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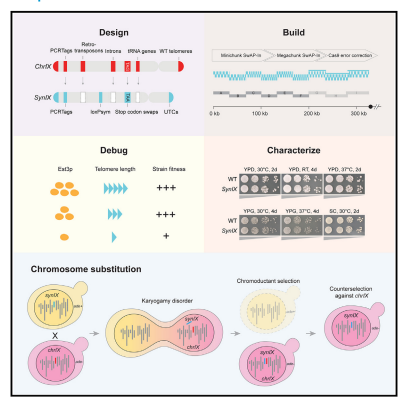
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Consequences of a telomerase-related fitness defect and chromosome substitution technology in yeast *synIX* strains

Graphical abstract



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In brief

McCulloch et al. describe complete synthesis of *synIX* and dissect a bug resulting from tRNA deletion upstream of telomerase-related *EST3* at the protein, RNA, and telomere length levels. A new genetic method, chromosome substitution, is used to swap *synIX* into alternative strains and can be broadly applied.

Highlights

- Following debugging, synlX shows high fitness under a variety of conditions
- An unusual bug resulting from a tRNA-associated deletion affects EST3 expression
- Chromosome substitution method moves chromosomes to "fresh" genetic background







Article

Consequences of a telomerase-related fitness defect and chromosome substitution technology in yeast *synlX* strains

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SUMMARY

We describe the complete synthesis, assembly, debugging, and characterization of a synthetic 404,963 bp chromosome, *synIX* (synthetic chromosome *IX*). Combined chromosome construction methods were used to synthesize and integrate its left arm (*synIXL*) into a strain containing previously described *synIXR*. We identified and resolved a bug affecting expression of *EST3*, a crucial gene for telomerase function, producing a *synIX* strain with near wild-type fitness. To facilitate future synthetic chromosome consolidation and increase flexibility of chromosome transfer between distinct strains, we combined chromoduction, a method to transfer a whole chromosome between two strains, with conditional centromere destabilization to substitute a chromosome of interest for its native counterpart. Both steps of this chromosome substitution method were efficient. We observed that wild-type *II* tended to co-transfer with *synIX* and was co-destabilized with wild-type *IX*, suggesting a potential gene dosage compensation relationship between these chromosomes.

INTRODUCTION

The Synthetic Yeast Genome Project (Sc2.0) marks a key milestone in the development of "designer" eukaryotic genomes. This global effort seeks to produce a modified version of the \sim 12 Mb Saccharomyces cerevisiae genome from the bottom

up that retains native strain fitness while eliminating repetitive elements to improve genome stability and adding custom features to endow new genome functionalities. These changes include deletion of tRNA genes and introduction of loxPsym sites (symmetric, nondirectional *loxP* sites that can recombine with each other in two distinct orientations) downstream of nonessential



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genes, enabling genome rearrangement via SCRaMbLE (synthetic chromosome rearrangement and modification by loxPmediated evolution) and subsequent study of how different DNA sequence changes affect organismal fitness and function.^{1–4} More broadly, by establishing a set of common design principles for all yeast chromosomes, Sc2.0 aims to provide a blueprint for future eukaryotic genome engineering endeavors. Each chromosome was redesigned and built from synthetic DNA blocks. Here, we report the assembly and characterization of synthetic chromosome IX (synIX), coinciding with completion of the assembly phase of Sc2.0 as a whole. 5-18

Our group initially assembled a circular version of the right arm of the chromosome, synIXR, as a proof of principle for feasibility of the Sc2.0 project⁶ and subsequently incorporated that chromosome arm into an otherwise wild-type and linear version of chromosome IX.2 Here, we describe construction and characterization of synIXL and, consequently, the final version of synIX. As with many other completed Sc2.0 chromosomes and in accordance with Sc2.0 project objectives, synlX-containing strains, following a "debugging" process to resolve deficiencies in fitness across multiple growth conditions, behave similarly to the parental yeast strain with respect to fitness and transcriptional profiling.

Fitness defects have emerged during completion of many of the Sc2.0 chromosome strains. We attribute these growth issues, or "bugs," to a still-incomplete understanding of the S. cerevisiae genome during the initial design phase. Identifying and resolving such bugs may improve future genome engineering efforts and help anticipate and avoid potential complications. Bugs in previous Sc2.0 strains have commonly arisen from design choices (sequence additions, deletions, or modifications) that unexpectedly affected protein or RNA levels. For example, PCRTag recoding (intended to alter DNA sequence without changing amino acid sequence, producing watermarks to distinguish synthetic and wild-type sequences) lowered protein expression or interfered with transcription factor binding sites in other synthetic chromosomes. 7,9,11,12 Addition of loxPsym sites and removal of introns have affected RNA expression of genes in Sc2.0 strains, altering strain fitness. 7,11 For synIX, we identify and characterize a fitness defect related to a sequence adjacent to EST3, encoding a crucial yeast telomerase holoenzyme component involved in yeast telomere replication. 19-21 This bug resulted from a specific intentional design step, removal of a tRNA gene and associated repetitive DNAs. Deletion of a tRNA gene, Ty1 retroelement, and the

DNA between them led to reduced average telomere length and impaired strain growth at higher temperature (37°C), a likely consequence of significant Est3p reduction. Reintroduction of part or all of this sequence dramatically improved fitness. Our experiments dissecting this bug show that Est3p expression levels were markedly reduced following deletion of an upstream tRNA gene and associated sequences per the chromosome's design; reinsertion of the tRNA gene and associated sequences in various configurations led to partial or total restoration of both Est3p levels and telomere length. We suggest that a minimum threshold Est3p level suffices to restore telomere length, supporting near-normal fitness at 37°C.

The ultimate goal of Sc2.0 is to create a high-fitness yeast strain with a fully synthetic genome. A previously described chromosome consolidation method relying on chromosome endoreduplication followed by sporulation becomes laborious and time consuming as synthetic chromosome number increases.^{2,22} Here, we present a strategy combining chromoduction of synIX and subsequent loss of native chromosome IX (chrIX) to accelerate consolidation. This approach takes advantage of a karyogamy defect arising when either parent in a cross carries a kar1-1 mutation.²³ Such crosses yield heterokaryons consisting of nuclear material from one yeast strain with mixed cytoplasm of the parent strains. In rare cases, whole chromosomes transfer between strains in a kar1-1 x KAR1 cross.²⁴ While chromoduction occurs more frequently with smaller chromosomes,²⁴ prior work has employed a selection strategy to identify chromoductants in which a particular chromosome has been transferred.^{25,26} Conversely, whole-chromosome destabilization can be achieved by activating a GAL promoter immediately upstream of yeast centromeres. 7,27 Here, we demonstrated the feasibility of synthetic chromosome transfer using synIX, producing a final haploid yeast strain in which synlX has replaced wild-type chrlX. This represents a promising proof of principle for targeted transfer of chromosomes among yeast strains; indeed, this method is used in final consolidation of Sc2.0 chromosomes into a single yeast strain.²²

RESULTS

Design and assembly of synlX

SynIX adheres to Sc2.0 project principles.2 The designed synIX sequence contains a variety of modifications, including deletion of ten tRNA genes (relocated to a separate tRNA

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neochromosome) and 11,632 bp repetitive DNA; recoding of 54 stop codons from TAG to TAA, 436 bp restriction enzyme sites, and 7,943 bp PCRTags; and addition of 142 loxPsym sites downstream of nonessential genes allowing for the Cre-mediated SCRaMbLE system (Figure 1A). 1-4 These changes reduced the chrIX length from 439,885 to 404,963 bp in synIX.

Following synthesis of synIXR, we began construction of syn-IXL using a strain containing a linear, partially synthetic chromosome synIXR with a native left arm (yLM461). Assembly of syn-IXL proceeded in several stages (Figure 1B). Initially, we assembled 750 bp building blocks from synthetic 60-70 bp oligonucleotides through polymerase cycling assembly (PCA) in the context of the Build-A-Genome undergraduate class (Figure S1A).^{5,6} We then combined these building blocks to form 2-4 kb minichunks. In later project stages, as DNA synthesis technology improved and costs dropped, we obtained the remaining minichunks directly from a vendor.

We divided synIXL into nine 30-60 kb megachunks, each comprising minichunks that overlapped adjacent ones. To assemble the first six megachunks (A-F), we transformed component minichunks into our entry strain one megachunk at a time, adding an auxotrophic marker to each distal minichunk (Figure 1C). Through yeast homologous recombination, each assembly step overwrote the native segment and adjacent auxotrophic marker with synthetic DNA and a new auxotrophic marker in accord with the switching auxotrophies progressively for integration (SwAP-In) method, 2,6 enabling selection of strains containing newly integrated DNA (Figure 1C; Table S1). After each integration round, the PCRTag watermarking system allowed identification of colonies with synthetic DNA and lacking wild-type DNA (Table S2).

The minichunk integration approach used in assembling megachunks A-F left some unwanted patches of wild-type chrlX sequence (Figure 1B, top). To improve efficiency and avoid such wild-type DNA patches, later synIXL sections used the "megachunk-BAC" integration strategy. 12 For these sections, we first assembled minichunks into 40-60 kb megachunks (megachunks G-I) as extrachromosomal bacterial artificial chromosomes (BACs) in yeast, with assembly success rates of 20%-45% (Figure 1D; Table S3), and subsequently transformed them into E. coli for plasmid extraction. Following megachunk-BAC sequence verification, these megachunks were released from their plasmid backbones using restriction enzyme digestion and delivered directly into the semisynthetic (A-F) synIX strain for SwAP-In. We verified successful integration of megachunks G-I via whole-genome sequencing; all three were fully integrated into the synIX strain and lacked novel mutations and wild-type sequences (Figure 1B, middle).

To convert wild-type DNA patches seen in megachunks A-F to synthetic, we used CRISPR-Cas9 editing to selectively target synIX at residual chrIX PCRTags within each patch (Figure S1B) and repaired them with appropriate synthetic donor DNAs. 9,10 Following replacement of ten minichunks over seven rounds of CRISPR-mediated editing, we obtained a version of synIX containing all synthetic segments as designed (Figure 1B, bottom; Table S4).

We continued CRISPR-mediated editing of this "draft" strain to correct point mutations, small deletions, and duplications in synIX, detailed in Table S5. Additionally, we discovered a discrepancy stemming from inaccuracies in the yeast reference genome sequence originally used in synIX design; the sequence for (nonessential) FAA3 contained an extra 7 bp absent from the updated reference sequence, resulting in a frameshift (Figure 1E).28,29 A previous genetic screen linked faa3 △ to decreased growth³⁰; consequently, we restored proper FAA3 function in our synIX strain by reestablishing the updated FAA3 reading frame (Figure S2).

During this process, we unexpectedly identified whole-chromosome disomy of synlX (Figure 1F, top). To restore the normal karyotype, we integrated a URA3-pGAL cassette upstream of CEN9 into one synIX copy²⁷ and induced chromosome destabilization (Figure S3). Whole-genome sequencing confirmed that the resulting synIX strain contained approximately equal read depth for each chromosome, as expected for a monosomic haploid yeast strain (Figure 1F, bottom). There were no overt phenotypic differences between the disomic and monosomic strains, suggesting that the disomy was an inadvertent consequence of CRISPR-mediated editing rather than selection for a second copy of synlX due to haploinsufficiency. This monosomic strain was used for further debugging.

