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Reduced Gene Dosage of Histone H4 Prevents CENP-A Mislocalization in Budding

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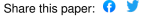
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Reduced Gene Dosage of Histone H4 Prevents CENP-A Mislocalization in Saccharomyces

2 *cerevisiae*

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17 ABSTRACT

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Mislocalization of the centromeric histone H3 variant (Cse4 in budding yeast, CID in flies, CENP-A in humans) to non-centromeric regions contributes to chromosomal instability (CIN) in yeast, fly, and human cells. Overexpression and mislocalization of CENP-A has been observed in cancers, however, the mechanisms that facilitate the mislocalization of overexpressed CENP-A have not been fully explored. Defects in ubiquitin-mediated proteolysis of overexpressed Cse4 (GALCSE4) leads to its mislocalization and synthetic dosage lethality (SDL) in mutants for E3 ubiquitin ligases (Psh1, Slx5, SCF^{Met30}, SCF^{Cdc4}), Doa1, Hir2, and Cdc7. In contrast, defects in sumoylation of GALcse4K215/216/A/R prevent its mislocalization and do not cause SDL in a $psh1\Delta$ strain. Here, we used a genome-wide screen to identify factors that facilitate the mislocalization of overexpressed Cse4 by characterizing suppressors of the psh1\Delta GALCSE4 SDL. Deletions of histone H4 alleles (*HHF1* or *HHF2*), which were among the most prominent suppressors, also suppress $slx5\Delta$, cdc4-1, $doa1\Delta$, $hir2\Delta$, and cdc7-4 GALCSE4 SDL. Reduced dosage of H4 contributes to defects in sumoylation and reduced mislocalization of overexpressed Cse4. We determined that the hhf1-20, cse4-102, and cse4-111 mutants, which are defective in the Cse4-H4 interaction, also exhibit reduced sumoylation of Cse4 and do not display $psh1\Delta$ GALCSE4 SDL. In summary, we have identified genes that contribute to the mislocalization of overexpressed Cse4 and defined a role for the gene dosage of H4 in facilitating Cse4 sumoylation and mislocalization to non-centromeric regions, contributing to SDL when Cse4 is overexpressed in mutant strains.

INTRODUCTION

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Centromeres are specialized chromosome loci that are essential for faithful chromosome segregation during mitosis and meiosis. The kinetochore (centromeric DNA and associated proteins) provides an attachment site for microtubules to promote proper segregation of sister chromatids during cell division (Allshire and Karpen 2008; Verdaasdonk and Bloom 2011; BURRACK AND BERMAN 2012; CHOY et al. 2012; MADDOX et al. 2012; MCKINLEY AND CHEESEMAN 2016). Despite the wide divergence of centromeric DNA sequence, establishment of centromeric chromatin is regulated by epigenetic mechanisms where incorporation of the essential and evolutionarily conserved centromeric histone H3 variant CENP-A (Cse4 in Saccharomyces cerevisiae, Cnp1 in Schizosaccharomyces pombe, CID in Drosophila melanogaster, and CENP-A in mammals) serves to nucleate kinetochore assembly (KITAGAWA AND HIETER 2001; BIGGINS 2013; McKinley and Cheeseman 2016). The evolutionarily conserved CENP-A-specific histone chaperones (Scm3 in S. cerevisiae and S. pombe, CAL1 in D. melanogaster, Holliday Junction Recognition Protein HJURP in humans) mediate the centromeric localization of CENP-A (CAMAHORT et al. 2007; MIZUGUCHI et al. 2007; STOLER et al. 2007; FOLTZ et al. 2009; PIDOUX et al. 2009; WILLIAMS et al. 2009; SHUAIB et al. 2010; CHEN et al. 2014). In budding yeast, other chaperones such as Chromatin Assembly Factor 1 (CAF-1), an evolutionarily conserved replication-coupled histone H3/H4 chaperone, can facilitate the deposition of overexpressed Cse4 when Scm3 is depleted (HEWAWASAM et al. 2018). The CAF-1 orthologues Mis16 in S. pombe and RbAp46/48 in humans and D. melanogaster also contribute to centromeric localization of CENP-A (FUJITA et al. 2007; Pidoux et al. 2009; Williams et al. 2009; Boltengagen et al. 2016).

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Restricting the localization of CENP-A to centromeres is essential for faithful chromosome segregation. However, overexpression of CENP-A leads to its mislocalization to non-centromeric chromatin and contributes to chromosomal instability (CIN) in yeast, flies, and humans (Collins et al. 2004; Heun et al. 2006; Moreno-Moreno et al. 2006; Au et al. 2008; MISHRA et al. 2011; LACOSTE et al. 2014; ATHWAL et al. 2015; SHRESTHA et al. 2017). Overexpression and mislocalization of CENP-A is observed in many cancers and is proposed to promote tumorigenesis (TOMONAGA et al. 2003; AMATO et al. 2009; LI et al. 2011; MCGOVERN et al. 2012; SUN et al. 2016). Thus, defining the molecular mechanisms that promote and prevent mislocalization of CENP-A is an area of active investigation. In budding yeast, post-translational modifications (PTMs) of Cse4, such as ubiquitination, sumoylation, and isomerization, are important for regulating steady-state levels of Cse4 and preventing its mislocalization to non-centromeric regions, thereby maintaining chromosome stability (COLLINS et al. 2004; HEWAWASAM et al. 2010; RANJITKAR et al. 2010; OHKUNI et al. 2014; OHKUNI et al. 2016; CHENG et al. 2017; AU et al. 2020). Ubiquitin-mediated proteolysis of Cse4 by E3 ubiquitin ligases such as Psh1 (HEWAWASAM et al. 2010; RANJITKAR et al. 2010), SUMO-targeted ubiquitin ligase (STUbL) Slx5 (OHKUNI et al. 2016), SCF^{Met30/Cdc4} (AU et al. 2020), SCF^{Rcy1} (CHENG et al. 2016), and Ubr1 (CHENG et al. 2017) and the proline isomerase Fpr3 (OHKUNI et al. 2014) regulate the cellular levels of Cse4. Psh1-mediated proteolysis of Cse4 has been well characterized and has been shown to be regulated by the FACT (Facilitates Chromatin Transcription/Transactions) complex (DEYTER AND BIGGINS 2014), CK2 (Casein Kinase 2) (HEWAWASAM et al. 2014), HIR (HIstone Regulation) histone chaperone complex (CIFTCI-YILMAZ et al. 2018), and DDK (Dbf4-Dependent Kinase) complex (EISENSTATT et al. 2020). In general, mutation or deletion of factors that prevent Cse4

mislocalization show synthetic dosage lethality (SDL) when Cse4 is overexpressed from a galactose-inducible promoter (*GALCSE4*).

In contrast to the many studies that have characterized pathways that prevent mislocalization of CENP-A to non-centromeric regions, mechanisms that facilitate the mislocalization of overexpressed CENP-A have not been fully explored. Studies from our laboratory and those of others show that the transcription-coupled histone H3/H4 chaperone DAXX/ATRX promotes mislocalization of CENP-A to non-centromeric regions in human cells (LACOSTE *et al.* 2014; SHRESTHA *et al.* 2017). In budding yeast, CAF-1 contributes to the mislocalization of overexpressed Cse4 to non-centromeric regions (HEWAWASAM *et al.* 2018). We have recently shown that sumoylation of Cse4K215/216 in the C-terminus of Cse4 facilitates its interaction with CAF-1 and this promotes the deposition of Cse4 to non-centromeric regions (OHKUNI *et al.* 2020). Notably, *psh1*Δ *cac2*Δ *GALCSE4* strains and *psh1*Δ *GALcse4K215/216R/A* strains do not exhibit SDL due to reduced mislocalization of Cse4 (HEWAWASAM *et al.* 2018; OHKUNI *et al.* 2020).

