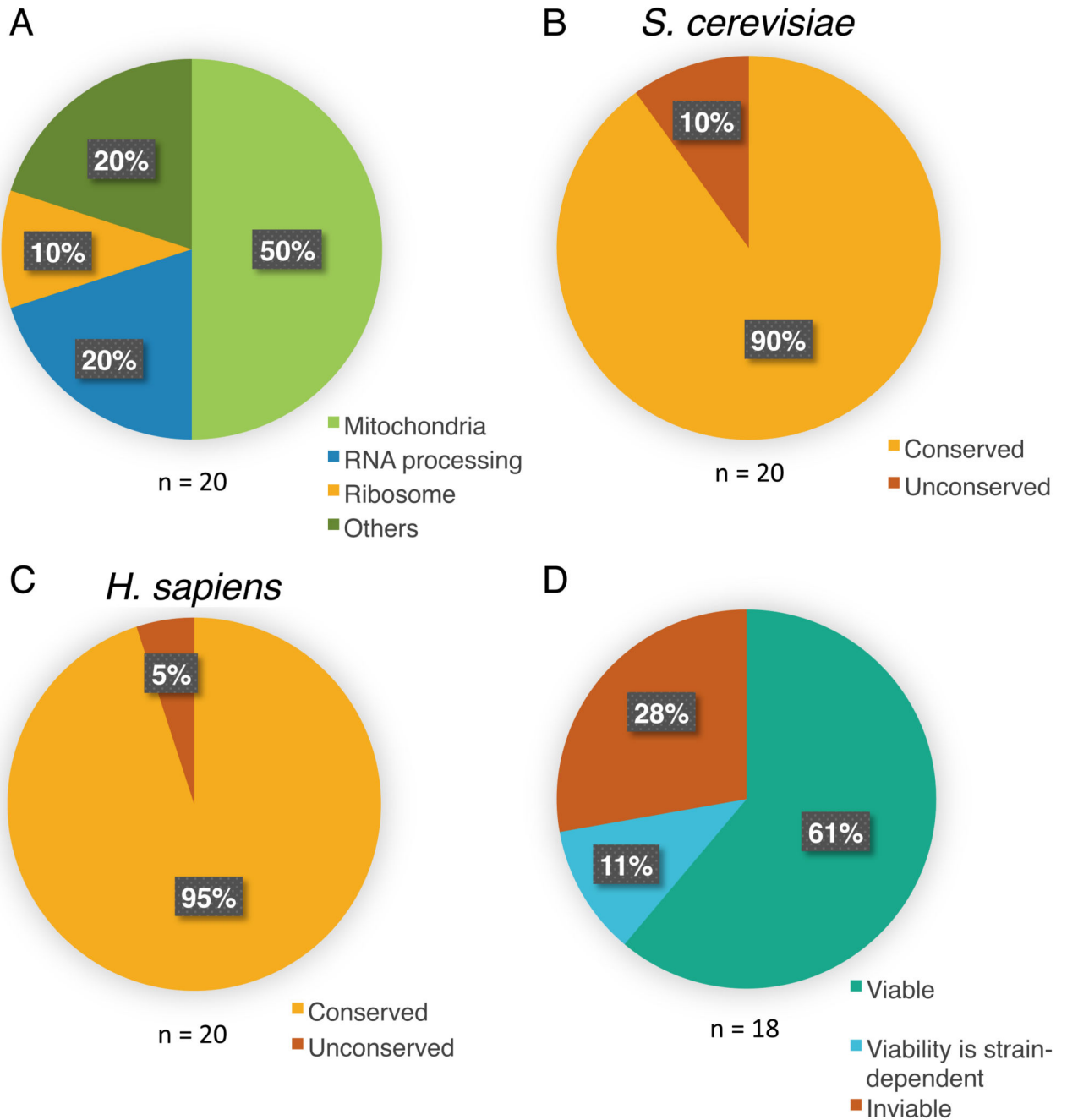
(A) Three viable strains after UV mutagenesis of *cut7* strain. In total, we obtained 38 BOE strains after UV mutagenesis of  $6 \times 10^7$  *cut7* spores. (B) PCR to verify the lack of *cut7* gene for 3 haploid strains that formed colonies. (C) Sequencing result to show the appearance of a premature stop codon in *pk11* gene in a *cut7* BOE strain. (D) 3 viable strains after UV mutagenesis of *top3* spores. (E) Summary of extragenic suppressor mutations of *cut7* and *top3*.