Debugging of synlX

After correcting the above synIX sequence issues, we found that this strain, coined yeast_chr09_9_1 (also referred to as yLHM1192), grew less well than the parental strain at 37°C on both YPD and YPG media (Figure 2A). A similar phenotype was observed in additional yLHM1192-equivalent strain isolates (Figure S4). Following a pooled sequencing bug mapping method,9 we backcrossed the synIX strain to a wild-type strain to generate a heterozygous diploid yeast strain for sporulation and subsequently dissected tetrads. Spores contained a randomized mixture of recombinant synIX and chrIX DNA resulting from meiotic recombination (Figure S5A). By conducting wholegenome sequencing of healthy and sick spores, we identified a region spanning \sim 15–20 kb upstream of CEN9 to 5 kb downstream of CEN9 associated with the fitness defect (Figures 2B and S5B). Consistent with this mapping, the synIX strain initially exhibited a marked decline in fitness following integration of megachunk I, containing the corresponding synthetic sequence (Figure S6). To determine the gene responsible, we cloned each gene in this region into a CEN plasmid for complementation testing (Figure S7). Strains containing a plasmid with EST3 showed a pronounced fitness improvement on both YPD and YPG media at 37°C compared with the buggy synlX strain (Figures 2B and S7).

To investigate which modifications of the synthetic EST3 region might reduce fitness, we replaced sections of the synthetic region with corresponding wild-type sequences in our synlX strain and monitored growth. Replacing EST3 coding sequences did not restore fitness (Figure 2C, yLHM1429); however, additionally restoring a deleted segment upstream of EST3 containing a tRNAAsp gene and a nearby Ty1 long terminal repeat (LTR) element dramatically improved synIX strain fitness (Figure 2C, yLHM1430 and yLHM1431). To better assess the contributions of these EST3-adjacent elements to fitness, we introduced smaller modifications to EST3's upstream region in the synIX strain, including different combinations of the tRNAAsp



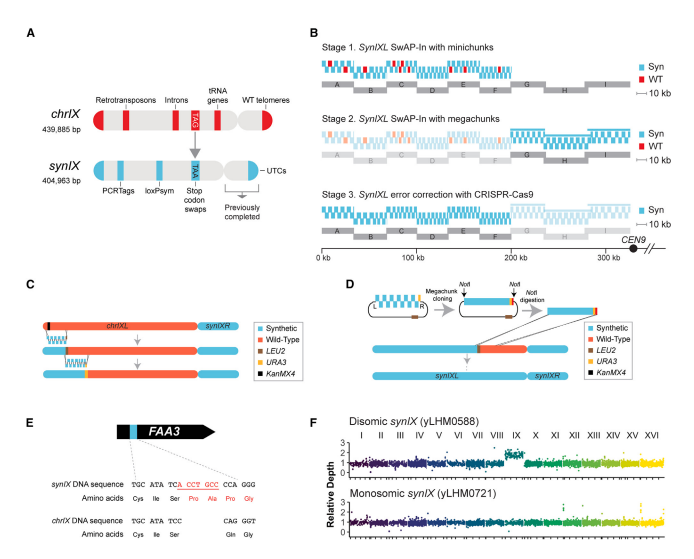


Figure 1. Design and assembly of synIX

(A) Diagram of the differences between wild-type (WT) chrlX and synlX. Designer features specific to synlX include addition of PCRTags and loxPsym sites, recoding of TAG stop codons to TAA, replacement of WT telomeres with universal telomere caps (UTCs), and removal of tRNA genes (relocated to a tRNA neochromosome), introns, retrotransposons, and subtelomeric repeats.

(B) Schematic of the synIXL construction process, reflecting synthetic and (unwanted) WT composition of chrIX in the synIX strain after each stage of assembly. Stage 1: SwAP-In with minichunks (A-F). Stage 2: SwAP-In with preassembled megachunks (G-I, opaque with blue lines). Stage 3: error correction with CRISPR-Cas9 (primarily in highlighted megachunks A-F). Blue: syn minichunks, red: WT minichunks, gray: megachunks. Red patches in stages 1 and 2 reflect segments of DNA that were not replaced by their expected synthetic minichunk counterparts during SwAP-In with minichunks in stage 1 (A-F); appropriate synthetic sequences were integrated at these sites during stage 3. kb, kilobase pairs.

(C) Minichunk integration strategy for assembling megachunks A-F of synlX in yeast (used for stage 1 in B). KanMX marker (black) was integrated into left end of WT chrlX (red). Individual minichunks (blue, synthetic) comprising one megachunk were then co-transformed into in-progress synlX strain and used to overwrite WT chrlX sequence (red) via homologous recombination. Alternating auxotrophic markers (LEU2, brown, and URA3, orange) were used and overwritten at

(D) Megachunk plasmid assembly and integration approach for building and integrating megachunks G-I in synIX (used for stage 2 in B). Minichunks (blue) were assembled in yeast, and an auxotrophic marker (here, URA3, orange) was added to the end of each assembly, followed by a segment of chrIX homology. Megachunk assembly overwrote prior round auxotrophic marker (brown, LEU2) and WT DNA (red) by homologous recombination.

(E) Schematic depicting frameshift mutation in FAA3 gene in synIX design. Synthetic sequence included 7 bp not present in S288C WT yeast reference genome. Added bases (red) cause a frameshift, with resulting synIX amino acid sequence (red) varying from the expected chrIX amino acid sequence (black, bottom). (F) Coverage plots showing read depth along each yeast chromosome in disomic synlX strain yLHM0588 (top) and monosomic synlX strain yLHM0721 (bottom). The x axis: yeast chromosome (not to scale). The y axis: relative depth based on number of reads at each position divided by average read depth across sixteen

See also Figures S1-S3 and Tables S2, S3-S5, and S8.

yeast chromosomes.

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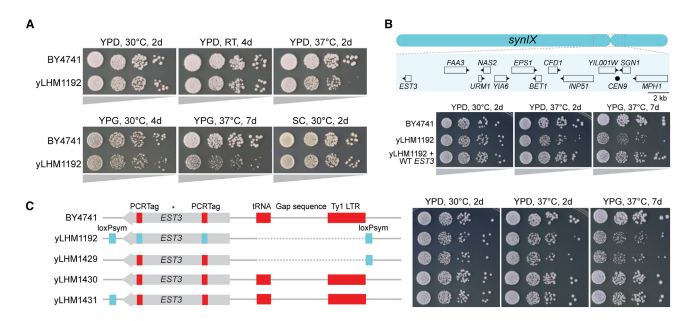


Figure 2. Identifying and mapping an EST3-related bug

(A) Spot assays comparing the growth of BY4741 (WT) to yLHM1192 (synthetic) across a variety of growth conditions. Each column represents a 10-fold dilution. RT, room temperature, ~22°C.

(B) (Top) Schematic of region near CEN9 identified as containing gene responsible for synlX fitness defect. Unhealthy synlX strain (yLHM1192) was transformed with plasmids containing transcription units for each gene shown. (Bottom) Spot assays comparing growth of BY4741 and yLHM1192 to yLHM1192 transformed with plasmid containing EST3 coding sequence plus 500 bp upstream sequence and 200 bp downstream sequence. Each column represents a 10-fold dilution. (C) Spot assays comparing growth of strains with different combinations of synthetic and WT features in and upstream of EST3. Each column represents a 10-fold dilution. Schematics on the left illustrate WT (red) and synthetic (blue) features present in each strain. Asterisk (*) marks site of known EST3 programmed +1 ribosomal frameshift.

See also Figures S4-S7.

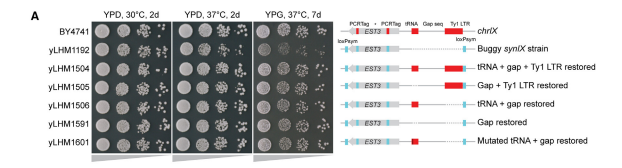
gene, a nearby Ty1 LTR, and an intervening single-copy "gap" DNA sequence with no known function (Figure 3A; Table S6). All of these edits improved strain fitness (Figure 3A). Similar phenotypes were observed in additional strains generated for each modification and did not noticeably change following extended strain passaging (Figure S8).

As EST3 is involved in telomere maintenance, 19-21 we hypothesized that EST3-associated changes in fitness might correlate with changes in average telomere length. We assessed telomere length of synIX and variant strains via southern blotting using a probe specific for telomeric repeats (Figure 3B). Following enzymatic digestion of yeast gDNA with XhoI, a smear of small telomeric fragments from chromosome ends containing Y' elements is seen in each lane; these fragments were used to estimate average telomere length, with other telomeric fragments appearing at larger sizes. As expected, yLHM1192's telomeres were markedly shorter than those seen in the wild-type BY4741 strain. Interestingly, the telomeres in the modified synIX strains split into two length groups; three groups of strains, yLHM1504 (containing the entire tRNA, intervening "gap" sequence between the tRNA and Ty1 with no known function, and Ty1 sequence), yLHM1506 (containing the tRNA and gap sequences but not the Ty1 sequence), and yLHM1601 (containing a tRNA with a transcription-inactivating point mutation^{31–34} and the aforementioned gap sequence) had telomeres similar to or slightly longer than wild-type yeast, while two other groups of strains, yLHM1505 (containing just the gap sequence and Ty1 LTR) and yLHM1591 (containing just the gap), had telomeres shorter than BY4741 but slightly longer than yLHM1192. Thus, our experiments support the hypothesis posed but with a slight revision: we suggest that extension of telomeres beyond a crucial minimum threshold length suffices to restore fitness.

Next, we assessed the role of upstream EST3 modifications on strain fitness and telomere length independent of other potential synIX-associated strain changes. We introduced each synthetic modification upstream of EST3 (from yLHM1192, yLHM1504-1506, yLHM1591, and yLHM1601) into wild-type yeast (Figure 4A). As in synIX strains, yLHM1192-equivalent wild-type edited strains showed decreased fitness, but other modified strains showed normal fitness (Figure S9). Fitness did not change appreciably following extended strain passaging (Figure S9). As in the synthetic strains, we saw three groups of telomere lengths: wild type (BY4741, as well as the strains with the yLHM1504-, yLHM1506-, and yLHM1601-equivalent modifications), shortened (strains with the yLHM1192-equivalent modifications), and intermediate (strains with the yLHM1505- and yLHM1591-equivalent modifications) (Figure 4B).

We next sought to understand why all modifications improved fitness but only some fully restored telomere length to wild-type levels. We hypothesized that differences in EST3 translation and/ or transcription might result in variable telomere length across strains. To examine Est3p expression in modified synlX strains,





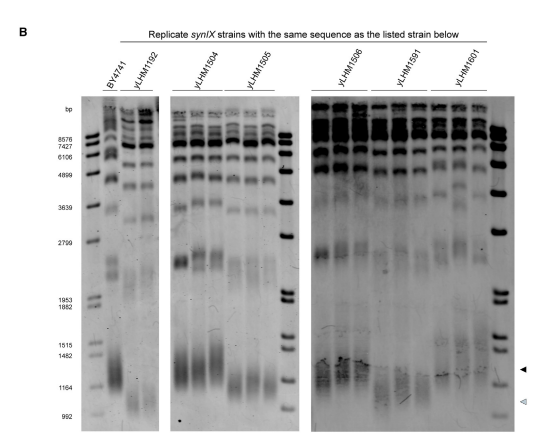


Figure 3. Modifying synlX strains upstream of EST3 alters fitness and telomere length

(A) Spot assays and accompanying schematics of the region upstream of EST3 for strains BY4741 (WT chrIX), yLHM1192 (synthetic), and variant strains derived from yLHM1192. WT PCRTags, tRNA^{Asp} gene, and Ty1 LTR, red; synthetic PCRTags and loxPsym sites, blue; mutation in tRNA^{Asp} gene, vertical black line. Each spot assay column represents a 10-fold dilution.

(B) Southern blot of *Xho*l-digested yeast gDNA derived from BY4741 (WT) and replicate strains with modifications equivalent to yLHM1192 (synthetic) and variant strains (yLHM1504, yLHM1505, yLHM1506, yLHM1591, and yLHM1601, as depicted in A). DNA was probed with a digoxigenin-labeled fragment specific for telomeric repeats. Black triangle (right): average size of Y'-containing telomeric fragments for BY4741. Gray triangle (right): average size of Y'-containing telomeric fragments for yLHM1192. Left and middle: left and right halves of one southern blot, cropped to remove one lane in the middle. Right: right side of second southern blot, cropped to remove left-hand ladder, BY4741, and yLHM1192 lanes.