Defining the mechanisms that facilitate the mislocalization of overexpressed Cse4 to non-centromeric regions is essential for understanding which pathways contributes to mislocalization of CENP-A in cancers with a poor prognosis. We performed a genome-wide screen using a synthetic genetic array (SGA) to identify genes that promote Cse4 mislocalization. We took advantage of the SDL of a $psh1\Delta$ GALCSE4 strain (HEWAWASAM et al. 2010; RANJITKAR et al. 2010; AU et al. 2013) to identify suppressors of the SDL phenotype. An SGA analysis was performed by combining mutants of essential genes and deletion of non-essential genes with $psh1\Delta$ GALCSE4. The screen identified mutations or deletions of genes encoding regulators of chromatin remodeling, RNA transcription/processing, nucleosome occupancy, ubiquitination,

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and histone H4. The budding yeast genome possesses two gene pairs which encode almost identical H3 and H4 proteins (HHT1/HHF1 and HHT2/HHF2) and two gene pairs which encode identical H2A and H2B proteins (HTA1/HTB1 and HTA2/HTB2). Deletion of the two alleles that encode histone H4 (HHF1 or HHF2) were among the most prominent suppressors of the $psh1\Delta$ GALCSE4 SDL. A role for the dosage of H4 in preventing mislocalization of Cse4 has not been previously examined. In this study, we focused on defining the molecular mechanisms that prevent the mislocalization of overexpressed Cse4 and suppress the psh1\(\Delta\) GALCSE4 SDL when the gene dosage of H4 is reduced. We showed that deletion of HHF1 or HHF2 also suppresses the GALCSE4 SDL in $slx5\Delta$, $doa1\Delta$, $hir2\Delta$, cdc4-1, and cdc7-4 strains. Deletion of HHF1 or HHF2 results in reduced Cse4 sumoylation and this correlates with reduced mislocalization to noncentromeric regions and rapid degradation of Cse4 in a psh1\(\Delta\) strain. Moreover, cse4-102, cse4-111, and hhf1-20, which have mutations in their histone fold domains and are defective for the formation of the Cse4-H4 dimer (SMITH et al. 1996; GLOWCZEWSKI et al. 2000), show reduced Cse4 sumoylation and do not cause SDL in psh1∆ GALCSE4 strains. In summary, our genomewide suppressor screen allowed us to identify genes that contribute to Cse4 mislocalization and to define a role for reduced gene dosage of H4 in preventing the mislocalization of Cse4 to non-

centromeric regions and suppression of the psh1\(\triangle \) GALCSE4 SDL.

MATERIALS AND METHODS

Strains and plasmids

Yeast strains used in this study are described in Table S2 and plasmids in Table S3. Yeast strains were grown in rich media (1% yeast extract, 2% bacto-peptone, 2% glucose) or synthetic medium with glucose or raffinose and galactose (2% final concentration each) and supplements to allow for selection of the indicated plasmids. Double mutant strains were generated by mating wild type or *psh1*∆ strains with empty vector or a plasmid containing *GAL1-6His-3HA-CSE4* to mutant strains on rich medium at room temperature for six hours followed by selection of diploid cells on medium selective for the plasmid and appropriate resistance markers. Diploids were sporulated for 5 days at 23°C and plated on selective medium without uracil, histidine, or arginine and with canavanine, clonNAT, and G418 to select for *MATa* double mutants. The synthetic genetic array (SGA) was performed as previously described (COSTANZO *et al.* 2016).

Growth assays

Growth assays were performed as previously described (EISENSTATT *et al.* 2020). Wild type and mutant strains were grown on medium selective for the plasmid, suspended in water to a concentration with an optical density of 1 measured at a wavelength of 600 nm (OD_{600} , approximately 1.0 X 10^7 cells per ml), and plated in five-fold serial dilutions starting with 1 OD_{600} on synthetic growth medium containing glucose or galactose and raffinose (2% final concentration each) selecting for the plasmid. Strains were grown at the indicated temperatures for 3-5 days.

Protein stability assays

Protein stability assays were performed as previously described (AU *et al.* 2008). Briefly, logarithmically growing wild type and mutant cells were grown for three to four hours in media selective for the plasmid containing galactose/raffinose (2% final concentration each) at 30°C followed by addition of cycloheximide (CHX, 10 µg/ml) and glucose (2% final concentration).

Protein extracts were prepared from cells collected 0, 30, 60, and 90 minutes after CHX addition with the TCA method as described previously (KASTENMAYER *et al.* 2006). Equal amount of protein as determined by the Bio-Rad DCTM Protein Assay were analyzed by Western blot. Proteins were separated by SDS-PAGE on 4-12% Bis-TRIS SDS-polyacrylamide gels (Novex, NP0322BOX) and analysis was done against primary antibodies α-HA (1:1000, Roche, 12CA5) or α-Tub2 (1:4500, custom made for Basrai Laboratory) in TBST containing 5% (w/v) dried skim milk. HRP-conjugated sheep α-mouse IgG (Amersham Biosciences, NA931V) and HRP-conjugated donkey α-rabbit IgG (Amersham Biosciences, NA934V) were used as secondary antibodies. Stability of the Cse4 protein relative to the Tub2 loading control was measured as the percent remaining as determined with the Image Lab Software (BioRad).

Ubiquitination pull-down assay

Levels of ubiquitinated Cse4 were determined with ubiquitin pull-down assays as described previously (AU *et al.* 2013) with modifications. Cells were grown to logarithmic phase, induced in galactose-containing medium for 3 hours at 30°C and pelleted. The cell pellet was resuspended in lysis buffer (20 mM Na₂HPO₄, 20 mM NAH₂PO₄, 50 mM NaF, 5 mM tetrasodium pyrophosphate, 10 mM beta-glycerolphosphate, 2 mM EDTA, 1 mM DTT, 1% NP-40, 5 mM N-Ethylmaleimide, 1 mM PMSF, and protease inhibitor cocktail (Sigma, catalogue # P8215)) and equal volume of glass beads (lysing matrix C, MP Biomedicals). Cell lysates were generated by homogenizing cells with a FastPrep-24 5G homogenizer (MP Biomedicals) and a fraction of the lysate was aliquoted for input. An equal concentration of lysates from wild type and mutant strains were incubated with tandem ubiquitin binding entities (Agarose-TUBE1, Life Sensors, Inc., catalogue # UM401) overnight at 4°C. Proteins bound to the beads were washed three times with TBS-T at room temperature and eluted in 2 x Laemmli buffer at 100°C for 10 minutes. The eluted protein was resolved on a 4-12% Bis-Tris gel (Novex, NP0322BOX) and

ubiquitinated Cse4 was detected by Western blot using anti-HA antibody (Roche Inc., 12CA5). Levels of ubiquitinated Cse4 relative to the non-modified Cse4 in the input were quantified using software provided by the Syngene imaging system. The percentage of ubiquitinated Cse4 levels is set to 100% in the wild type strain.

In vivo sumoylation assay

Cell lysates were prepared from 50 ml culture of strains grown to logarithmic phase in raffinose/galactose (2% final concentration each) medium at 30°C for 4 hours to induce expression of Cse4 from the galactose-inducible promoter. Cells were pelleted, rinsed with sterile water, and suspended in 0.5 ml of guanidine buffer (0.1 M Tris-HCl at pH 8.0, 6.0 M guanidine chloride, 0.5 M NaCl). Cells were homogenized with Matrix C (MP Biomedicals) using a bead beater (MP Biomedicals, FastPrep-24 5G). Cell lysates were clarified by centrifugation at 6,000 rpm for 7 min and protein concentration was determined using a DC protein assay kit (Bio-Rad). Samples containing equal amounts of protein were brought to a total volume of 1 ml with appropriate buffer.

In vivo sumoylation was assayed in crude yeast extracts using nickel-nitrilotriacetic acid (Ni-NTA) agarose beads to pull down His-HA-tagged Cse4 as described previously (OHKUNI *et al.* 2015) with modifications. Cell lysates were incubated with 100 μl of Ni-NTA superflow beads (Qiagen, 30430) overnight at 4 °C. After being washed with guanidine buffer one time and with breaking buffer (0.1 M Tris-HCl at pH 8.0, 20 % glycerol, 1 mM PMSF) five times, beads were incubated with 2x Laemmli buffer including imidazole at 100°C for 5 min. The protein samples were analyzed by SDS-PAGE and western blotting. Primary antibodies were anti-HA (12CA5) mouse (Roche, 11583816001) and anti-Smt3 (y-84) rabbit (Santa Cruz Biotechnology, sc-28649). Secondary antibodies were ECL Mouse IgG, HRP-Linked Whole Ab (GE Healthcare Life Sciences, NA931V) or ECL Rabbit IgG, HRP-linked Whole Ab (GE Healthcare Life

Sciences, NA934V). Protein levels were quantified using Image Lab software (version 6.0.0)

from Bio-Rad Laboratories, Inc (Hercules, CA).

ChIP-qPCR

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Chromatin immunoprecipitations were performed with two biological replicates per strain as previously described (COLE et al. 2014; CHEREJI et al. 2017; EISENSTATT et al. 2020) with modifications. Logarithmic phase cultures were grown in raffinose/galactose (2% final concentration each) media for 4 hours and were treated with formaldehyde (1% final concentration) for 20 minutes at 30°C followed by the addition of 2.5 M glycine for 10 minutes at 30°C. Cell pellets were washed twice with 1 X PBS and resuspended in 2 mL FA Lysis Buffer (1 mM EDTA pH8.0, 50 mM HEPES-KOH pH7.5, 140 mM NaCl, 0.1% sodium deoxycholate, 1% Triton X-100) with 1 x protease inhibitors (Sigma) and 1 mM PMSF (final concentration). The cell suspension was split into four screw top tubes with glass beads (0.4-0.65 mm diameter) and lysed in a FastPrep-24 5G (MP Biosciences) for 40 seconds three times, allowed to rest on ice for 5 minutes, and lysed two final times for 40 seconds each. The cell lysate was collected, and the chromatin pellet was washed in FA Lysis Buffer twice. Each pellet was resuspended in 600 µl of FA Lysis Buffer and combined into one 5 ml tube. The chromatin suspension was sonicated with a Branson digital sonifer 24 times at 20% amplitude with a repeated 15 seconds on/off cycle. After 3 minutes of centrifugation (13000 rpm, 4°C), the supernatant was transferred to another tube. Input sample was removed (5%) and the average size of the DNA was analyzed. The remaining lysate was incubated with anti-HA-agarose beads (Sigma, A2095) overnight at 4°C. The beads were washed in 1 ml FA, FA-HS (500 mM NaCl), RIPA, and TE buffers for five minutes on a rotor two times each. The beads were suspended in ChIP Elution Buffer (25 mM Tris-HCl pH7.6, 100 mMNaCl, 0.5% SDS) and incubated at 65°C overnight. The beads were treated with proteinase K (0.5 mg/ml) and incubated at 55°C for four hours followed by

Phenol/Chloroform extraction and ethanol precipitation. The DNA pellet was resuspended in a total of 50 µl sterile water. Samples were analyzed by quantitative PCR (qPCR) performed with the 7500 Fast Real Time PCR System with Fast SYBR Green Master Mix (Applied Biosystems). qPCR conditions used: 95°C for 20 sec; 40 cycles of 95°C for 3 sec, 60° for 30 sec. Primers used are listed in Table S4.