Gene name	Gene product	Function	Viable colony #	Identified mutation	Li et al (2019) BOE result	Li et al (2019) C-BOE mutation	<i>S. cerevisiae</i> (S288C) essentiality
mrps17	mitochondrial ribosomal protein subunit S17		7		C-BOE, OP-BOE	atp3, mts4	non-essential
mrp21	mitochondrial ribosomal protein subunit Mrp21		5		C-BOE, OP-BOE	atp3, mts4	non-essential
mrp18	mitochondrial ribosomal protein subunit L8		4	atp1, rpt3, hu15	C-BOE, OP-BOE	atp3, mts4	non-essential
cw17	Prp19 complex subunit		4		non-bypassable		non-essential
plp2	NURS complex subunit, Zf-C2H2 type zinc finger protein		3		C-BOE	dis3	(unconserved)
lip5	mitochondrial lipic acid synthetase Lip5		1		C-BOE	atd1	non-essential
slm5	mitochondrial asparagine-tRNA ligase		0		C-BOE, OP-BOE	atp3, mts4	non-essential
mug89	phospholipid-translocating ATPase complex Lem3 family subunit		0		C-BOE, OP-BOE	itr2	non-essential
bms1	GTP binding protein Bms1		0		non-bypassable		essential
idi1	isopentenyl-diphosphate delta-isomerase		0		non-bypassable		essential
rps7	19S proteasome regulatory subunit		0		non-bypassable		essential
pac1	double-strand-specific ribonuclease		0		non-bypassable		essential
cog2	Golgi transport complex subunit		0		non-bypassable		essential
erg1	squalene monooxygenase		0		non-bypassable		essential
tor2	serine/threonine protein kinase		0		non-bypassable		essential
erg9	farnesyl-diphosphate:farnesyl-diphosphate transferase		0		non-bypassable		essential
tsz13	enoyl reductase		0		non-bypassable		essential
orc5	origin recognition complex subunit		0		non-bypassable		essential
cut3	condensin complex SMC subunit Smc4		0		non-bypassable		essential
prp2	U2 small nuclear RNA auxiliary factor small subunit, U2AF-59		0		non-bypassable		non-essential
rp4	DNA-directed RNA polymerase II complex subunit		0		non-bypassable		non-essential
cdc27	DNA polymerase delta subunit		0		non-bypassable		essential
pop6	RNase P and RNase MRP subunit		0		non-bypassable		essential
mis12	Kinetochore component		0		non-bypassable		essential
psf2	GIN5 complex subunit		0		non-bypassable		essential
rm11	HNA polymerase I general transcription initiation factor subunit		0		non-bypassable		essential
sec5	SNARE		0		non-bypassable		essential
shq1	box H/ACA snoRNP assembly protein		0		non-bypassable		essential
tim50	TIM23 translocase complex subunit		0		non-bypassable		essential
nup44	nucleoporin		17		N.D.		essential
sec22	SNARE		16		N.D.		non-essential
plp2	phosducin family protein		15		N.D.		essential
ram23	mitochondrial ribosomal protein subunit S23		10		N.D.		non-essential
rrs1	ribosome biogenesis protein		8		N.D.		essential
pgs1	CDP-diaclyglycerol-glycerol-3-phosphate-3-phosphatidyltransferase		7		N.D.		essential
SPBC2F12.10	mitochondrial ribosomal protein subunit L35		5		N.D.		non-essential
rps1601	40S ribosomal protein S16		5		N.D.		non-essential
SPBC24C5.03	mitochondrial proline-tRNA ligase		5		N.D.		non-essential
esf2	U3 snoRNP-associated protein		4		N.D.		essential
lsm3	Lsm2-8 complex subunit		3		N.D.		essential
SPBC1A4.11c	Schizosaccharomyces specific protein		3		N.D.		(unconserved)
SPBC2D10.08c	mitochondrial ribosomal protein subunit Yml6		3		N.D.		essential
tu1	mitochondrial translation elongation factor EF-Tu		1		N.D.		non-essential
cc8	chaperonin-containing 1-complex theta subunit		0		N.D.		essential
oag1	mRNA guanylyltransferase		0		N.D.		essential
mrp17	mitochondrial ribosomal protein subunit L7		0		N.D.		non-essential
trs20	TRAPP complex subunit Trs20		0		N.D.		essential
trf5	translation initiation factor eIF-3g		0		N.D.		essential