See also Figure S8 and Table S6.

we tagged the 3' end of *EST3* with 3× FLAG and evaluated protein levels at 37°C (Figure 5A; Table S7). FLAG-tagged synthetic strains were crossed to an untagged wild-type strain of opposite mating type (BY4742) to complement any potential recessive *synIX*-related effects on strain growth while allowing monitoring

of synlX-derived Est3p levels. Consistent with telomere length data, yLHM1504-, yLHM1506-, and yLHM1601-derived strains expressed higher levels of FLAG-tagged Est3p than yLHM1192-, yLHM1505-, and yLHM1591-derived strains (Figure 5A). Follow-up comparisons of yLHM1192-, yLHM1505-, and

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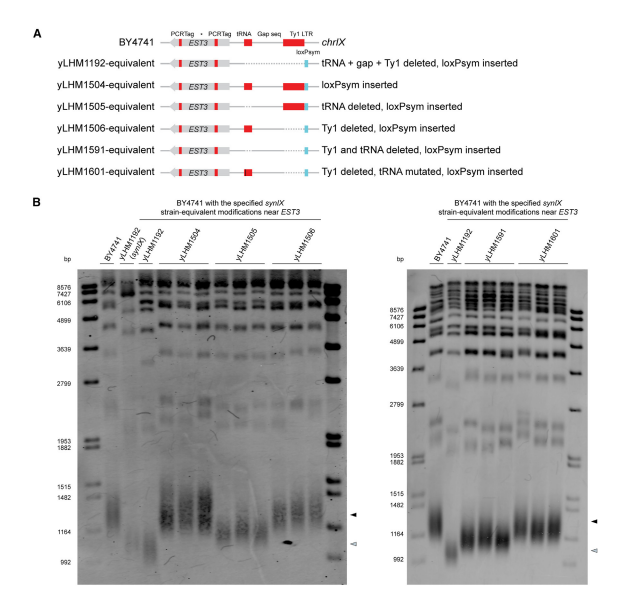


Figure 4. Modifying WT yeast strains upstream of EST3 alters telomere length

(A) Schematics of region upstream of EST3 for strains BY4741 (WT chrlX), yLHM1192-equivalent (synthetic modification matching yLHM1192 in otherwise WT yeast strain), and variant strains derived from the yLHM1192-equivalent strain. WT PCRTags, tRNAAsp gene, and Ty1 LTR, red; loxPsym sites, blue; mutation in tRNAAsp gene, vertical black line.

(B) Southern blot of Xhol-digested yeast gDNA derived from BY4741 (WT), yLHM1192 (synthetic), and replicate strains with synthetic-equivalent modifications made upstream of EST3, as depicted in (A). DNA was probed with a digoxigenin-labeled fragment specific for telomeric repeats. Black triangle (right): average size of Y'-containing telomeric fragments for BY4741. Gray triangle (right): average size of Y'-containing telomeric fragments for BY4741 + yLHM1192-equivalent modification strain.

See also Figure S9 and Table S6.

yLHM1591-derived strains revealed higher Est3p levels in yLHM1505- and yLHM1591-derived strains than in yLHM1192derived strains, although all three strains had markedly lower Est3p levels than did FLAG-tagged wild-type yeast (Figure 5B). We saw neither longer nor shorter isoforms of FLAG-tagged Est3p associated with low-abundance strains (Figures 5B and \$10). Similar results were observed in FLAG-tagged heterozygous diploid strains generated by crossing FLAG-tagged versions of the wild-type EST3 variant strains from Figures 4A and S9 to untagged BY4742 and monitoring Est3p levels (Figures S11A and S11B). Collectively, the protein-level data support the hypothesis that Est3p levels were low and telomeres short in synIX strains, and modifications that increase telomere length appeared to do so by improving expression of Est3p. Moreover, a moderate level of Est3p gave rise to "medium"-length telomeres (longer than those in the synIX strain but shorter than those in wild-type strains), whereas higher (near-normal) levels of Est3p gave rise to longer telomeres. The tRNA^{Asp} gene affected Est3p levels the most.



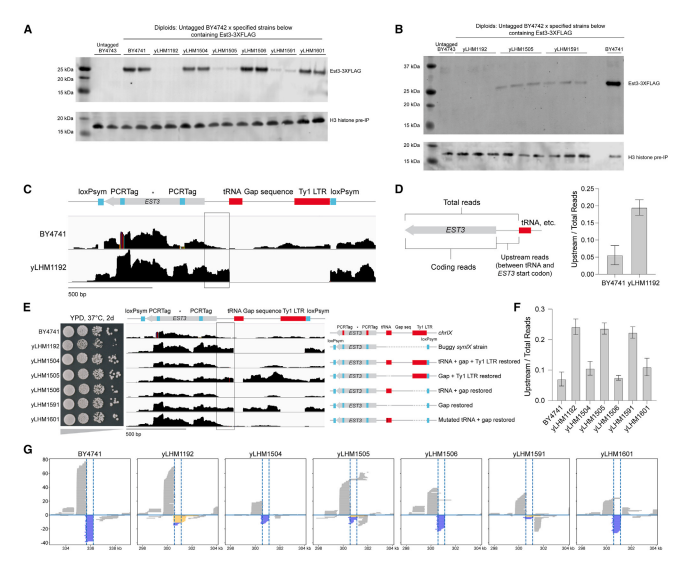


Figure 5. Dissecting effects of EST3-adjacent features on Est3 protein and EST3 RNA expression

(A) Immunoblotting immunoprecipitated (IP/IB) Est3p (top) and H3 histone pre-IP loading control (bottom). Est3p was C-terminally tagged with 3× FLAG in each synthetic strain, and diploids were generated via mating with WT BY4742. Two independent colonies were grown in YPD at 37°C for each strain. Right-most lane (second ladder) was cropped out of image.

- (B) IP/IB Est3p (top) and H3 histone pre-IP loading control (bottom) for low-expressing Est3p strains plus controls, as in (A), except that three independent colonies were grown for each yLHM1192-, yLHM1505-, and yLHM1591-derived diploid strain.
- (C) Strand-specific RNA sequencing alignments for the reverse strand of BY4741 and yLHM1192 (strand from which EST3 transcription is expected, i.e., "reverse-strand reads") grown at 37°C in YPD to a composite reference containing both synthetic and native IX features. Schematic illustrates WT chrIX (red) and synthetic (blue) features. Black box on RNA sequencing alignment plot highlights region between expected EST3 start codon and upstream tRNA. Asterisk (*) marks site of known EST3 programmed +1 ribosomal frameshift.
- (D) Quantification of reverse-strand reads mapping directly upstream of EST3 (between tRNA and start codon, corresponding to black box in C) as a fraction of total reverse-strand EST3 coding and upstream reads for each strain shown in (C). Read counts were based on alignment to native (BY4741) or synthetic (yLHM1192) reference sequences. Error bars represent SD of three biological replicates.
- (E) Spot assays and strand-specific RNA sequencing alignments for reverse-strand reads aligning to EST3 region for samples grown at 37°C in YPD. WT PCRTags, tRNAAsp gene, and Ty1 LTR, red; synthetic PCRTags and loxPsym sites, blue; mutation in tRNAsp gene, vertical black line. Asterisk (*) marks site of known EST3 programmed +1 ribosomal frameshift.
- (F) Quantification of reverse-strand reads mapping directly upstream of EST3 (between tRNA and start codon, corresponding to black box in E) as fraction of total reverse-strand EST3 coding and upstream reads for each strain shown in (E). Read counts were based on alignment to native (BY4741) or synthetic (yLHM1192) reference sequences. Error bars represent SD of three biological replicates.
- (G) Nanopore direct RNA sequencing reads aligned to EST3 region (±3 kb). The x axis: chromosome coordinate. The y axis: number of reads. Forward-strand reads are above the y axis, and reverse-strand reads are below the y axis. Dotted lines indicate boundaries of EST3 coding region. Yellow reads:

(legend continued on next page)

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To further elucidate what contributes to Est3p level variations in engineered synIX strains, we examined RNA sequencing data. Strand-specific RNA sequencing data revealed differences in the pattern of EST3 transcripts in wild-type and synlX strains at 30°C and 37°C (Figures 5C, 5D, S12A, and S12B). In particular, while both strains showed sequencing read depth throughout the EST3 coding region, synIX strains contained transcripts upstream of the expected EST3 transcription start site (TSS) that were absent from BY4741. Importantly, since these transcripts were transcribed from the same strand as native EST3, some EST3 transcripts might have had an extended 5' UTR, predicted to encode upstream AUG codons that would not produce functional Est3p. In our modified synIX strains, we saw two classes of upstream transcript profiles (Figures 5E, 5F, S13A, and S13B). yLHM1504, yLHM1506, and yLHM1601, the three strains with wild-type-length or slightly longer telomeres, showed a relative reduction in RNA sequencing reads directly upstream of EST3 compared with strand-specific total reads aligned to the upstream plus coding sequence of EST3 (Figures 5E, 5F, S13A, and S13B). By contrast, yLHM1505 and yLHM1591, the two strains with intermediate-length telomeres, still showed substantial transcript read depth upstream of EST3. The wild-type strains with modifications upstream of EST3 yielded similar transcriptional patterns (Figures S14A and S14B).

Nanopore direct RNA sequencing, which evaluates full-length transcript patterns, revealed a distinct population of EST3-spanning transcripts in yLHM1192 containing at least one out-offrame AUG upstream of the expected EST3 TSS (Figure 5G). Many of these transcripts started 250-500 bp upstream of the AUG that normally initiates Est3p translation (Figure S15). We hypothesize that such "nontranslatable" upstream transcripts fail to yield properly translated Est3p, contributing to reduced telomerase function and decreased strain fitness (Figure S16). By contrast, whereas yLHM1505 and yLHM1591 contained a few nontranslatable transcripts spanning EST3, most transcripts upstream of EST3 observed in these strains terminated prior to the expected EST3 TSS (gray in Figure 5G). yLHM1192, yLHM1505, and yLHM1591 all had relatively fewer full-length, presumably translatable EST3 transcripts than did the tRNA-containing strains (yLHM1504, yLHM1506, and yLHM1601).

These results collectively point to a model in which upstream initiated transcripts normally "silenced" by the tRNAAsp are unable to produce Est3p and only reads initiating at or near the native TSS produce functional EST3 transcripts and resulting protein. The original synIX strain (yLHM1192) appears to yield almost no functional Est3p, while variant strains produce either low (yLHM1505 and yLHM1591) or near-wild-type (yLHM1504, yLHM1506, and yLHM1601) Est3p levels. However, strains with even low levels of functional Est3p and moderate reduction in telomere length appear to surpass a threshold for maintaining normal strain fitness. Thus, three classes of Est3p abundance/

telomere length yield only two fitness profiles: healthy and sick

Characterization of the healthy synlX strain

We ultimately proceeded with the strain in which a mutated version of the deleted tRNA gene and the adjacent gap sequence were reintroduced upstream of EST3 (yLHM1601 in Figure 5E). The tRNA sequence contained a point mutation previously shown to block tRNA transcription, 31-34 compatible with functional tRNA relocation to a tRNA neochromosome¹⁵ while maintaining Est3p levels similar to those of wild-type yeast (Figure 5A). This strain is most consistent with Sc2.0's design principles of maximizing strain fitness while eliminating repetitive elements and tRNA genes. This final synIX version, coined yeast_chr09_9_2, grew similarly to wild-type yeast across a variety of media types and temperature conditions, as did additional sequence-matched replicate strains (Figures 6A and S17-S19). The final strain largely matched the expected synthetic reference sequence, with slight deviations from the original design reflecting changes made during bug fixing and other minor alterations during the assembly process (Table S8). The final strain also showed even coverage across synlX and normal chromosomal copy number (Figure 6B). SynIX migrated slightly faster on pulsed-field gel electrophoresis (PFGE) than did native IX (Figure 6C). We observed an unusual banding pattern among a couple of the higher bands in yLHM1601, possibly related to an expanded repetitive sequence near the telomere of chrXVI. This difference did not appear to affect strain fitness and disappeared following synlX transfer to an alternative matched-background wild-type yeast strain via the chromosome substitution method discussed in the next section (Figure 6C, lanes 3 and 4).