Data availability

Strains and plasmids are available upon request. Supporting figures S1-S7 are available as JPG files. Supporting Table S1 is an Excel file that describes mutations that suppress the *psh1*\(\triangle \) *GALCSE4* SDL, the gene systematic name, the gene name, the functional category, growth and colony scores, and validation information if applicable. File S1 contains Tables S2, S3, and S4 which describe the yeast strains, plasmids, and primers used in this study, respectively. Supporting information is available at FigShare.

235 RESULTS

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A genome-wide screen identified suppressors of the SDL in a psh1\(\triangle \) GALCSE4 strain

Identifying pathways that facilitate the deposition of overexpressed Cse4 to noncentromeric regions will provide insight into the mechanisms that promote chromosomal instability (CIN) in CENP-A overexpressing cancers. Deletion of PSH1, which regulates ubiquitin-mediated proteolysis of overexpressed Cse4, results in synthetic dosage lethality (SDL) when Cse4 is overexpressed (GALCSE4) (HEWAWASAM et al. 2010; RANJITKAR et al. 2010). We reasoned that strains with deletions or mutations of factors that promote Cse4 mislocalization would rescue the SDL of a psh1 \triangle GALCSE4 strain. Therefore, we generated a psh1 \triangle query strain overexpressing CSE4 from a galactose-inducible plasmid and mated it to arrays of 3,827 nonessential gene deletion strains and of 786 conditional mutant alleles, encoding 560 essential genes, and 186 non-essential genes for internal controls (Costanzo et al. 2016). Growth of the haploid meiotic progeny plated in quadruplicate was visually scored on glucose-and galactosecontaining media grown at 30°C for non-essential and 26°C for essential gene mutant strains (Figure 1A). Highlighted in the figure are all four replicates of deletion of histone H4 ($hhf1\Delta$) and Hap3 (hap3\(amplie)\) showing better growth on galactose media compared to the control strains along the perimeter and other deletion strains on the plate (Figure 1B, bottom and top square, respectively). Strains that suppress the $psh1\Delta$ GALCSE4 SDL on galactose-containing media were given a growth score of one (low suppression) to four (high suppression) (Table S1). The number of replicates within the quadruplicate that displayed the same growth were given a colony score of one (one out of four replicates) to four (all four replicates). We identified ninetyfour deletion and mutant alleles encoding ninety-two genes that suppressed the psh1\(\triangle \) GALCSE4 SDL and the majority (81%) of quadruplicates had all four colonies displaying the same level of suppression, indicated by a colony score of four (Table S1).

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Of the ninety-four alleles, we selected thirty-eight candidate mutants (fourteen nonessential deletion strains and twenty-four conditional mutants) to confirm the suppression of the psh1\(\triangle GALCSE4\) SDL (Table 1). These candidates displayed a growth score of three or four where most of the replicates displayed high suppression and represent pathways involved in RNA processing and cleavage, DNA repair, chromatin remodeling, histone modifications, and DNA replication (Table 1). Secondary validation of the SDL suppressors was done by independently generating double mutant strains of psh1\(\Delta\) GALCSE4 with candidate mutants. Growth assays were performed on media selective for the GALCSE4 plasmid and containing either glucose or raffinose and galactose. We used a $hir2\Delta psh1\Delta$ strain as a negative control because hir2\(\Delta\) psh1\(\Delta\) GALCSE4 strains display SDL (CIFTCI-YILMAZ et al. 2018). Of the thirtyeight strains tested, twenty-nine showed almost complete suppression, five strains showed a partial suppression, and four did not suppress the SDL on galactose media (Tables 1 and S1 and Figures S1A and S1B). We further tested a subset of the thirty-eight genes to confirm overexpression of CSE4 and found that strains with mutations in genes involved in RNA processing and transcription do not show galactose-induced expression of CSE4 (Table S1 and Figure S1C), indicating that these are false positive hits. Through secondary validation, we confirmed that 89% of the candidate mutants tested suppressed the psh1\(\triangle GALCSE4 \) SDL.

We initiated our studies with the INO80 chromatin remodeling complex as our screen identified deletion and mutant alleles corresponding to three components of the INO80 complex, Ies2, Arp8, and Act1 (POCH AND WINSOR 1997; SHEN *et al.* 2000; SHEN *et al.* 2003; Tosi *et al.* 2013). Secondary validation assays showed that *ies2∆* and *act1-132* do not suppress the SDL of

the $psh1\Delta$ GALCSE4 strain (Figures S1A and S1B). In contrast, $arp8\Delta$ did suppress the $psh1\Delta$ GALCSE4 SDL (Figures S1A and S2A) however, the $arp8\Delta$ strain displayed polyploidy when analyzed by Fluorescent Activated Cell Sorting (FACS) (Figure S2B) and we consequently did not pursue further studies with the INO80 complex.

Deletion of histone H4 alleles suppresses the SDL of a psh1\(\textit{D}\) GALCSE4 strain

Two nonallelic loci, HHT1/HHF1 and HHT2/HHF2, encode identical H3 and H4 proteins in budding yeast. The screen identified the deletion of either one of the histone H4 alleles, $HHT1/hhf1\Delta$ ($hhf1\Delta$) or $HHT2/hhf2\Delta$ ($hhf2\Delta$), as among the most prominent suppressors of the $psh1\Delta$ GALCSE4 SDL. A role for the dosage of histone H4-encoding genes in mislocalization of Cse4 has not yet been reported. We confirmed that the $hhf1\Delta$ and $hhf2\Delta$ strains do not exhibit defects in ploidy or cell cycle by FACS analysis (Figure S3). Growth assays confirmed that $psh1\Delta$ $hhf1\Delta$ GALCSE4 and $psh1\Delta$ $hhf2\Delta$ GALCSE4 strains plated on galactose media do not exhibit SDL (Figure 2A). We determined that the phenotype was linked to deletion of the H4 alleles because transformation of a plasmid with the respective wild type histone H4 gene into the $psh1\Delta$ $hhf1\Delta$ or $psh1\Delta$ $hhf2\Delta$ strains restored the SDL observed in the $psh1\Delta$ GALCSE4 strain (Figure 2B).

We next investigated if deletion of a single allele for either histone H3 or H2A genes could suppress the SDL of a $psh1\Delta$ GALCSE4 strain. Note that the two nonallelic loci, HTA1/HTB1 and HTA2/HTB2, encode almost identical H2A and H2B proteins. Deletion of HTA1 $(hta1\Delta/HTB1)$, HTA2 $(hta2\Delta/HTB2)$, HHT1 $(hht1\Delta/HHF1)$, or HHT2 $(hht2\Delta/HHF2)$ did not suppress the SDL of a $psh1\Delta$ GALCSE4 strain on galactose media (Figures 2C and 2D and Table 2). Based on these results we conclude that the suppression of $psh1\Delta$ GALCSE4 SDL is specific to the reduced gene dosage of H4.

Reduced gene dosage of H4 suppresses the SDL of $slx5\Delta$, $doa1\Delta$, $hir2\Delta$, cdc4-1, and cdc7-4 GALCSE4 strains

To determine if the SDL suppression by reduced H4 gene dosage is limited to the $psh1\Delta$ GALCSE4 strain, we deleted HHF1 or HHF2 in deletion or mutant strains encoding Slx5, Doa1, Hir2, Cdc4, and Cdc7 as deletion or mutation of these factors show SDL with GALCSE4 and mislocalization of transiently overexpressed Cse4 (Au et~al. 2013; OHKUNI et~al. 2016; CIFTCI-YILMAZ et~al. 2018; Au et~al. 2020; EISENSTATT et~al. 2020). Growth on galactose media revealed that the SDL of $doa1\Delta$, $slx5\Delta$, cdc4-1, and cdc7-4 GALCSE4 strains is suppressed when either HHF1 or HHF2 is deleted (Figures 2E and 2F and Table 2), while the SDL of $hir2\Delta$ GALCSE4 is suppressed only when HHF2 is deleted (Figure 2E and Table 2). These results suggest that the gene dosage of H4 contributes to the SDL of mutants that exhibit defects in Cse4 proteolysis and mislocalize Cse4 to non-centromeric regions.