SPBC18H10.17c	mitochondrial recombinase Mhr1		0		N.D.		non-essential
clt1	clathrin light chain		0		N.D.		non-essential
spo14	GDP/GTP exchange factor, WD repeat protein		0		N.D.		essential
nop8	ribosome biogenesis protein		0		N.D.		essential
apc10	anaphase-promoting complex substrate recognition subunit		0		N.D.		non-essential
lam41	mitochondrial phosphatidate cytidylyltransferase		0		N.D.		essential
sof1	U3 snoRNP-associated protein		0		N.D.		essential
cct3	chaperonin-containing 1-complex gamma subunit		0		N.D.		essential
alp5	actin-like protein Arp4		0		N.D.		essential
psf1	GIN5 complex subunit		0		N.D.		essential
qc1	mitochondrial processing peptidase (MPP) complex beta subunit Mas1		0		N.D.		essential
utp18	GCl-48 family		0		N.D.		essential
lcb1	serine palmitoyltransferase complex subunit		0		N.D.		essential
sim4	CENP-K ortholog		0		N.D.		non-essential
mas2	mitochondrial processing peptidase (MPP) complex alpha subunit		0		N.D.		essential
cut2	securin, sister chromatid separation inhibitor		0		N.D.		essential
saq1	spindle pole body SUN domain protein		0		N.D.		essential
cc11	chaperonin-containing 1-complex alpha subunit		0		N.D.		essential
ned8	ubiquitin-like protein modifier for cullin		0		N.D.		non-essential
cdc14	SIN component		0		N.D.		(unconserved)
tb1	DNA binding factor		0		N.D.		essential
rad50	DNA repair protein, SUMO-related		0		N.D.		non-essential
ucp3	GTPase activating protein		0		N.D.		non-essential
pim1	RCC1 family Ran GDP/GTP exchange factor		0		N.D.		essential
slu7	splicing factor		0		N.D.		essential
kin17	human KIN ortholog		0		N.D.		essential
psm1	mitotic cohesin complex ATPase subunit Pam1/Smc1		0		N.D.		essential
orc1	origin recognition complex subunit		0		N.D.		essential
SPBC2G5.05	transketolase		0		N.D.		non-essential
mrs1	mitochondrial and cytoplasmic arginine-tRNA ligase Rrs1/Mrs		0		N.D.		essential
mot1	TATA-binding protein-associated transcription initiation repressor		0		N.D.		essential
smf1	Sm snRNP core protein		0		N.D.		essential
hsp78	mitochondrial heatshock protein		0		N.D.		non-essential
spc24	Kinetochore component		0		N.D.		essential
pic1	INCENP ortholog		0		N.D.		non-essential
gtb1	gamma-tubulin		0		N.D.		essential
gcv3	glycine decarboxylase complex subunit H		0		N.D.		non-essential
pop4	RNase P and RNase MRP subunit		0		N.D.		essential
rio2	protein kinase, RIO family		0		N.D.		essential
yp11	GTPase		0		N.D.		essential
orc6	origin recognition complex subunit		0		N.D.		essential
gp2	UDP-N-acetylglucosamine-dolchyl-3-phosphate N-acetylglucosaminophosphotransferase		0		N.D.		essential
taf73	transcription factor TFIID complex subunit Taf5-like		0		N.D.		essential
brf1	transcription factor TFIIB complex subunit		0		N.D.		essential

■ Mitochondria  
■ RNA processing  
■ Ribosome  
■ Others  
 C-BOE: Chemical mutagenesis  
 T-BOE: Transposon-based mutagenesis  
 OP-BOE: Gene library expression

### Figure 3. Summary of BOE screening

The gene product was listed based on the information found in PomBase. Information on other BOE screens (Li et al. 2019) and *S. cerevisiae* orthologues (*Saccharomyces Genome Database (SGD)*) are also listed. Bold letters indicate bypassable essential genes identified in our study and/or Li et al. (2019).



**Figure 4. Features of bypassable essential genes**

(A) Classification of the function of 20 bypassable essential genes identified in this study.

(B, C) Conservation of the identified genes in *S. cerevisiae* (B) or *H. sapiens* (C).

(D) Essentiality of the *S. cerevisiae* orthologues (based on *Saccharomyces Genome Database (SGD)*).