Transcriptional analysis of the final synlX strain (yLHM1601, or yeast_chr09_9_2) identified a small number of differentially expressed genes compared to the wild-type strain (Figures 6D, 6E, \$20A, and \$20B). Several hits found on synIX were located in or near telomeres, with some of these genes upregulated and others downregulated. On the left arm of synIX, SOA1 (YIL166C) showed decreased expression in the synIX strain at 37°C (~5× lower than in BY4741). Several subtelomeric genes found between SOA1 and TEL09L in native chrIX were deleted during synIX design. We hypothesize that SOA1 expression decreased due to increased silencing associated with its new position near the left end UTC (universal telomere cap), as has been observed on other synthetic chromosomes where genes moved closer to UTCs due to subtelomeric DNA deletion. 12 Conversely, right-arm subtelomeric genes YIR042C and YIR043C showed expression increases at 37°C (~12x increase for YIR042C) and both 30°C and 37°C (\sim 15× and \sim 34× increases for YIR043C), respectively. Unlike SOA1, YIR042C and YIR043C were already located very close to TEL09R in native chrlX. Consequently, the UTCs introduced to synlX per Sc2.0 design principles may have failed to silence nearby genes to

[&]quot;nontranslatable" reads containing at least one AUG upstream of expected EST3 start codon plus a minimum of 20 bp 5' UTR sequence. Blue reads: reads mapping to EST3 that start no more than 20 bp upstream of first AUG upstream of expected EST3 start codon, including "translatable/functional" reads spanning the entirety of EST3 and reads starting downstream of EST3-initiating AUG. See also Figures S10-S16 and Tables S6-S7.



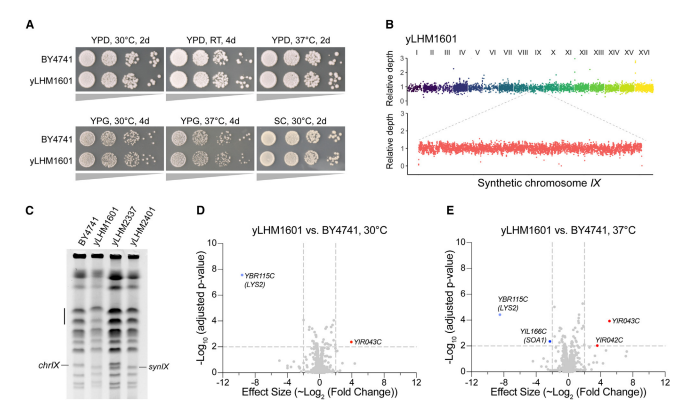


Figure 6. SynIX characterization

(A) Spot assays comparing growth of BY4741 (WT) to yLHM1601 (synthetic) across several conditions. Each column represents a 10-fold dilution. RT, room temperature, ~22°C.

(B) DNA sequencing coverage plot for *synIX* strain (yLHM1601). The x axis: yeast chromosome (not to scale) or coordinate along *synIX*. Relative depth (y axis) based on reads mapped to yeast_chr09_3_55 reference sequence divided by average read depth across sixteen yeast chromosomes (top) or *synIX* (bottom). (C) Pulsed-field gel showing chromosomes from BY4741 (*chrIX*), yLHM1601 (*synIX*), yLHM2337 (BY4742-background strain with copies of *chrIX* and *synIX*), and yLHM2401 (yLHM2337 after loss of *chrIX*) strains. Vertical black line indicates location of potential discrepancy in band intensity for larger chromosomes between BY4741 and yLHM1601; *synIX* was moved to an alternative strain (yLHM2337) with larger bands resembling BY4741 in intensity, and *chrIX* was subsequently lost from this strain (producing yLHM2401) via strategy described in next section.

(D and E) Volcano plots of differentially expressed genes obtained from RNA sequencing data for syn/X strain (yLHM1601) vs. BY4741 measured at 30°C (D) and 37°C (E) in YPD. Upregulated genes in yLHM1601 depicted in red (chr/X) and downregulated genes in medium blue (chr/X) or light blue (other chromosomes). Transcript counts based on alignment to S288C reference transcriptome. The auxotrophic gene LYS2 is present in BY4741 but not in yLHM1601. Fold change cutoff is 4, and adjusted p value cutoff is 0.01. Three biological replicates were used for each strain. For corresponding unfiltered plots of RNA sequencing data depicting all measured genes, see Figures S20A and S20B.

See also Figures S17-S19 and Table S8.

the same degree as the native telomeres. Of note, the earliest version of the UTC, used at the right telomere, lacked the X sequence included at the left telomere and elsewhere in the Sc2.0 project. Since the X sequence is present in native *TEL09R*, its absence might help explain the selectively elevated expression of these *TEL09R*-adjacent genes.

Chromosome substitution of synIX

Here, we sought to combine chromoduction with subsequent loss of native *chrlX* from a recipient strain to seamlessly move *synlX* between yeast strains while maintaining normal karyotype in a strain of interest (Figure 7A). As a proof of concept, a disomic *synlX* strain was used as donor, as an extra copy of *synlX* was deemed likely to produce higher chromosome transfer efficiency.²⁵ A recipient *kar1-1* mutant strain blocked karyogamy. The recipient strain also contained two mutations, *can1* and

cyh2, recessive selectable markers ensuring robust exclusion of donor cells and potential diploid cells on drug-containing medium. To facilitate selection for recipient strains containing synlX transferred from the donor, we deleted the LYS12 gene in the recipient strain (Figure S21A). This allowed selection for lysine prototrophy to ensure synlX's presence in the chromoductants. Additionally, to enable eventual loss of chrlX from the recipient, we subsequently knocked in a URA3-pGAL cassette upstream of CEN9 in the recipient for chrlX destabilization and counterselection (Figure S21B).²⁷

After modifying the recipient, we mated donor and recipient strains (yLHM0387 and yWZ601, respectively) and selected recipients that had acquired at least one copy of *synIX* via chromoduction on SC–Lys–Arg+Can+Cyh plates. A PCRTagging assay was performed using selected PCRTags across *chrIX* and *synIX*. Four chromoductants (yWZ610–613) containing both *chrIX* and

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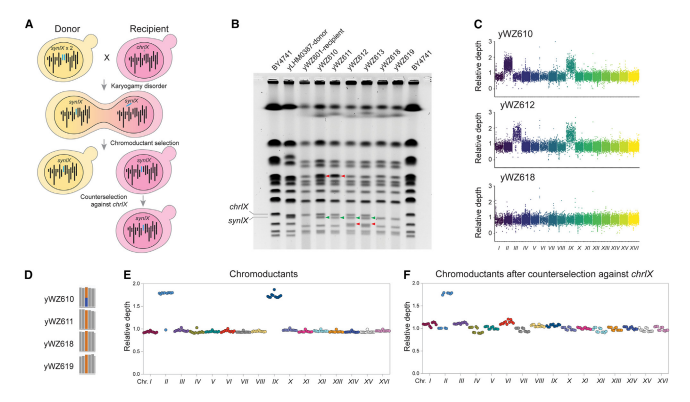


Figure 7. Chromosome substitution of synIX

(A) Chromosome substitution occurs in two steps, chromoduction, in which synlX is transferred to recipient strain, followed by counterselection against WT chrlX to remove native version of chromosome and restore normal chromosome copy number in chromoductant strain. Schematic diagram of substitution steps. Black bars, WT chromosomes. Blue bars, synIX. Recipient strain harbors an ade2 mutation and thus forms pink colonies for easily distinguishing chromoductants. (B) PFGE of donor strain, recipient strain, disomic chromoductants, and chromoductants after counterselection against chrIX. BY4741 serves as control. chrIX and synIX chromosomes indicated with black arrows. Green triangles indicate synIX in disomic chromoductants. Red triangles indicate extra chrII or chrIII in disomic chromoductants.

(C) Whole-genome sequencing read depth plots of disomic chromoductants (yWZ610, yWZ612) and chrlX counterselected (i.e., fully substituted) chromoductants (yWZ618). The x axis: sixteen yeast chromosomes, not drawn to scale. The y axis: relative read depth, calculated as number of reads at each position divided by average read depth across sixteen yeast chromosomes.

(D) chrll SNPs in chromoductant strains before and after counterselection against chrlX. Recipient strain SNP: orange, donor strain SNP: blue. Strains: yWZ610: chromoductant strain with one copy of chrll from donor strain and one from recipient strain (mixed SNP population); yWZ611: chromoductant strain with two copies of chrll from recipient strain (endoreduplicated chrll); yWZ618: strain derived from yWZ610 after counterselection against chrlX, with one copy of chrll harboring recipient strain SNP; yWZ619: strain derived from yWZ611 after counterselection against chrlX, with one copy of chrlI harboring recipient strain SNP. (E) Relative depth plot for chromosome substitution strains prior to counterselection against chrIX. The x axis: sixteen yeast chromosomes, not drawn to scale. The v axis: relative read depth.

(F) Relative depth plot for chromosome substitution strains after counterselection against chrlX. The x axis: sixteen yeast chromosomes, not drawn to scale. The y axis: relative read depth.

See also Figure S21 and Table S9.

synIX PCRTag amplicons were isolated from 14 candidates (Figure S21C). Disomic chromoductants stably maintained both chrlX and synlX upon restreaking onto SC-Lys plates. Genotyping validation of the four disomic chromoductants suggested that yWZ612 and yWZ613 had additionally acquired the HIS4 gene on chrlll from the donor (the recipient was his440), suggesting that chrlll was co-transferred with synlX (Figure S21D).

To destabilize chrIX in the disomic chromoductants to generate haploids with a single copy of synlX, disomic chromoductants (yWZ610, yWZ611) were grown in galactose-containing medium to activate GAL transcription, destabilizing CEN9 and consequently leading to chrlX loss.²⁷ PCRTagging analysis of the chromoductants (yWZ618, yWZ619) showed that only synIX was present (Figure S21E). PFGE analysis showed that (1) synIX was slightly smaller than chrIX in both BY4741 and the recipient strain; (2) both synlX and chrlX were present in disomic chromoductants; (3) chrlll was co-transferred with synIX in yWZ612 and yWZ613; and (4) chrIX was lost in chromoductants following counterselection against chrlX (yWZ618, yWZ619) (Figure 7B). However, two chromoductants, yWZ610 and yWZ611, surprisingly contained an extra copy of chrll in addition to both chrlX and synlX (Figure 7B), confirmed by whole-genome sequencing (Figure 7C). Further examination of SNPs differing between donor and recipient *chrll* revealed that yWZ610 contained one donor chrll and one recipient chrll, while yWZ611 contained two copies of recipient chrll (Figure 7D). This finding suggested that chrll endoreduplicated in yWZ611, perhaps to balance a gene dosage perturbation caused by synlX



chromoduction. Consistent with this hypothesis, one copy of chrll was lost from yWZ610 and yWZ611 during inactivation of chrlX (Figure 7B, yWZ618 and yWZ619).

To investigate whether co-transfer and simultaneous loss of chrlX and chrll was coincidental, we isolated 11 additional disomic chromoductants. Whole-genome sequencing revealed that 10 of the 11 disomic chromoductants gained an extra copy of chrll (Figure 7E). Following counterselection against chrlX, four disomic chromoductants lost chrlI, while six chromoductants remained disomic (Figure 7F). To rule out that this was specific to our disomic synlX donor strain, we performed chromosome substitution using an alternative disomic synlX donor strain and three monosomic synIX donor strains. As in earlier experiments, chrll frequently co-transferred with synlX, and the extra copy of chrll often disappeared after culturing strains in galactose (Table S9). In conclusion, we successfully used chromosome substitution to transfer synlX from different donor strains to a kar1-1 recipient strain via chromoduction and to subsequently destabilize chrlX. More in-depth diagnosis of chromoductants revealed that chrll was preferentially and consistently co-transferred with synIX and then frequently lost during counterselection against chrlX.