Reduced gene dosage of H4 reduces the mislocalization of Cse4 in psh1\(\trace2\) strains

The SDL phenotype of $psh1\Delta$ GALCSE4 strains is correlated with the mislocalization of Cse4 to non-centromeric regions (HEWAWASAM et al. 2010; RANJITKAR et al. 2010). We examined if the suppression of SDL in the $psh1\Delta$ $hhf1\Delta$ GALCSE4 or $psh1\Delta$ $hhf2\Delta$ GALCSE4 strains is due to reduced mislocalization of Cse4. We performed ChIP-qPCR to assay the localization of Cse4 using chromatin from wild type, $psh1\Delta$, $hhf1\Delta$, $hhf2\Delta$, $psh1\Delta$ $hhf1\Delta$, and $psh1\Delta$ $hhf2\Delta$ strains transiently overexpressing CSE4. In agreement with previously published data (HILDEBRAND AND BIGGINS 2016; HEWAWASAM et al. 2018; OHKUNI et al. 2020), we found that Cse4 enrichment at non-centromeric regions such as the promoters of RDS1, SLP1, GUP2, and COQ3 is higher in the $psh1\Delta$ strain compared to the wild type strain (Figures 3A and 3B and S4A and S4B). In contrast, deletion of HHF2 in a wild type strain or when combined with $psh1\Delta$ showed reduced levels of Cse4 enrichment at these regions (Figures 3A and 3B). Results for

ChIP-qPCR with the $hhf1\Delta$ strain also showed reduced levels of Cse4 at non-centromeric loci (Figure S4A and S4B). Consistent with previous studies (HILDEBRAND AND BIGGINS 2016), we observed higher levels of Cse4 at peri-centromeric regions in a $psh1\Delta$ strain (Figures 3C and S4C). However, we observed reduced levels of Cse4 at peri-centromeric regions in $psh1\Delta$ $hhf1\Delta$ and $psh1\Delta$ $hhf2\Delta$ strains when compared to the $psh1\Delta$ strain (Figures 3C and S4C). Localization of Cse4 to the centromere was not significantly altered in $hhf1\Delta$, $hhf2\Delta$, $psh1\Delta$ $hhf1\Delta$, and $psh1\Delta$ $hhf2\Delta$ strains (Figures 3C and S4C). Based on these results, we conclude that reduced gene dosage of H4 contributes to reduced levels of Cse4 at non-centromeric and peri-centromeric regions in $psh1\Delta$ strains.

Scm3 is the primary chaperone for centromeric deposition of Cse4 and strains depleted for Scm3 are not viable (CAMAHORT *et al.* 2007). However, overexpression of Cse4 can rescue the growth defect of Scm3-depleted cells, suggesting that non-Scm3-based mechanisms can promote centromeric deposition of overexpressed Cse4 (HEWAWASAM *et al.* 2018). Our studies so far have shown that reduced gene dosage of *H4* contributes to suppression of Cse4 mislocalization to non-centromeric regions. We next asked if the reduced gene dosage of *H4* would affect the Scm3-independent centromeric deposition of Cse4 by assaying the growth of Scm3-depleted cells that overexpress *CSE4*. In these strains, expression of Scm3 is regulated by a galactose-inducible promoter and is only expressed when grown in galactose medium, but not in glucose medium. However, overexpression of Cse4 from a copper-inducible promoter can suppress the growth defect caused by depletion of Scm3 on copper-containing medium (HEWAWASAM *et al.* 2018). We constructed *hhf2A GAL-SCM3 Cu-CSE4* strains and performed Western blot analysis to confirm the induced overexpression of Cse4 in these strains when grown in copper-containing medium (Figure 3D). Growth assays showed that deletion of *HHF2*

resulted in poor growth of cells when Cse4 is overexpressed in Scm3-depleted strains (Figure 3E, glucose + 0.5mM Cu). We conclude that physiological levels of histone H4 are required for centromeric association of Cse4 in cells depleted of Scm3 and for mislocalization of Cse4 to peri-centromeric and non-centromeric regions in $psh1\Delta$ strains.

Deletion of *HHF2* contributes to reduced stability of Cse4 in a *psh1* △ strain

The SDL phenotype and mislocalization of Cse4 in a $psh1\Delta$ GALCSE4 strain is associated with a higher stability of Cse4 (Hewawasam et al. 2010; Ranjitkar et al. 2010). The suppression of the $psh1\Delta$ GALCSE4 SDL and the reduced mislocalization of Cse4 by $hhf2\Delta$ led us to hypothesize that the stability of Cse4 would be reduced in a $psh1\Delta$ $hhf2\Delta$ strain. Protein stability assays showed that, in agreement with previous studies (Hewawasam et al. 2010; Ranjitkar et al. 2010), transiently overexpressed Cse4 is highly stable in the $psh1\Delta$ strain when compared to that observed in a wild type strain. The stability of Cse4 was not significantly affected in the $hhf2\Delta$ strain when compared to the wild type strain. Consistent with our hypothesis, we observed reduced stability of Cse4 in the $psh1\Delta$ $hhf2\Delta$ strain compared to the $psh1\Delta$ strain (Figure 4A). These results show a correlation between suppression of SDL of a $psh1\Delta$ GALCSE4 strain, lower levels of mislocalized Cse4 at non-centromeric regions, and reduced stability of Cse4 due to reduced gene dosage of H4.

Since defects in the ubiquitin-proteasome mediated proteolysis of Cse4 contribute to its mislocalization and increased stability (HEWAWASAM *et al.* 2010; RANJITKAR *et al.* 2010), we investigated if deletion of *HHF2* affects ubiquitination of Cse4 (Ub_n-Cse4) in a $psh1\Delta$ strain. Ubiquitin pull-down assays were done to determine the levels of Ub_n-Cse4 in wild type, $psh1\Delta$, hhf2, and $psh1\Delta$ $hhf2\Delta$ strains transiently overexpressing *CSE4*. Wild type strains expressing a non-tagged Cse4 or a mutant form of Cse4 (cse4^{16KR}) that cannot be ubiquitinated, where the 16

lysine residues are mutated to arginine, were used as negative controls. As previously reported (HEWAWASAM *et al.* 2010; RANJITKAR *et al.* 2010), levels of Ub_n-Cse4 were greatly reduced in the $psh1\Delta$ strain (38.2%±12.7) when compared to the wild type strain. The levels of Ub_n-Cse4 in the $psh1\Delta$ hhf2 Δ strain (31.7%±12.3) were similar to the $psh1\Delta$ strain, however Ub_n-Cse4 levels were decreased in the $hhf2\Delta$ strain (65.3%±23.9) compared to the levels in the wild type strain (Figure 4B). We propose that reduced mislocalization of Cse4 and ubiquitin-independent proteolysis of Cse4 contribute to reduced stability of Cse4 in a $psh1\Delta$ hhf2 Δ GALCSE4 strain.

Reduced dosage of *H4* is associated with defects in sumoylation of Cse4

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We recently reported that Cse4 is sumoylated and that the sumoylation status of Cse4 at residues K215/216 correlates with the SDL of $psh1\Delta$ GALCSE4 strains (OHKUNI et al. 2020). Overexpression of the sumoylation-defective cse4K215/216R/A does not cause SDL in $psh1\Delta$, $slx5\Delta$, or $hir2\Delta$ strains; the lack of an SDL phenotype in the $psh1\Delta$ strain is due to reduced mislocalization and lower protein stability of cse4K215/216R/A. The phenotypic consequences related to defects in Cse4 sumoylation are similar to the ones we have observed due to reduced dosage of H4. We examined if sumovlation of Cse4 is affected due to reduced dosage of H4. Wild type, hhf1∆, and hhf2∆ GALCSE4 strains were assayed for Cse4 sumoylation. Consistent with previous results (OHKUNI et al. 2016; OHKUNI et al. 2018; OHKUNI et al. 2020), we detected sumoylated Cse4 as a pattern of three high molecular weight bands in wild type cells overexpressing wild type Cse4 but not in wild type cells expressing vector alone or overexpressing cse4^{16KR} (Figure 5A). Deletion of either histone H4 allele resulted in reduced levels of sumoylated Cse4 (Figures 5A and 5B; p-value WT vs $hhf1\Delta = 0.0006$, p-value WT vs $hhf2\Delta = 0.0007$). To confirm that the reduction of sumoylated Cse4 is linked to deletion of the histone H4 allele, we assayed the levels of sumoylated Cse4 in hhf2\(\Delta\) GALCSE4 strains

transformed with an empty vector or with a plasmid borne *HHF2*. As expected, plasmid borne *HHF2* restored the levels of sumoylated Cse4 to that observed in wild type cells (Figures 5C and 5D). We conclude that physiological levels of histone H4 are required for Cse4 sumoylation.