DISCUSSION

We successfully conclude the synthesis, debugging, and characterization of synIX. synIX's complex developmental history reflects advances in DNA synthesis technology over more than a decade. When the project first began, each DNA segment was painstakingly assembled by undergraduates from oligonucleotides into minichunks, which in turn were integrated iteratively into the yeast genome to overwrite native chrlX. While such methods enabled successful incorporation of much of the synthetic sequence of interest, drawbacks included inadvertent retention of wild-type patches and introduction of new point mutations. Newer approaches, including direct minichunk synthesis and megachunk cloning and sequencing prior to integration, greatly improved assembly efficiency and reduced construction error rates. Ultimately, we produced a yeast strain with a fully synthetic version of chrlX displaying near-wild-type fitness.

While debugging synIX, we identified a fitness defect related to deletion of a DNA segment upstream of the EST3 gene containing a tRNA Asp gene, a Ty1 LTR, and an intervening single-copy DNA sequence. Strains with this deletion showed a reduction in telomere length as well as reduced Est3p levels and many RNA species mapping upstream of EST3's normal TSS. Reinserting all or part of the deleted sequence sufficed to restore normal fitness. However, only a subset of the edited strains, specifically those containing the tRNA gene, showed reduced transcriptional activity upstream of EST3 and restoration of normal or near-normal Est3p levels and normal telomere length. We hypothesize that the tRNA sequence normally provides a buffer against production of aberrant transcripts upstream of EST3, and thus the transcripts that are produced in tRNA-containing strains more effectively give rise to functional, full-length Est3p. Indeed, tRNAs suppress transcription from nearby RNA polymerase II promoters in a process called tRNA-mediated

gene silencing.34-36 Consistent with this idea, nanopore RNA sequencing data suggests that a subset of transcripts include one or more AUGs upstream of EST3 and thus presumably fail to undergo proper translation; strains with an abundance of such transcripts also appear to have fewer full-length, translatable transcripts and lower Est3p levels. Other changes made upstream of EST3, such as reinsertion of the gap sequence between the tRNA and the Ty1 LTR or Ty1 LTR itself, seemingly trigger small improvements in Est3p levels vs. those found in strains with the original synIX design. These strains have intermediate-length telomeres, longer than those seen in the original synIX design but shorter than those seen in native yeast, and maintain stable fitness after repeated passaging. We speculate that the changes introduced to these strains may be sufficient to maintain Est3p production and consequent telomere length above a threshold level needed for normal strain fitness, despite not fully returning either to native levels.

Interestingly, a similar occurrence of abnormal transcriptional patterns resulting from tRNA deletion and loxPsym site insertion was previously observed at the HIS2 locus in synVI. Here, the authors suggested that a "cryptic start site" upstream of HIS2, generated by the altered sequence in this region, might yield a subset of transcripts incapable of producing functional His2 protein or that this transcription might impair normal HIS2 promoter activity. It would be useful to examine whether common principles underlie multiple Sc2.0 bugs following tRNA deletion.

Finally, we demonstrated a chromosome substitution technique that combines selective chromoduction, or movement of a desired chromosome between yeast strains, with subsequent destabilization and loss of the corresponding native chromosome. During this process of chromosome substitution, we found that certain chromosomes appeared to co-transfer with synIX at higher frequencies than did others. Specifically, we saw multiple instances of transfer of chrll or chrlll alongside synIX. These extra chromosomes often disappeared following destabilization of chrlX. Prior work suggested that smaller chromosomes are more likely to transfer during chromoduction than larger ones and that co-transfer of multiple chromosomes commonly occurs.²⁴ However, as chrll is not small, it seems unlikely that chromosome size fully explains specific chrll cotransfer.

It is likely that an extra copy of chrll compensates for a gene dosage imbalance arising from the transient extra copy of chrlX introduced by chromoduction. In one chromoductant strain, analysis of SNPs from the recipient and donor strains' copies of chrll revealed that both copies of the chromosome came from the recipient strain. The presence of an endoreduplicated chrll is consistent with the hypothesis that there is selective pressure for a higher dosage of chrll fitness in the presence of two copies of chrlX and that co-migration of chrll and synlX in the other chromoductant strains may not have occurred by chance. Moreover, multiple donor strains, including strains disomic and monosomic for synIX, showed co-transfer of chrII alongside synlX. It may be informative to ascertain mechanisms that dictate which chromosomes tend to show such cochromoduction.

Ultimately, chromosome substitution has numerous applications, including transfer and consolidation of chromosomes

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from other wild-type yeast strains, engineered strains, and non-laboratory strains. Existing work profiling collections of hundreds or thousands of yeast strains has shed light on yeast diversity and evolution ^{37–39}; chromosome substitution will facilitate further probing of how changes on individual chromosomes or combinations of chromosomes affect strain fitness and function. Similarly, this technology could assist in analyzing why genomic changes in SCRaMbLEd yeast strains and other engineered strains produce particular phenotypic results. ^{1–4} More immediately, chromosome substitution is already in use to bring the initial phase of Sc2.0 to a close via consolidation of all of the synthetic chromosomes into a single strain, marking a key eukaryotic genome engineering milestone. ²²

Limitations of the study

SynIX is but one of sixteen synthetic yeast chromosomes assembled as part of Sc2.0. Consequently, the efforts made here are limited compared to what can be done following chromosome consolidation of most or all synthetic chromosomes into a single yeast strain. The chromosome substitution method described here will facilitate generation of such consolidated chromosome strains for further study. Additionally, our analysis of the EST3 bug could be made even more comprehensive by making additional variants of the region upstream of that gene and testing those modifications in additional backgrounds.

CONSORTIA

This work is part of the international Synthetic Yeast Genome (Sc2.0) consortium. The chromosome design and building consortium includes research groups worldwide: Boeke Lab at Johns Hopkins University and New York University (led chromosomes I, III, IV, VI, VIII, IX); Chandrasegaran lab at Johns Hopkins (led chromosomes III and IX); Cai Lab at University of Edinburgh and University of Manchester (led chromosomes II and VII and tRNA neochromosome); Yue Shen's team at BGI-Research SHENZHEN (led chromosomes II, VII, and XIII); Y.J. Yuan's team at Tianjin University (led chromosomes V and X); Dai Lab at Tsinghua University and Shenzhen Institute of Advanced Technology, CAS (led chromosome XII); Ellis Lab at Imperial College London (led chromosome XI); Sakkie Pretorius's team at Macquarie University (led chromosomes XIV and XVI); Matthew Wook Chang's team at National University of Singapore (led chromosome XV); Bader and Boeke Labs at Johns Hopkins University (led design and workflow); and Build-A-Genome undergraduate teams at Johns Hopkins University and Loyola University Maryland (contributed to chromosomes I, III, IV, VIII, and IX). The Sc2.0 consortium includes numerous other participants who are acknowledged on the project website, www. syntheticyeast.org.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.xgen.2023.100419.

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AUTHOR CONTRIBUTIONS

J.D.B., S.C., and W.Z. conceptualized the study. W.Z., S.C., L.H.M., and J.D.B. designed the experiments. J.D.B. and J.S.B. designed synlX. The Build-A-Genome class members constructed building blocks. V.S., N.A., and S.R. constructed minichunks and incorporated them for megachunks A-F. L.H.M. designed, constructed, and delivered megachunks G-I and performed error correction, bug mapping, bug correction, and functional studies of synlX. L.H.M. and W.Z. developed and evaluated the chromosome substitution method. L.H.M., A.L.H., J.C., V.F., E.L., and G.S. performed sequencing and data analyses. L.H.M., W.Z., and J.D.B. wrote the manuscript. L.A.M., H.J., J.L., M.S.T., and W.R.B. provided advice. S.C., J.D.B., J.S.B., and L.M.S. provided lab supervision. All authors reviewed and edited the manuscript.



DECLARATION OF INTERESTS

J.D.B. is a founder and director of CDI Labs, Inc., a founder of and consultant to Neochromosome, Inc, a founder, SAB member of, and consultant to ReOpen Diagnostics, LLC, and serves or served on the scientific advisory board of the following: Logomix, Inc., Sangamo, Inc., Modern Meadow, Inc., Rome Therapeutics, Inc., Sample6, Inc., Tessera Therapeutics, Inc., and the Wyss Institute. J.S.B. is a founder of Neochromosome, Inc., consultant to Opentrons Labworks, Inc., and advisor to Reflexion Pharmaceuticals, Inc.

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Digoxigenin Recombinant Rabbit	Invitrogen	Thermo Fisher Scientific Cat#
Monoclonal Antibody (9H27L19)		700772; RRID:AB_2532342
Monoclonal ANTI-FLAG® M2 antibody produced in mouse	Sigma-Aldrich	Sigma-Aldrich Cat# F1804; RRID:AB_262044
Rabbit anti-Histone H3 antibody	Abcam	Abcam Cat# ab1791; RRID:AB_302613
RDye 800CW Goat anti-Mouse gG1-Specific	LI-COR Biosciences	LI-COR Biosciences Cat# 926–32350; RRID:AB_2782997
RDye 680RD Goat anti-Rabbit IgG	LI-COR Biosciences	LI-COR Biosciences Cat# 926–68071; RRID:AB_10956166
Bacterial and virus strains		
PI300 <i>E. coli</i> strain	Lucigen	EC300150
OP10 <i>E coli</i> strain	Invitrogen	C4040
Chemicals, peptides, and recombinant protein	ns	
Zymolyase 20T	US Biological	Z1000
ymolyase 100T	US Biological	Z1004
DTA-free protease inhibitor	Roche	11873580001
ithium acetate dihydrate	Sigma-Aldrich	L6883
olyethylene glycol	Sigma-Aldrich	81188
erring sperm DNA	Promega	D1816
otassium acetate	Fisher	BP364
inc acetate dihydrate	Sigma-Aldrich	Z0625
S)-(+)-Camptothecin	Sigma-Aldrich	C9911
-Sorbitol	Sigma-Aldrich	S1876
-Azauracil	Sigma-Aldrich	A1757
lydroxyurea	Sigma-Aldrich	H8627
Methyl methanesulfonate	Sigma-Aldrich	129925
flethyl 1-(butylcarbamoyl)-2- enzimidazolecarbamate (Benomyl)	Sigma-Aldrich	381586
cycloheximide	Sigma-Aldrich	01810
-Canavanine sulfate	Sigma-Aldrich	C9758
lydrogen peroxide	Millipore	88597
JLTRAhyb TM Ultrasensitive lybridization Buffer	Invitrogen	AM8669
SPE Buffer 20X Concentrate	Sigma-Aldrich	S2015
ntercept (TBS) Blocking Buffer	LI-COR Biosciences	927–60001
WEEN 20	Sigma-Aldrich	P1379
riton TM X-100	Sigma-Aldrich	X100
-Mercaptoethanol	Sigma-Aldrich	M6250
oTaq Green Master Mix	Promega	M7123
kb Plus DNA Ladder	NEB	N3200L
NA Molecular Weight Marker VII, NG-labeled	Sigma-Aldrich	11669940910
Precision Plus Protein TM All Blue Prestained Protein Standards	Bio-Rad	1610373
RNase A	Thermo Fisher Scientific	EN0531
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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Notl-HF	NEB	R1389L
BbsI-HF	NEB	R3539L
Afel	NEB	R0652L
Xhol	NEB	R0146L
Amino-terminal FLAG-BAP [™]	Sigma-Aldrich	P7582
Fusion Protein	•	
Dynabeads [™] M-270 Epoxy	Invitrogen	14302D
Dynabeads TM Oligo(dT) ₂₅	Invitrogen	61005
SuperScript TM IV Reverse Transcriptase	Invitrogen	18090010
Agencourt RNAClean XP beads	Beckman Coulter	A63987
Critical commercial assays		
PureLink TM HiPure Plasmid Midiprep Kit	Invitrogen	K2100-15
Zyppy Plasmid Miniprep Kit	Zymo Research	D4037
Fungi/Yeast Genomic DNA Isolation Kit	Norgen	27300
Qubit dsDNA HS Assay Kit	Thermo Fisher Scientific	Q32854
Qubit dsDNA BR Assay Kit	Thermo Fisher Scientific	Q32850
NEBNext Ultra II FS DNA Library Kit	NEB	E7805
Zymo-Spin I Columns	Zymo Research	C1003
PALL 60207 Biodyne B High Sensitivity and Low Background Nylon Transfer Membrane, 0.45 μm Pore Size, 30 cm W x 3 m L Roll	Pall Corporation	60207
RNeasy Mini Kit	QIAGEN	74106
Qubit RNA BR Assay Kit	Thermo Fisher Scientific	Q10210
QIAseq Stranded Total RNA Library Kit	QIAGEN	180745
QIAseq FastSelect -rRNA Yeast Kit	QIAGEN	334217
NextSeq 500/550 High Output Kit v2.5 (75 Cycles)	Illumina	20024906
NextSeq 500/550 High Output Kit v2.5 (150 Cycles)	Illumina	20024907
MasterPure Yeast RNA Purification Kit	Lucigen	MPY03100
Agilent RNA 6000 Nano Kit	Agilent	5067–1511
Qubit RNA HS Assay Kit	Thermo Fisher Scientific	Q32852
Direct RNA Sequencing Kit	Oxford Nanopore Technologies	SQK-RNA002
Flow Cell Priming Kit	Oxford Nanopore Technologies	EXP-FLP001
MinION Flow Cell (R9.4.1)	Oxford Nanopore Technologies	FLO-MIN106D
Deposited data	· · · · · · · · · · · · · · · · · · ·	
SynIX DNA and RNA sequences	This study	Sc2.0 umbrella: BioProject PRJNA351844 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA351844); synlX: BioProject PRJNA900304 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA900304)
Experimental models: Organisms/strains		
Saccharomyces cerevisiae: BY4741	Boeke laboratory	BY4741
All other strains used in this study are listed in Table S1	Boeke laboratory	Various, detailed in Table S1
Oligonucleotides		
PCRTag primers are listed in Table S2	Boeke laboratory	Various, detailed in Table S2