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A histone H4 mutant defective for interaction with Cse4 suppresses the *psh1* \(GALCSE4 \) SDL and shows defects in Cse4 sumoylation

Our results so far have shown that reduced gene dosage of H4 contributes to the suppression of the SDL phenotype, reduced stability of Cse4, decreased mislocalization of Cse4 in psh1\(\triangle \) GALCSE4 strains, and defects in Cse4 sumoylation. We hypothesized that strains with defects in the interaction of H4 with Cse4 will display the same phenotypes that are observed due to reduced dosage of H4 in $psh1\Delta$ strains. To test our hypothesis, we used HHT1/hhf1 $hht2\Delta/hhf2\Delta$ strains with mutations either in the N-terminal lysines (HHT1/hhf1-10) or in the histone fold domain (HHT1/hhf1-20) (Figure 6A) that have been well characterized by genetic and biochemical analysis (SMITH et al. 1996; GLOWCZEWSKI et al. 2000). The temperature sensitivity of the HHT1/hhf1-20 strain, but not the HHT1/hhf1-10 strain, is suppressed by overexpression of Cse4 and the HHT1/hhf1-20 strain is proposed to have defects in the formation of the Cse4-H4 dimer (SMITH et al. 1996; GLOWCZEWSKI et al. 2000). We deleted PSH1 in the same genetic background as the HHT1/HHF1, HHT1/hhf1-10, and HHT1/hhf1-20 strains, transformed these strains with CSE4 on a galactose-inducible plasmid, and performed growth assays. Compared to wild type strains with a single copy of genes encoding histones H3/H4, HHT1/HHF1 psh1∆ strains display SDL on galactose medium when Cse4 is overexpressed, though to a less prominent degree compared to strains expressing both alleles encoding H3/H4 (compare Figure 6B to Figure 2A, psh1\(\triangle \) GALCSE4). The relative decrease in SDL may be due to the expression of a single copy of the genes encoding histones H3/H4 in the strain background. The HHT1/hhf1-20 mutant suppresses the SDL of psh1\(\triangle \) GALCSE4 strains while

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the HHT1/hhf1-10 mutant does not (Figure 6B). These findings suggest that the defect in the Cse4-H4 interaction contributes to the suppression of the psh1\(\Delta\) GALCSE4 SDL in the HHT1/hhf1-20 strain. We next examined the stability of Cse4 in HHT1/HHF1, HHT1/HHF1 psh1\(\Delta\), HHT1/hhf1-10, HHT1/hhf1-20, HHT1/hhf1-10 $psh1\Delta$, and HHT1/hhf1-20 $psh1\Delta$ strains transiently overexpressing CSE4. In agreement with previous findings (Figure 4A), overexpressed Cse4 is rapidly degraded in HHT1/HHF1 cells and is stabilized in the HHT1/HHF1 psh1∆ strain (HEWAWASAM et al. 2010; RANJITKAR et al. 2010) (Figure S5, top panels). Interestingly, degradation of overexpressed Cse4 in both HHT1/hhf1-10 and HHT1/hhf1-20 strains was faster compared to the HHT1/HHF1 strain. The HHT1/hhf1-20 psh1∆ strain showed rapid degradation of Cse4 when compared to the HHT1/HHF1 psh1∆ and HHT1/hhf1-10 $psh1\Delta$ strains (Figure S5). The rapid degradation of overexpressed Cse4 in the HHT1/hhf1-20 $psh1\Delta$ strain is consistent with previous studies for a correlation between higher protein stability and GALCSE4 SDL (HEWAWASAM et al. 2010; RANJITKAR et al. 2010; CIFTCI-YILMAZ et al. 2018; AU et al. 2020; EISENSTATT et al. 2020) and suggests that defective Cse4-H4 interaction contributes to the lack of *GALCSE4* SDL in $psh1\Delta$ strains. To examine the effect of the HHT1/hhf1-20 and HHT1/hhf1-10 alleles on the levels of Cse4 sumoylation, we used HHT1/HHF1, HHT1/hhf1-10, and HHT1/hhf1-20 strains overexpressing CSE4 to examine the sumoylation status of Cse4. Western blot analysis was performed after equal amounts of protein (5 mg) for each strain were pulled down with Ni-NTA agarose beads and normalized to the levels of non-modified Cse4 in the pull down (Figure 6C and D). Sumoylated Cse4 was observed in the HHT1/HHF1 and the HHT1/hhf1-10 strains (Figures 6C and 6D). Levels of sumoylated Cse4 were normalized to non-modified Cse4 in the

pull down samples. The low expression of Cse4 in the HHT1/hhf1-10 strain (Figure 6C, input) contributes to the higher levels of Cse4 sumoylation due to normalization to the low levels of non-modified Cse4 in this strain (Figure 6D). In contrast, the levels of Cse4 sumoylation were barely detectable in the HHT1/hhf1-20 strain when compared to the HHT1/HHF1 strain (Figures 6C and 6D). The reduced sumoylation of Cse4 in the HHT1/hhf1-20 strain is consistent with the rescue of SDL in the HHT1/hhf1-20 psh1 Δ GALCSE4 strain. We conclude that defects in the interaction of hhf1-20 with Cse4 contributes to reduced Cse4 sumoylation and suppression of psh1 Δ GALCSE4 SDL due to rapid degradation of Cse4.

Cse4 mutants defective in the Cse4-H4 interaction do not cause SDL in a $psh1\Delta$ strain and exhibit defects in Cse4 sumoylation

To further confirm that the Cse4-H4 interaction contributes to SDL in a $psh1\Delta$ GALCSE4 strain and Cse4 sumoylation, we investigated if Cse4 residues that are essential for the Cse4-H4 dimer formation (Figure 7A) affect the SDL of a $psh1\Delta$ strain and sumoylation of Cse4. Like the HHT1/hhf1-20 mutant, the cse4 mutants cse4-102 (L176S M218T) and cse4-111 (L194Q) exhibit defects in the Cse4-H4 dimer formation, while cse4-110 (L197S) likely impairs formation of the (Cse4-H4)₂ tetramer (GLOWCZEWSKI et~al.~2000). We hypothesized that overexpression of these cse4 mutants will not lead to SDL in a $psh1\Delta$ strain and these mutants will show defects in Cse4 sumoylation. To test these hypotheses, we generated galactose-inducible plasmids expressing cse4-102 (L176S M218T), $cse4-107^{MB}$ (L176S), cse4-108 (M218T), cse4-110 (L197S), and cse4-111 (L194Q). To test the effect of the cse4 mutants on SDL in a $psh1\Delta$ strain, we performed growth assays. We first determined that overexpression of mutant cse4 from these plasmids did not result in growth defects in a wild type strain (Figure S6). In agreement with our hypothesis, overexpression of all cse4 mutants did not cause SDL in a

 $psh1\Delta$ strain (Figure 7B). We conclude that the Cse4-H4 dimerization is essential for the SDL phenotype of a $psh1\Delta$ GALCSE4 strain.

Next, we generated galactose-inducible plasmids expressing cse4Y193A/F and cse4D217A/E for growth assays in a $psh1\Delta$ strain. The rationale for cse4Y193A/F is that Y193 is next to the mutated residue in cse4-111 (L194Q), is located at the center of the α 2 helix of Cse4, and interacts with the α 2 helix of H4 in the context of Scm3 (ZHOU et~al.~2011). For cse4D217A/E, the D217 residue is adjacent to the residue mutated in cse4-108 (M218T), is part of the K215/216 sumoylation consensus site, 214-MKKD-217 (Ψ -K-x-D/E), and is essential for dimerization of Cse4 (CAMAHORT et~al.~2009). Growth assays on galactose media showed that cse4Y193A and cse4D217A/E do not cause SDL in a $psh1\Delta$ strain (Figures 7C and 7D). Note that cse4Y193F showed partial lethality in a $psh1\Delta$ strain when compared to cse4 (Figure 7C). Taken together, these results show that overexpression of the cse4 mutants with defects in the formation of the Cse4-H4 dimer, do not lead to a SDL phenotype in a $psh1\Delta$ strain.

The lack of SDL in $psh1\Delta$ strains overexpressing cse4-102, cse4-107^{MB}, cse4-108, cse4-110, cse4-111, cse4Y193A, and cse4D217A/E is similar to the suppression of $psh1\Delta$ GALCSE4 SDL when combined with $hhf1\Delta$, $hhf2\Delta$, and hhf1-20 strains. Defects in Cse4 sumoylation in $hhf1\Delta$, $hhf2\Delta$, and hhf1-20 strains led us to hypothesize that cse4-102, cse4-107^{MB}, cse4-108, cse4-110, cse4-111, cse4Y193A, and cse4D217A/E strains will also show defects in Cse4 sumoylation. Thus, we examined the sumoylation status of the cse4 mutants used in the growth assays (Figure 7E). Consistent with our hypothesis, levels of Cse4 sumoylation were reduced in all cse4 mutants except cse4Y193F, which showed only a partial reduction of Cse4 sumoylation (Figures 7E and 7F). The reduced sumoylation of cse4Y193F is consistent with the partial lethality observed in a $psh1\Delta$ strain expressing cse4Y193F. Our results demonstrate that

overexpression of cse4 mutants defective for the Cse4-H4 dimer formation lead to defects in Cse4 sumoylation. We conclude that the Cse4-H4 dimer formation regulates Cse4 sumoylation and this contributes to $psh1\Delta$ GALCSE4 SDL.

493 DISCUSSION

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Mislocalization of overexpressed CENP-A and its homologs contributes to chromosomal instability (CIN) in yeast, fly, and human cells (HEUN et al. 2006; AU et al. 2008; MISHRA et al. 2011; LACOSTE et al. 2014; ATHWAL et al. 2015; SHRESTHA et al. 2017) and overexpression and mislocalization of CENP-A are observed in many cancers (TOMONAGA et al. 2003; AMATO et al. 2009; LI et al. 2011; McGovern et al. 2012; Sun et al. 2016; ZHANG et al. 2016). In this study, we performed the first genome-wide screen to identify deletion or temperature sensitive (ts) mutants that suppress the synthetic dosage lethality (SDL) due to mislocalization of overexpressed Cse4 in $psh1\Delta$ GALCSE4 strains. Deletion of either allele that encodes histone H4 (HHF1 and HHF2) were among the most prominent suppressors of $psh1\Delta$ GALCSE4 SDL. We determined that reduced gene dosage of H4 contributes to defects in Cse4 sumoylation and this prevents mislocalization of overexpressed Cse4 at peri-centromeric and non-centromeric regions, leading to suppression of the psh1∆ GALCSE4 SDL. We also determined that the Cse4-H4 interaction contributes to Cse4 sumoylation and psh1\(\triangle GALCSE4\) SDL as hhf1-20, cse4-102, and cse4-111 mutants, which are defective for the Cse4-H4 interaction, exhibit reduced sumoylation of Cse4 and do not exhibit psh1\(\triangle \) GALCSE4 SDL. Taken together, our genome-wide screen identified genes that contribute to Cse4 mislocalization and provides mechanistic insights into how reduced gene dosage of H4 prevents mislocalization of Cse4 into non-centromeric regions.

The suppressor screen was performed under a condition with high levels of Cse4 expression induced from a GAL1-GHis-GAL1-GHis-GAL1-GHis-GAL1-GHis-GAL1-GHis-GAL1-GII-GI

overexpression. These growth conditions limited us from identifying partial suppressors such as deletion of *NHP10*, which encodes a subunit of the INO80 chromatin remodeling complex and was previously shown to suppress the $psh1\Delta$ *GALCSE4* SDL on medium with a lower concentration of galactose (0.1%) (HILDEBRAND AND BIGGINS 2016). While our screen did not identify $nhp10\Delta$, it did identify two deletions and one mutant allele for genes encoding INO80 subunits, Ies2, Arp8, and Act1, respectively, that are evolutionarily conserved between yeast and human cells (POCH AND WINSOR 1997; SHEN *et al.* 2000; SHEN *et al.* 2003; Tosi *et al.* 2013). Secondary growth validation showed that $arp8\Delta$, but not act1-132 or $ies2\Delta$, suppresses the $psh1\Delta$ *GALCSE4* SDL. However, the polyploid nature of the $arp8\Delta$ strain precluded further study with this suppressor. The stringent growth conditions of the screen also prevented the identification of deletion of Cac2, a subunit of the CAF-1 complex, which promotes Cse4 incorporation at noncentromeric regions (HEWAWASAM *et al.* 2018). We determined that $cac2\Delta$ cannot suppress the $psh1\Delta$ *GALCSE4* SDL under the conditions used in our screen (data not shown).

Previous studies have shown that mislocalization of Cse4 to non-centromeric regions contributes to the *GALCSE4* SDL in *psh1* Δ , *slx5* Δ , *doa1* Δ , *hir2* Δ , *cdc4-1*, and *cdc7-4* strains (Hewawasam *et al.* 2010; Ranjitkar *et al.* 2010; Au *et al.* 2013; Ohkuni *et al.* 2016; Ciftci-Yilmaz *et al.* 2018; Au *et al.* 2020; Eisenstatt *et al.* 2020). We sought to define mechanisms that prevent lethality due to mislocalization of overexpressed Cse4. The identification of both *hhf1* Δ and *hhf2* Δ as suppressors of *psh1* Δ *GALCSE4* SDL led us to examine how reduced gene dosage of *H4* contributes to preventing mislocalization of Cse4. A role for histone H4 in centromeric localization of Cse4 has been examined previously (Deyter *et al.* 2017) however, the effect of gene dosage of *H4* in non-centromeric chromosome localization of Cse4 has not yet been explored. We determined that suppression of the *GALCSE4* SDL phenotype by *hhf1* Δ and

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 $hhf2\Delta$ is not restricted to $psh1\Delta$ strains and is also observed in $slx5\Delta$, $doa1\Delta$, cdc4-1, and cdc7-4 strains. The SDL phenotype of the $hir2\Delta$ GALCSE4 strain showed better suppression with $hhf2\Delta$ than with $hhf1\Delta$. This may be due to differential expression of H4 mRNA, which is five to seven times more abundant from the HHT2-HHF2 allele than from the HHT1-HHF1 allele (CROSS AND SMITH 1988) or due to the role of the HIR complex in histone gene expression (PROCHASSON et al. 2005; FILLINGHAM et al. 2009; KURAT et al. 2014).

We used several approaches to understand the molecular mechanism for suppression of the psh1 \triangle GALCSE4 SDL phenotype by hhf1 \triangle and hhf2 \triangle . These include ChIP-qPCR at regions of known Cse4 association, protein stability assays, and determining the status of Cse4 ubiquitination and sumoylation. Genome-wide studies have shown that overexpressed Cse4 is significantly enriched at promoters and peri-centromeric regions in a $psh1\Delta$ strain (HILDEBRAND AND BIGGINS 2016). Our ChIP-qPCR data showed reduced levels of Cse4 at peri-centromeric and non-centromeric regions in $psh1\Delta \ hhf1\Delta$ and $psh1\Delta \ hhf2\Delta$ strains when compared to the $psh1\Delta$ strain. The mislocalization of overexpressed Cse4 to non-centromeric regions contributes to highly stable Cse4 in $psh1\Delta$, $slx5\Delta$, $doa1\Delta$, $hir2\Delta$, cdc4-1, and cdc7-4 strains (HEWAWASAM et al. 2010; RANJITKAR et al. 2010; Au et al. 2013; OHKUNI et al. 2016; CIFTCI-YILMAZ et al. 2018; AU et al. 2020; EISENSTATT et al. 2020). We reasoned that reduced mislocalization of Cse4 to non-centromeric regions in $psh1\Delta hhf2\Delta$ strains may contribute to faster degradation of Cse4 in these strains. Our results showed that the proteolysis of Cse4 was indeed faster in $psh1\Delta hhf2\Delta$ strains when compared to the $psh1\Delta$ strain. Intriguingly, this was not due to increased ubiquitination of Cse4 (Ub_n-Cse4) in $psh1\Delta$ $hhf2\Delta$ strains. These results suggest a ubiquitinindependent mechanism that may contribute to the proteolysis of Cse4 in $hhf2\Delta psh1\Delta$ strains.

Ubiquitin-independent proteolysis has also been reported previously as cse4^{16KR}, in which all lysine residues are mutated to arginine, is still degraded (COLLINS *et al.* 2004).

Our results showing that reduced dosage of H4 contributes to the suppression of GALCSE4 SDL in $psh1\Delta$ strains, reduced mislocalization of Cse4, and lower protein stability of Cse4 are similar to the phenotypes of the sumoylation-defective cse4K215/216R/A strains (OHKUNI et~al.~2020). Consistent with these results, deletion of either histone H4 allele resulted in reduced levels of sumoylated Cse4. We therefore propose that physiological levels of H4 regulate the sumoylation of Cse4 and that this in turn facilitates mislocalization of overexpressed Cse4 to non-centromeric regions and GALCSE4 SDL in mutant such as $psh1\Delta$. Importantly, in contrast to histone H4, reduced dosage of genes encoding other canonical histones such as histones H2A or H3 does not suppress the $psh1\Delta$ GALCSE4 SDL.