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Continued				
REAGENT or RESOURCE	SOURCE	IDENTIFIER		
Recombinant DNA				
See Tables S3–S7 for plasmids	Boeke laboratory	Various, detailed in Tables S3-S7		
used in this study				
Software and algorithms				
GeneDesign	Boeke and Bader laboratories	Richardson et al. ⁴⁰		
R	R Core Team	https://www.R-project.org/		
Rstudio v1.3.1093	Rstudio	https://www.rstudio.com		
Integrated Genomics Viewer (IGV) v2.7.2	Broad Institute ⁴¹	http://software.broadinstitute.org/software/igv/		
Image Studio Lite	LI-COR Biosciences	https://www.licor.com/bio/image-studio/		
GraphPad Prism v9.4.0 for macOS	GraphPad Software	http://www.graphpad.com		
Trimmomatic v0.39	Bolger et al. ⁴²	http://www.usadellab.org/cms/		
		index.php?page=trimmomatic		
FastQC v0.11.4	Andrews ⁴³	http://www.bioinformatics.babraham.		
		ac.uk/projects/fastqc/		
Bowtie 2 v2.2.9	Langmead and Salzberg ⁴⁴	http://bowtie-bio.sourceforge.net/		
	A45	bowtie2/index.shtml		
ggplot2 v3.3.5	Wickham ⁴⁵	https://ggplot2.tidyverse.org/		
Seqtk v1.3	N/A	https://github.com/lh3/seqtk/		
Kallisto v0.46.0	Bray et al. ⁴⁶	https://pachterlab.github.io/kallisto/		
STAR v2.5.2a	Dobin et al. ⁴⁷	https://github.com/alexdobin/STAR/		
SAMtools v1.9	Li et al. ⁴⁸	http://samtools.sourceforge.net/		
BEDtools v2.26.0	Quinlan et al. ⁴⁹	https://github.com/arq5x/bedtools2/		
sleuth v0.30.0	Pimentel et al. ⁵⁰	https://pachterlab.github.io/sleuth/		
Other				
Resource website for Sc2.0	Bader laboratory	https://syntheticyeast.github.io/		

RESOURCE AVAILABILITY

Lead contact

Further information should be directed to and will be fulfilled by the lead contact, Jef D. Boeke (jef.boeke@nyulangone.org).

Materials availability

All unique/stable reagents generated in this study are available from the lead contact with a completed Materials Transfer Agreement.

Data and code availability

- Data: All data are available under the overarching Sc2.0 umbrella BioProject (PRJNA351844). The data for synIX are provided under BioProject PRJNA900304. The specific data reported here were deposited to Gene Expression Omnibus accession number GSE244153 for the RNA-seq data and Genbank Accession number CP125382 for the synIX sequence.
- Code: This work did not generate any code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT PARTICIPANT DETAILS

Yeast strains used in this work are listed in Table S1.

METHOD DETAILS

SynIX design

Design sequences for synIX were developed using BioStudio, according to the same guidelines used for designing the other synthetic chromosomes for Sc2.0. Designer sequences yeast_chr09_3_54 and yeast_chr09_3_55 were used as references for subsequent steps in the project. Version yeast_chr09_3_56 was developed after completion of yeast_chr09_9_02, and was used as a reference for final strain verification.

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Yeast media

Yeast strains were cultured in rich medium (YPD) or synthetic complete (SC) medium with appropriate components "dropped out" e.g., SC–Ura lacks uracil. Yeast transformations were performed using lithium acetate protocols. YPG plates used for growth assays contained 3% glycerol in place of dextrose as a carbon source. For expanded spot assays, the following media types were used: pH 4 and pH 8: pH of 2X YEP + dextrose adjusted using HCl and NaOH, respectively, before adding agar; camptothecin (Sigma-Aldrich, C9911): 0.1 μ g/mL, 0.5 μ g/mL, or 1.0 μ g/mL in YPD; sorbitol (Sigma-Aldrich, S1876): 0.5 M, 1.0 M, 1.5 M, or 2.0 M in YPD; 6-azauracil (Sigma-Aldrich, A1757): 100 μ g/mL in YPD; medium; hydroxyurea (Sigma-Aldrich, H8627): 0.2 M in YPD; MMS: methyl methanesulfonate (Sigma-Aldrich, 129925), 0.05% in YPD; benomyl (Sigma-Aldrich, 381586): 15 μ g/mL in YPD; cycloheximide (Sigma-Aldrich, 01810): 10 μ g/mL in YPD liquid medium for 2 h followed by plating to YPD; H₂O₂ (Millipore, 88597): 1 mM in YPD liquid medium for 2 h followed by plating to YPD. For chromoduction experiments, SC–Lys–Arg or SC–Leu–Arg plates were used with cycloheximide (10 μ g/mL) and canavanine (12 μ g/mL) added. For chromosome destabilization, strains were grown in YEP with 2% galactose and 1% raffinose added. Sporulation medium was prepared using a 50X base consisting of 50 g potassium acetate and 0.25 g zinc acetate dihydrate in 100 mL H₂O. Final 1X sporulation media was prepared from 2 mL of 50X sporulation medium base plus 300 μ L of 10% yeast extract, 200 μ M uracil, 2 mM leucine 300 μ M histidine, and H₂O to 100 mL.

Minichunk assembly

Minichunks were originally assembled according to the following process, similar to the one previously described for *synIll*⁵: First, approximately 70-mer oligonucleotides with 20-mer overlaps were designed using GeneDesign software⁴⁰ and assembled by members of the Build-A-Genome class at Johns Hopkins University using polymerase cycling assembly (PCA) to produce 750 bp building blocks. Building blocks were verified by sequencing. Overlapping building blocks with approximately 40 bp overlaps were then assembled into 2–4 kb minichunks by direct homologous recombination in yeast. Minichunks were recovered from yeast into *E. coli*, and the recovered minichunk plasmids were verified by sequencing.

Later minichunks were ordered directly from DNA synthesis vendors with flanking *Not*I or *Bbs*I sites, verified by sequencing, excised from the vectors in which they were built via restriction digestion or, for those containing internal *Not*I or *Bbs*I sites, amplified by PCR, and integrated directly into the *synIX* strain or used for megachunk assembly.

Megachunk assembly and verification

Unlike the subsequent chromosomes, *synIX* was initially designed in separate "left and right" halves, which were subsequently merged. During the design process, *synIXL* was originally divided into megachunks A-F and V-Z. Following the integration of megachunks A-F as described in the "Minichunk integration by SwAP-In" section, the minichunks comprising megachunks V-Z in the original *synIX* design were reapportioned into three new megachunks, designated G, H, and I. Each megachunk plasmid was designed with flanking *Not*I sites for eventual excision of the assembled megachunk, 13–16 minichunks, a terminal selectable marker (*LEU2* or *URA3*, for use with SwAP-In), and homology arms between the first and last minichunks and the plasmid backbone. Overlaps between segments ranged from approximately 100-800 bp. Minichunks were released from their backbone plasmids or amplified by PCR, and were co-transformed with linkers and megachunk backbone plasmid into wild-type yeast (BY4741). Following plasmid assembly in yeast by homologous recombination and two days of growth on appropriate selective medium (SC–Leu or SC–Ura) at 30°C, single colonies were isolated and assessed by PCR to detect the presence or absence of each desired assembly junction. Colonies containing all desired junctions were recovered from yeast by phenol extraction and isopropanol precipitation, and subsequently electroporated into EPI300 *E. coli* cells (Lucigen, EC300150). Megachunk plasmids were extracted from *E. coli* using the PureLink HiPure Plasmid Midiprep Kit (Invitrogen, K2100-15).

To verify the sequences of the recovered megachunk plasmids, DNA sequencing libraries were prepared using 300 ng of plasmid DNA as input for the NEBNext Ultra II FS DNA Library Kit (NEB, E7805). DNA libraries were sequenced via Illumina NextSeq 500 using 36 bp pair-end reads. Following sequencing, reads were first processed using Trimmomatic v0.39⁴² and FastQC v0.11.4, aligned to an appropriate reference sequence using bowtie2 v2.2.9, and visualized using IGV v2.7.2.

Minichunk integration by SwAP-In

Minichunks were released from their backbones by restriction enzyme digestion, and the minichunks corresponding to one megachunk in the *synIX* design sequence (approximately 11–16 minichunks) were cotransformed into the partially completed *synIX* strain. Integration began at the left telomere of *chrIX* and proceeded in a stepwise fashion toward the centromere. At each step, the rightmost minichunk integrated contained either a *URA3* or a *LEU2* auxotrophic marker. These markers were alternated at each step according to the principles of Switching Auxotrophies Progressively for Integration (SwAP-In), ^{2,6} and colonies containing the new marker and lacking the previous strain's marker were identified by replica plating. Strains were assessed by PCRTag analysis after each round of integration.

Megachunk integration by SwAP-In

Sequence-verified megachunks were released from their assembly backbones via NotI digestion. Approximately 1 μ g of digest product was transformed into the semisynthetic synIX strain, and cells were grown for two days at 30°C on appropriate SC dropout plates to select for the desired integration product. As with minichunk integration by SwAP-In, alternating LEU2 and URA3 auxotrophic





markers were used each round, enabling selection of colonies containing the desired new integration marker and a lack of the previous round's integration marker, as determined by replica plating. Colonies were assessed by PCRTag analysis, and colonies containing the desired PCRTags of interest were used for subsequent megachunk integration rounds. After integration of all three megachunks introduced by this method, yeast strain sequences were verified by whole genome sequencing, as described in the "Whole genome sequencing" section. To remove the URA3 marker introduced during megachunk I integration, a PCR product with homology arms flanking URA3 was transformed into the synIX strain, and colonies lacking the URA3 marker were obtained following growth on 5-FOA medium.

PCRTag analysis

Cells from a single yeast colony were scraped from a plate using a pipette tip and resuspended in 30 μL of 20 mM NaOH in a 96-well PCR plate. The PCR plate was sealed and placed in a thermal cycler using the following boiling cycle: 3 cycles of 98°C for 3 min and 4°C for 1 min 1 μL of boiled product and 0.5 μM each primer were used in a 5 μL reaction using the GoTaq Green PCR system (Promega, M7123). An acoustic liquid handler (Labcyte, Echo 550) transferred primers and DNA template. Samples were run through the following protocol in a 384-well thermal cycler: 95°C for 5 min, 30 cycles of 95°C for 30 s, 55°C for 90 s, and 72°C for 60 s, and a final extension step of 72°C for 7 min. Samples were visualized following electrophoresis on a 1% agarose in 1X Tris-Taurine-EDTA gel containing ethidium bromide using the ChemiDoc XRS+ System (Bio-Rad).

Growth assays

A single yeast colony for each strain was inoculated into 5 mL YPD and incubated at 30°C overnight with rotation. For assays monitoring strain growth across multiple passages, overnight cultures were diluted 1:1000 in YPD and incubated at 30°C overnight with rotation until reaching the desired passage number. Overnight yeast culture was diluted to A₆₀₀ = 0.15 into 6 mL of fresh YPD liquid medium and grown at 30° C with rotation until A_{600} reached 0.5–0.6. Cultures were serially diluted in 10-fold increments in H_2 O, with a target OD of approximately 0.1 in 100 µL total volume for the first row of the wild-type control strain used to determine the starting culture volume in H₂O for each sample. 5 μL (YPD and YPG plates) or 10 μL (SC plates) of each dilution was spotted to each plate. Plates were grown at appropriate temperatures (room temperature of approximately 22°C on YPD, 30°C on YPD, YPG, and SC, and 37°C on YPD and YPG) and photographed daily. For additional media types (Figures S18 and S19), plates were prepared as described in the "Yeast media" section. Strains were serially diluted in 10-fold increments of H₂O as described above, and 5 μL of each dilution was spotted to each plate. Plates were grown at 30°C and photographed daily.

Plate reader growth assays in liquid culture

Three single yeast colonies per strain of interest were resuspended in 1 mL YPD in a 2 mL deep well plate and incubated for 24 h at 30°C with shaking at 800 rpm. Saturated cultures were diluted 1:100 in 200 μL YPD in a glass bottom 96-well plate with lid (Greiner Bio-One, 655892) and loaded into a Cytation 5 cell imaging multimode reader (Agilent). The following program was run using the Cytation 5 Gen5 software: 32 h at 37°C with continuous shaking, with A₆₀₀ measured every 10 min. Data analysis and visualization were conducted using Microsoft Excel and GraphPad Prism 9.

Genome editing using CRISPR-Cas9

A two-plasmid system with one guide RNA (gRNA) plasmid and one Cas9 plasmid was used for most editing.⁵² The Cas9 coding sequence was assembled together with a TEF1 promoter and CYC1 terminator in a pRS415 plasmid backbone (with LEU2 marker). This plasmid was pre-transformed into yeast strains prior to the introduction of the gRNA plasmid and donor DNA. Guide RNA target sequences were selected by identifying 20 bp upstream of the desired protospacer adjacent motif (PAM) sequence (NGG), gRNAs were cloned into a 2-micron plasmid (pRS426, with URA3 marker) under the control of a SNR52 promoter using Gibson assembly. A modified version of the gRNA plasmid was used for editing two loci simultaneously. In this case, a second gRNA was cloned into the plasmid under control of a RPR1 promoter and RPR1 terminator. gRNA sequences were confirmed by Sanger sequencing (Genewiz). To edit the synIX strain, approximately 50 ng of gRNA plasmid and 200-300 fmol (for minichunk patch replacement) or 1 pmol (for point mutation fixing) of donor DNA were transformed into yeast cells containing the Cas9 plasmid. Cells were grown on dropout medium to select for the presence of both the Cas9 and gRNA plasmids, and editing was confirmed using either PCRTag analysis or PCR followed by Sanger sequencing (Genewiz).

Sporulation and tetrad dissection experiments for bug mapping

The synIX strain yLHM1192 was crossed to a wild-type yeast strain (BY4742 with a URA3-pGAL-CEN9 cassette integrated near the centromere, also known as yLHM0539) to generate a heterozygous diploid (yLHM1233) with one copy of wild-type chrlX and one copy of synlX. To prepare strains for sporulation, a single colony of each strain was inoculated into 5 mL YPD and incubated at 30°C overnight with rotation. Overnight cultures were diluted to an OD of \sim 1 in YPD and grown to an OD of \sim 4, washed five times with H₂O, and resuspended in 2 mL of 1X sporulation medium. Strains were incubated at room temperature for 7–10 days with rotation, and monitored for the presence of tetrads. For tetrad dissection, 100 µL of these resuspended yeast cells in sporulation medium were washed and incubated with 25 μL of 0.5 mg/mL zymolyase in 1M sorbitol for 8 min 200 μL of 1M sorbitol was added to the cells, and 10 µL of the resulting mixture was added to a YPD plate. Tetrads were separated and picked using a dissection microscope

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(Singer Instruments). Spores were grown for 2–3 days on YPD until visible colonies emerged. These strains were used for growth assays as described in the "Growth assays" section to classify each spore as healthy (similar fitness to the wild-type parent) or sick (similar fitness to the *synIX* parent). Strains from eighteen four-spore tetrads (72 strains) were prepared for whole genome sequencing, as described in the "Whole genome sequencing" section below.

Plasmid cloning and transformation for synlX bug mapping

Wild-type genes of interest plus approximately 500 bp of upstream flanking sequence and 200 bp of downstream flanking sequence were amplified from BY4741 by PCR. Genes were cloned into plasmid pRS413 linearized with Afel by Gibson assembly and transformed into TOP10 cells. Colonies were grown up in 2 mL LB + 75 μ g/mL carbenicillin and miniprepped using the Zyppy Plasmid Miniprep Kit (Zymo Research, D4037). Plasmid insert sequences were verified by Sanger sequencing. Plasmids were subsequently transformed into the synIX strain and grown for two days on SC–His plates. Yeast colonies containing each plasmid of interest were isolated, grown in SC–His liquid medium, and used for growth assays (as described in the "Growth assays" section) to assess their relative fitness compared to the starting synIX strain.

Selective destabilization and loss of one copy of synlX from disomic strain

A cassette containing a galactose-inducible centromere and a *URA3* marker (*URA3-pGAL-CEN9*) with homology arms matching *synIX* was amplified by PCR and transformed into the *synIX* strain. After two days of growth at 30°C on SC–Ura plates, single colonies were purified and assessed by junction PCR for the insertion of the *URA3-pGAL-CEN9* fragment. Strains were then grown in YEP +2% galactose +1% raffinose for two days and diluted to 5-FOA plates to obtain single colonies lacking *URA3*. Strains were single-colony purified and prepared for whole genome sequencing to assess chromosome copy number.

Whole genome sequencing

A single yeast colony was inoculated into 5 mL of YPD liquid medium at 30°C overnight with rotation. Overnight cultured yeast cells were harvested by centrifugation at 3000 x g for 3 min, and yeast genomic DNA was extracted using the Fungi/Yeast Genomic DNA Isolation Kit (Norgen, 27300). Genomic DNA concentration was determined using the Qubit dsDNA HS Assay Kit (Thermo Fisher Scientific, Q32854). DNA sequencing libraries were prepared using 300 ng of yeast genomic DNA as input for the NEBNext Ultra II FS DNA Library Kit (NEB, E7805). DNA libraries were sequenced via Illumina NextSeq 500 using either 75 bp or 36 bp pair-end reads. Following sequencing, reads were first processed using Trimmomatic v0.39⁴² and FastQC v0.11.4, ⁴³ aligned to an appropriate reference genome using bowtie2 v2.2.9, ⁴⁴ and visualized using IGV v2.7.2. ⁴¹ Chromosome coverage plots were prepared in Rstudio version 1.3.1093 using the ggplot2 v3.3.5 package. ⁴⁵

Southern blot analysis

A single yeast colony was inoculated into 10 mL of YPD medium at 30°C overnight with rotation. Overnight cultured yeast cells were harvested by centrifugation at 3000 x g for 3 min, and yeast genomic DNA was extracted using the MasterPure Yeast DNA Purification Kit (Lucigen, MPY80200) according to the manufacturer's instructions, with an RNase A treatment step. DNA was quantified using the Qubit dsDNA BR Assay Kit (Thermo Fisher Scientific, Q32850). 2 μg of DNA were digested with XhoI at 37°C overnight. Digests were electrophoresed on a 1% Tris-Borate-EDTA gel in 0.5X Tris-Borate-EDTA running buffer for 10 h at 50V. Following electrophoresis, the gel was washed in depurination solution (0.25 M HCl) for 15 min, denaturation solution (0.5 M NaOH, 1.5 M NaCl) for 60 min, and neutralization solution (0.5 M Tris, 1.5 M NaCl, pH 7.5) for 30 min. Chromosomes were transferred by capillarity to a nylon membrane (Pall Corporation, 60207) using 10X SSC buffer (20X SSC: 1.5 M NaCl, 0.3 M sodium citrate, pH 7) for 18 h. After transfer, the membrane was washed in 2X SSC buffer for 5 min and baked at 80°C for 30 min. Prehybridization was performed for 60 min in ULTRAhyb Ultrasensitive Hybridization Buffer (Invitrogen, AM8669) at 55°C with rotation. For hybridization, probes were prepared by labeling a ∼360 bp PCR fragment of universal telomere cap sequence from plasmid pJS160 using the PCR DIG Probe Synthesis Kit (Roche, 11636090910) with oligos 5'- GCTATACGAAGTTATTAGGGTAGTGTG-3' and 5'- CTGCAGGTCGACTCTAGAGGATC-3', according to the manufacturer's instructions. The following thermocycler conditions were used: 1 cycle of 95°C for 2 min, 30 cycles of 95°C for 30 s, 60°C for 30 s, and 72°C for 40 s, and a final elongation step of 72°C for 7 min. Probes were purified using Zymo-Spin I Columns (Zymo Research, C1003) and denatured at 95°C for 5 min. Hybridization was performed at 55°C overnight with rotation in 10 mL ULTRAhyb Ultrasensitive Hybridization Buffer (Invitrogen, AM8669) using 500 ng of probe per experiment. The blot was washed twice with 2X SSPE buffer (20X SSPE buffer: 0.02 M EDTA and 2.98 M NaCl in 0.2 M phosphate buffer, pH 7.4; Sigma-Aldrich, S2015) for 5 min at room temperature, twice with 2X SSPE +1% SDS for 30 min at 55°C, and twice with 0.1X SSPE for 15 min at 55°C. Blot was incubated overnight in 1:1 Intercept (TBS) Blocking Buffer (LI-COR Biosciences, 927-60001): 1X Tris-Buffered Saline (TBS) with 0.1% TWEEN 20 (Sigma-Aldrich, P1379) (1X TBST) and 1% SDS at room temperature. The next day, the membrane was washed with 1X TBST for 5 min and incubated for 1 h at room temperature with rabbit anti-DIG antibody (working concentration 1:2500 in 1:1 Intercept (TBS) Blocking Buffer: 1X TBST; Invitrogen, 700772). The primary antibody solution was washed out three times with 1X TBST for 15 min at room temperature. The membrane was incubated in goat anti-rabbit IgG secondary solution (LI-COR Biosciences, 926-68071, used at 1:10,000 in 1:1 Intercept (TBS) Blocking Buffer (LI-COR Biosciences, 927-60001): 1X TBST with 1% SDS) for 1.5 h. The blot was washed three times with 1X TBST for 15 min at room temperature plus one time with 1X TBS for 30 min at room temperature. An LI-COR Odyssey Instrument was used to develop the blot images. St