To further examine the role of H4 in regulating the mislocalization of Cse4, we pursued studies using well characterized separation of function alleles of H4 (hhf1-20) and CSE4 (cse4-102 and cse4-111) with defects in the Cse4-H4 interaction (SMITH et~al.~1996; GLOWCZEWSKI et~al.~2000). Consistent with a role of H4 for its interaction with Cse4, we observed suppression of the $psh1\Delta$ GALCSE4 SDL in a hhf1-20 strain and lack of SDL when cse4-102 or cse4-111 were overexpressed in a $psh1\Delta$ GALCSE4 strain. The hhf1 mutant strains lack the HHT2/HHF2 allele and express only a single copy of H3/H4 (HHT1/HHF1). In this strain background, the $psh1\Delta$ GALCSE4 SDL was less severe compared to results in our strains with wild type copies of both HHT1/HHF1 and HHT2/HHF2 (Figure 2). Despite this, we were able to unambiguously establish that HHT1/hhf1-20, but not HHT1/hhf1-10, suppresses the $psh1\Delta$ GALCSE4 SDL. Interestingly, the hhf1-10 $psh1\Delta$ GALCSE4 strain displayed a more lethal phenotype than the wild type HHT1/HHF1 $psh1\Delta$ GALCSE4 strain. The N-terminal lysine residues on histone H4

(K5, 8, 12, 16) are acetylated and the *HHT1/hhf1-10* mutations mimic the hyperacetylated state of the lysine residues (K to Q). We have previously shown that levels of acetylated H4 are low at centromeres and that the maintenance of hypoacetylated H4 at the centromere is essential for kinetochore function and faithful chromosome segregation (CHOY *et al.* 2011). We propose that the hyperacetylated state of H4 in the *HHT1/hhf1-10* strain contributes to the more severe SDL that we observed. A recent study showed that strains with a mutation of histone H4 arginine 36 to alanine (H4R36A) display SDL when Cse4 is overexpressed and that this is due to defects in the interaction of H4R36A with Psh1, thereby leading to enrichment of Cse4 and Psh1 at noncentromeric regions in these cells (DEYTER *et al.* 2017).

Consistent with our previous studies (OHKUNI et al. 2020), we observed a correlation between the suppression of GALCSE4 SDL and reduced sumoylation of Cse4 in HHT1/hhf1-20, cse4-102, and cse4-111 strains. Similar results were observed with cse4Y193A/F, which is adjacent to the mutated site in cse4-111 (L194Q), and with cse4D217D/E, which is adjacent to the residue mutated in cse4-108 (M218T) and a part of the K215/216 sumoylation consensus site (CAMAHORT et al. 2009). Accordingly, low levels of sumoylated cse4Y193F correlate with a partial lethality of a psh1\(\Delta\) GALcse4Y193F strain and severe defects in sumoylated cse4Y193A correlate with a lack of SDL in a psh1\(\Delta\) GALcse4Y193A strain. Phenylalanine (F) is identical to tyrosine (Y) except for the hydroxyl group present on Y. It is possible that the structural similarity between Y and F allows at least partial formation of the Cse4-H4 dimer, resulting in partial sumoylation of cse4Y193F. In contrast, we observed a reduction of Cse4 sumoylation of both cse4D217A and cse4D217E mutants compared to wild type. The D217 residue of Cse4 is essential for growth and is important for the Cse4 dimerization. Since the cse4D217E mutant, which is part of the intact sumoylation consensus site, shows reduction of Cse4 sumoylation and

does not complement the null mutation (Figure S7), we propose that D217 has a role besides regulating sumoylation of Cse4K215/216. Sumoylation of Cse4 is not essential for centromeric localization of Cse4 because a $cse4^{16KR}$ strain with all 16 lysine (K) residues mutated to arginine (R) is viable in the context of the wild type centromeric chaperone Scm3 (AU *et al.* 2008). Sumoylation of Cse4K215/216 or physiological levels of H4 are indispensable only when Scm3 is not expressed (OHKUNI *et al.* 2020). Our results show that defects in Cse4 sumoylation contribute to reduced levels of non-centromere associated Cse4 with no significant effect on levels of centromere associated Cse4 in $psh1\Delta hhf1\Delta$ and $psh1\Delta hhf2\Delta$ strains. We propose that reduced dosage of H4 serves to protect the cells from the detrimental effects of overexpressed Cse4 due to defects in Psh1, SCF^{Cdc4}, Cdc7, Slx5/8, HIR, and Doa1-mediated proteolysis of Cse4. We define a previously undefined role for histone H4 gene dosage and the Cse4-H4 interaction as key upstream events for the sumoylation of Cse4, which facilitates non-centromeric localization of overexpressed Cse4 and SDL in a $psh1\Delta GALCSE4$ strain.

In summary, our genome-wide screen identified suppressors of $psh1\Delta$ GALCSE4 SDL with deletions of either allele that encodes histone H4 (HHF1 and HHF2) as among the most prominent suppressors. We present several experimental evidences to support our conclusions that reduced gene dosage of H4 contributes to defects in Cse4 sumoylation and reduced mislocalization of overexpressed Cse4 at peri-centromeric and non-centromeric regions, which in turn results in faster degradation of Cse4 and suppression of the $psh1\Delta$ GALCSE4 SDL. The suppression of SDL by $hhf1\Delta$ and $hhf2\Delta$ is not limited to a $psh1\Delta$ GALCSE4 background but is also observed in other mutants that exhibit GALCSE4 associated SDL. Most importantly, our results with the hhf1-20, cse4-102, and cse4-111 mutants, which are defective in the Cse4-H4 interaction, showed that the Cse4-H4 interaction is essential for non-centromeric association of

Cse4. These studies are important from a clinical standpoint given the poor prognosis of CENP-A overexpressing cancers (Tomonaga *et al.* 2003; Amato *et al.* 2009; Li *et al.* 2011; McGovern *et al.* 2012; Sun *et al.* 2016; Zhang *et al.* 2016). Future studies with histone H4 and other mutants identified in our screen will provide insights into mechanisms that promote mislocalization of overexpressed Cse4 and how defects in these mechanisms may safeguard the cell from the lethal effect due to mislocalization of overexpressed Cse4 in mutants such as $pshl\Delta$.

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FIGURE LEGENDS Figure 1. A genome-wide screen identified suppressors of the psh1\(\Delta\) GALCSE4 SDL. A. Schematic for the genome-wide screen. A psh1\(\triangle\) strain (YMB8995) transformed with GAL1-6His-3HA-CSE4 (pMB1458) was mated to an array of non-essential gene deletions and an array of conditional alleles of essential genes. Growth of the haploid meiotic progeny plated in quadruplicate was visually scored on glucose-and galactose-containing media grown at 30°C for non-essential and 26°C for essential gene mutant strains. Ninety-two genes were identified as growing better on galactose-containing media than the psh1\(\Delta\) GALCSE4 strain. Thirty-eight candidate genes were selected for confirmation of suppression of lethality. **B. Representative** plates from the genome-wide screen. Shown is Plate 01 of the non-essential gene deletion array. The mutant strains were spotted in quadruplicate on selective media plates containing glucose (top) or galactose (bottom). Red boxes (top box is $hap3\Delta$; bottom box is $hhf1\Delta$) highlight mutant strains that displayed improved growth on galactose-containing plates compared to the psh1\(\triangle \) GALCSE4 control strain (perimeter of plate) and did not show a growth defect or improved growth on the glucose plates. Figure 2. Deletion of H4 genes suppresses GALCSE4 SDL. Three independent isolates for each strain were assayed and shown is a representative for each. A. The psh1\(\Delta \) GALCSE4 SDL is suppressed by deletion of HHF1 or HHF2. Growth assays of wild type, $psh1\Delta$, $hhf1\Delta$, $hhf2\Delta$, $psh1\Delta$ $hhf1\Delta$, and $psh1\Delta$ $hhf2\Delta$ strains with empty vector (pMB433; YMB9802, YMB10478, YMB10825, YMB11166, YMB10821, and YMB10823, respectively) or GALI-6His-3HA-CSE4 (pMB1458; YMB9803, YMB10479, YMB10937, YMB10938, YMB10822, and YMB10824, respectively). Cells were spotted in five-fold serial dilutions on glucose (2% final concentration) or raffinose/galactose (2% final concentration each) media selective for the