RNA extraction, sequencing, and analysis for short read sequencing

A single yeast colony was inoculated into 5 mL of YPD liquid medium at 30°C overnight with rotation. Overnight yeast culture was diluted to A₆₀₀ = 0.15–0.2 into 6 mL of fresh YPD liquid medium and grown at 30°C or 37°C with rotation until A₆₀₀ reached 0.8– 1.0. Yeast cells were harvested from 4 mL of culture, and total RNA was extracted using the RNeasy Mini Kit (QIAGEN, 74106) according to the manufacturer's instructions. RNA was quantified using the Qubit BR RNA Assay Kit (Thermo Fisher Scientific, Q10210). 1 µg total RNA was used as input for the QIAseq Stranded Total RNA Library Kit (QIAGEN, 180745) plus QIAseq FastSelect -rRNA Yeast Kit (QIAGEN, 334217). RNA libraries were sequenced via Illumina NextSeq 500 using 75 bp paired-end reads. For applications requiring similar numbers of reads per sample, downsampling was performed using segtk v1.3. Samples were first processed using Trimmomatic v0.39⁴² and FastQC v0.11.4.⁴³ Processed reads were aligned to the S288C transcriptome and a custom synIX transcriptome (in which chrlX transcripts were replaced with their synthetic versions) using Kallisto v0.46.0⁴⁶ and STAR v2.5.2a.⁴⁷ STAR-aligned results were further processed using SAMtools v1.948 and BEDtools v2.26.0,49 and visualized in IGV v2.7.2.41 Kallisto pseudoalignment data was further processed in Rstudio version 1.3.1093 using the sleuth v0.30.0⁵⁰ and ggplot2 v3.3.5⁴⁵ packages. For volcano plots, we calculated effect scores (approximately log2 fold change values), tested results for significance using Wald's test, and corrected for multiple testing with the false discovery rate adjusted p value using the Benjamini-Hochberg method. The following genes deleted from the synIX design were removed from the plots in Figures 6D and 6E: YIL173W (VTH1), YIL171W (part of HTX12 pseudogene), YIL170W (part of HTX12 pseudogene), YIL169C (CSS1), YIL167W (SDL1, blocked reading frame), YIL082W, YIL060W, YIL059C, YIL058W, YILWTy3-1, YILCdelta3. The following dubious ORFs, retrotransposable elements, and subtelomeric genes were also removed: YBL100C, YGR296C-B, YHL050W-A, YHR219C-A, YIL020C-A, YJR029W, YLL066C, YML133W-B, YMR046C, YNL339W-B, YNL339C, YOR277C, YPL283W-B, YPR158W-B, YPR204C-A, YBLWdelta4, YCLWdelta3. Unfiltered versions of the volcano plots are depicted in Figure S20.

RNA extraction, sequencing, and analysis for nanopore direct RNA sequencing

Methods recently described by us were used to perform nanopore sequencing to analyze RNA isoforms. 54 Total RNA was extracted from 50 mL flash-frozen cell pellets grown to mid-log (OD₆₀₀ \sim 0.65–0.85) using the MasterPure Yeast RNA Purification Kit (Lucigen) including a DNase I treatment step. RNA (diluted 1:10) quality and concentration were measured by Agilent 2100 Bioanalyzer with the Agilent RNA 6000 Nano Kit and Qubit RNA High Sensitivity Kit (Thermo Fisher Scientific), respectively. Poly(A) mRNA was enriched from 50 μg total RNA on 132 μL Dynabeads oligo(dT)₂₅ beads. The Direct RNA Sequencing Kit (SQK-RNA002, Oxford Nanopore Technologies) was used to generate libraries from 500 ng poly(A) RNA. An optional reverse transcription was performed at 50°C for 50 min using SuperScript IV Reverse Transcriptase (Invitrogen) in between the ligation of the RTA and RMX adaptors. Following reverse transcription the RNA:cDNA was cleaned up with 1.8 volumes of Agencourt RNAClean XP beads and washed with 70% ethanol. Following RMX ligation only one volume of beads was used in the clean-up, and WSB (SQK-RNA002) was used in the wash steps. Direct RNA libraries (typically 150-200 ng) were loaded onto primed (EXP-FLP001) MinION flow cells (FLO-MIN106D, R9 version) in RRB buffer and run on the GridION with MinKNOW 5.2.2 for up to 72 h. Nanopore long reads were base-called, trimmed of adapter sequences, and filtered for quality, retaining only those with the best alignment scores for multi-mapping reads, as previously described. 54 For TSS distributions, long-reads that overlapped at least 75% of the annotated gene were used.

Construction of FLAG-tagged EST3 strains

A DNA fragment containing a six-glycine linker plus a 3X FLAG tag (DYKDHDG-DYKDHDI-DYKDDDDK) was designed for integration at the 3' end of EST3, in-frame with the rest of the protein and inserted just before the EST3 stop codon. The segment containing the tag was flanked by 70-100 bp homology arms matching the yeast genome sequence adjacent to the desired insertion site. Genome editing using CRISPR-Cas9 in yeast (as describe above) was used to cut the genome near the end of the EST3 coding sequence and insert the tag of interest. Colony PCR and Sanger sequencing (Genewiz) were used to confirm in-frame insertion of the tag.

Immunoblot analysis

A single yeast colony was inoculated into 5 mL of YPD liquid medium at 37°C overnight with rotation. Overnight culture was diluted to A₆₀₀ = 0.15–0.2 into 50 mL of fresh YPD liquid medium and grown at 37°C with shaking at 200 rpm until A₆₀₀ reached 0.8–1.0. Yeast were harvested by centrifugation, and pellets were frozen overnight at -80° C. Pellets were subsequently resuspended in 500 μ L of lysis buffer (20 mM HEPES pH 7.4, 1% Triton X-100 (Sigma-Aldrich, X100), 2 mM MgCl₂, 500 μM NaCl, and 1X protease inhibitor (Roche)). Cell suspension was added to MP Biomedicals FastPrep Lysing Matrix C and subjected to the following shaking program at 4°C using a FastPrep-24 5G (MP Biomedicals): 7 cycles of 15 s shaking at 6.0 m/s + 30 s rest between cycles. Samples were centrifuged at 13000 x g for 10 min at 4°C, and 30 μL of supernatant was transferred to a new tube and mixed with 10 μL of 4X LDS loading buffer (Invitrogen, NP0007) (pre-immunoprecipitation sample). The remaining supernatant was transferred to pre-equilibrated FLAG M2 (Sigma F1804)-conjugated M-270 Dynabeads (Invitrogen, 14302D) for immunoprecipitation. 55 Samples with beads were incubated on ice for 60 min, washed 3 times with lysis buffer, and eluted at 70°C with shaking at 700 rpm in 50 μL 1.1X LDS loading buffer (Invitrogen, NP0007) in lysis buffer for 10 min 2-mercaptoethanol (1.43 M final) was added, and samples were heated to 95°C for 5 min and loaded onto an SDS-PAGE gel for immunoblotting. Mouse anti-FLAG M2 antibody (Sigma F1804, used at 1:5000) and rabbit antihistone H3 antibody (Abcam, 1791, used at 1:5000) were used for blotting proteins of interest overnight at 4°C. Goat anti-mouse IgG1 (LI-COR Biosciences, 926-32350, used at 1:20,000) and goat anti-rabbit IgG (LI-COR Biosciences, 926-68071, used at 1:20,000)

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were used for blotting FLAG and H3 primary antibodies, respectively, during a 1-h incubation at room temperature. An LI-COR Odyssey Instrument was used to develop the blot images.⁵³

Pulsed-field gel electrophoresis

A single yeast colony was inoculated into 7 mL of YPD liquid medium and grown at 30°C for two days with rotation. Approximately 40 mg of yeast cells were harvested by centrifugation. To prepare plugs, 16 μ L of zymolyase solution (25 mg/mL zymolyase 20T in 10 mM KH₂PO₂ pH 7.5) and 360 μ L of 0.5% melted low melting point agarose (Bio-Rad, 1620137) in 100 mM EDTA pH 7.5 were added to the cell pellet and mixed by pipetting. 90 μ L of this suspension was added per well of a BioRad plug mold, and the plugs were cooled for 30 min at 4°C. Solidified plugs were then transferred into 15 mL tubes containing 1 mL of digestion buffer (0.5 M EDTA, 10 mM Tris, pH 7.5), mixed by inversion, and incubated at 37°C with rotation overnight. The next day, 400 μ L of proteinase K solution (5% sarcosyl and 5 mg/mL proteinase K in 0.5 M EDTA, pH 7.5) were added to the tube. Samples were incubated at 50°C for 5 h, and then washed once with H₂O and three times with TE (2 mM Tris, 1 mM EDTA, pH 8.0). Plugs were stored at 4°C until ready for gel loading. Half of one plug per sample was embedded per well of a gel consisting of 1% low melting point agarose in 0.5X Tris-Borate-EDTA buffer. Samples were separated in running buffer of 0.5X Tris-Borate-EDTA by clamped homogeneous electric field gel electrophoresis using the CHEF Mapper XA Pulsed-Field Electrophoresis System (Bio-Rad), as previously described. The following program was used: auto-algorithm with size range of 200 kb to 2.5 Mb, temperature of 14°C, voltage of 6 V/cm, included angle of 120°. After electrophoresis, gels were stained using 5 μ g/mL ethidium bromide in H₂O for 30 min, destained in H₂O for 30 min, and then imaged using the ChemiDoc XRS+ System (Bio-Rad).

Chromosome substitution and conditional centromere destabilization

For initial experiments, disomic synIX strain yLHM0387 was used as the donor and yWZ601 as the recipient for kar1-1-mediated chromoduction. For follow-up experiments, disomic strain yLHM0604 and monosomic strains yLHM0721, yLHM1591, and yLHM1601 were used as donors and yWZ602 as the recipient for kar1-1-mediated chromoduction. yWZ601 and yWZ602 were derived from DMy04425 following deletion of LYS12 from wild-type chrlX and insertion of a URA3-pGAL promoter cassette upstream of CEN9.²⁷ Strains were grown independently on YPD plates, and subsequently mated on YPD plates via replica plating. After 14 h of growth, these plates were replica plated to SC-Lys-Arg plates with 12 μg/mL canavanine and 10 μg/mL cycloheximide added. Plates were grown until individual colonies emerged; if a patched formed instead, patches were replica plated to the same selective medium and grown for additional time until individual colonies emerged. Single colonies were restreaked to fresh selective plates and analyzed by PCRTagging, pulsed-field gel electrophoresis, and whole genome sequencing. To lose wild-type chrlX, strains were grown in YPD overnight, diluted 1:1000 in YEP +2% galactose medium for 24 h, and plated on 5-FOA medium to select for strains that had lost wild-type chrlX. Single colonies were genotyped by PCRTag analysis and further characterized by pulsed-field gel electrophoresis and whole genome sequencing. For chromosome substitution to move syn/X from yLHM1601 to recipient strain yLHM2231 (derived from BY4742), a LEU2 marker was integrated into synIX near the end of megachunk H, in minichunk 101, generating strain yLHM1754. The same protocol as listed above was followed using donor strain yLHM1754 and recipient strain yLHM2231, except after mating and 14 h of growth, plates were replica plated to SC-Leu-Arg (rather than SC-Lys-Arg) plates with 12 μg/mL canavanine and 10 μg/mL cycloheximide added.

QUANTIFICATION AND STATISTICAL ANALYSIS

Information on the number of biological replicates, statistical tests, and p values used is provided in the Figure legends and "Method Details" section.