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plasmid and grown at 30°C for three to five days. B. The psh14 GALCSE4 SDL suppression is **linked to the hhf1** Δ and hhf2 Δ alleles. Growth assays of psh1 Δ hhf1 Δ (YMB10822) and psh1 Δ hhf2∆ (YMB10824) strains with GAL1-6His-3HA-CSE4 (pMB1458) transformed with empty vector (pRS425) or a plasmid containing wild type HHF1 (pMB1928) or HHF2 (pMB1929). Strains were assayed as described above in (A). C. and D. Deletion of genes encoding histones H2A (C) or H3 (D) does not suppress the SDL of a psh1\(\textit{A}\) GALCSE4 strain. Growth assays of wild type, $psh1\Delta$, and (C) $hta1\Delta$, $hta2\Delta$, $psh1\Delta$ $hta1\Delta$, $psh1\Delta$ $hta2\Delta$, or (D) $hht1\Delta$, $hht2\Delta$, $psh1\Delta hht1\Delta$, and $psh1\Delta hht1\Delta$ strains with empty vector (pMB433; YMB9802, YMB10478, YMB11258, YMB11266, YMB11260, YMB11268, YMB11274, YMB11282, YMB11276, and YMB11284, respectively) or GAL1-6His-3HA-CSE4 (pMB1458: YMB9803, YMB10479, YMB11262, YMB11270, YMB11264, YMB11272, YMB11278, YMB11286, YMB11280, and YMB11288, respectively). Strains were assayed as described above in (A). E. Reduced gene dosage of H4 suppresses the SDL of slx5\(\Delta\), doa1\(\Delta\), and hir2\(\Delta\) GALCSE4 strains. Growth assays of wild type (YMB9804), $hhf1\Delta$ (YMB10937), $hhf2\Delta$ (YMB10938), $slx5\Delta$ (YMB10963), $slx5\Delta hhfl\Delta$ (YMB11046), $slx5\Delta hhf2\Delta$ (YMB11047), $doal\Delta$ (YMB11032), $doal\Delta hhfl\Delta$ (YMB11050), doa1\(\Delta\) hhf2\(\Delta\) (YMB11053), hir2\(\Delta\) (YMB8332), hir2\(\Delta\) hhf1\(\Delta\) (YMB11105), hir2\(\Delta\) hhf2∆ (YMB11107) strains expressing GAL1-6HIS-3HA-CSE4 (pMB1458). Strains were assayed as described above in (A) and grown at 30°C for three to five days. F. Deletion of HHF1 or HHF2 suppresses the SDL of cdc4-1 and cdc7-4 GALCSE4 strains. Growth assays of wild type (YMB9804), hhfl\(\Delta\) (YMB10937), hhf2\(\Delta\) (YMB10938), cdc4-1 (YMB9756), cdc4-1 $hhf1\Delta$ (YMB11051), cdc4-1 $hhf2\Delta$ (YMB11054), cdc7-4 (YMB9760), cdc7-4 $hhf1\Delta$ (YMB11052), and cdc7-4 hhf2 Δ (YMB11055) with GAL1-6His-3HA-CSE4 (pMB1458). Strains were assayed as described above in (A) and grown at 23°C for three to five days.

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Figure 3. Deletion of HHF2 reduces enrichment of Cse4 at peri-centromeric and noncentromeric regions. (A-C) ChIP-qPCR was performed on chromatin lysate from wild type (YMB9804), $psh1\Delta$ (YMB10479), $hhf2\Delta$ (YMB10938), and $psh1\Delta$ $hhf2\Delta$ (YMB10824) strains transiently overexpressing GAL1-6His-3HA-CSE4 (pMB1458). Enrichment of 6His-3HA-Cse4 is shown as a fold over wild type. Displayed are the mean of two independent experiments. Error bars represent standard deviation of the mean. **p-value < 0.0099, *p-value < 0.09, ns=not significant. A. and B. Levels of Cse4 enrichment at non-centromeric regions are reduced in a hhf2\(\text{strain}\). Enrichment of 6His-3HA-Cse4 at (A) RDS1, SLP1, COO3, GUP2, and (B) ACT1, SAP1, PHO5, FIG4, and UGA3. C. Levels of Cse4 at peri-centromeric regions, but not at the core centromere, are significantly reduced when HHF2 is deleted. Top: A diagram of the peri-centromere and centromere of Chromosome III analyzed by ChIP-qPCR. Horizontal lines represent the regions amplified. Bottom: Enrichment of 6His-3HA-Cse4 at the core centromere and at the left and right peri-centromeric regions on Chromosome III. D. Cse4 is expressed from a copper-inducible promoter in hhf2\(\Delta\) strains depleted of Scm3. Strains from (D) were grown to logarithmic phase in liquid media selective for the plasmid. Cells were induced with 0.5 mM copper for 2 hours and protein lysates were collected and analyzed by Western blot against Cse4 and Tub2 as a loading control. E: empty vector; C: copper inducible Cse4; -: no copper; +: 0.5 mM copper. E. Deletion of HHF2 reduces Cse4 deposition at the centromere in cells depleted of Scm3. Growth assays of strains in which Scm3 is expressed from a galactose inducible promoter and Cse4 is expressed from a copper-inducible promoter. Wild type and hhf2∆ with empty vector (pSB17; JG1589 and YMB11252, respectively) or a plasmid with copper inducible Cse4 (pSB873; JG1690 and YMB11254, respectively) were

plated in five-fold serial dilutions on media plates selective for the plasmid with raffinose/galactose (2% final concentration each) or glucose (2% final concentration) and with or without copper (0.5 mM final concentration). Plates were grown for three to five days at 30°C. Two independent transformants were tested and a representative image is shown.

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Figure 4. Deletion of HHF2 contributes to reduced stability and ubiquitin-independent proteolysis of Cse4 in a psh1 Δ strain. A. hhf2 Δ strains contribute to reduced stability of Cse4 in a psh1\(\text{strain}\). Western blot analysis of protein extracts from wild type (YMB9804), $psh1\Delta$ (YMB10479), $hhf2\Delta$ (YMB10938), and $psh1\Delta$ $hhf2\Delta$ (YMB10824) strains transiently overexpressing GAL1-6His-3HA-CSE4 (pMB1458). Cells were grown to logarithmic phase in media selective for the plasmid and containing raffinose (2% final concentration) and induced with galactose (2% final concentration) for 4 hours. Cultures were treated with cycloheximide (CHX, 10 µg/mL) and glucose (2%) and analyzed at the indicated time points. Extracts were analyzed by Western blot against HA (Cse4) and Tub2 as a loading control. Levels of 6His-3HA-Cse4 were normalized to Tub2 and the quantification of the percent remaining 6His-3HA-Cse4 after CHX treatment is shown in the graph. Error bars represent the SEM of two independent experiments. B. Deletion of HHF2 does not increase ubiquitination of Cse4 in a psh1\(\triangle \) strain. Ubiquitin-pull down assays were performed using protein extracts from wild type strains (BY4741) with no tag (pMB433) or overexpressing cse4^{16KR} (pMB1892) and from wild type (YMB9804), $psh1\Delta$ (YMB10479), $hhf2\Delta$ (YMB10938), and $psh1\Delta$ $hhf2\Delta$ (YMB10824) strains overexpressing 6His-3HA-CSE4 (pMB1458). Lysates were incubated with Tandem Ubiquitin Binding Entity beads (LifeSensors) prior to analysis of ubiquitin-enriched samples by Western blot against HA and input samples against HA and Tub2 as a loading control. Polyubiquitinated Cse4 (Ub_n-Cse4) is indicated by the bracket. HA levels in input samples were normalized to Tub2 levels and quantification of levels of Ub_n-Cse4 were normalized to the levels of Cse4 in the input. The percentage of Ub_n-Cse4 from two independent experiments with standard error is shown.

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Figure 5. Histone H4 contributes to the sumoylation of Cse4. A. Levels of sumoylated Cse4 are decreased in histone H4 deletion strains. Sumoylation levels were assayed on wild type (BY4741) strains transformed with empty vector (pYES2), pGAL-8His-HA-CSE4 (pMB1345), or pGAL-8His-HA-cse4 (pMB1344) and $hhf1\Delta$ (YMB10766) and $hhf2\Delta$ (YMB10767) strains transformed with pGAL-8His-HA-Cse4 (pMB1345). Sumoylated and nonmodified Cse4 were detected using cell lysates that were incubated with Ni-NTA beads followed by western blot analysis with antibodies against Smt3 and HA (Cse4), respectively. Arrows indicate the three high molecular weight bands that represent sumoylated Cse4. Asterisk indicates nonspecific sumoylated proteins that bind to beads. B. Quantification of relative levels of sumoylated Cse4 in histone H4 deletion strains. Levels of sumoylated Cse4 were normalized to non-modified Cse4 probed against HA in the pull-down sample. Statistical significance from two independent experiments was assessed by one-way ANOVA (p value = 0.0004) followed by Tukey post test (all pairwise comparisons of means). Error bars indicate average deviation from the mean. C. The Cse4 sumovlation defect in a $hhf2\Delta$ strain is linked to the HHF2 allele. Sumovlation levels were determined from lysates from a hhf2Δ (YMB10767) strain with pGAL-8His-HA-CSE4 (pMB1345) transformed with vector (pRS425) or HHF2 (pMB1929) as described in (A). Arrows indicate the three high molecular weight bands that represent sumoylated Cse4. Asterisk indicates nonspecific sumoylated proteins that bind to beads. D. Quantification of relative

levels of sumoylated Cse4. Relative levels of sumoylated Cse4 were normalized to non-modified Cse4 probed against HA in the pull-down sample. Error bars indicate average deviation from the mean from two biological replicates.

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Figure 6. Mutation in the histone fold domain of H4 histone suppresses the SDL phenotype of a psh1\(\triangle GALCSE4\) strain and causes defects in Cse4 sumoylation. A. Schematic of *HHF1*. Displayed is a cartoon of the *HHF1* gene with mutations in *hhf1-10* indicated by an 'x' and hhf1-20 with a '^' in the histone fold domain (HFD, blue). The specific residues mutated in each allele are indicated below the schematic. B. Mutations in the histone fold domain of histone H4 suppress the SDL phenotype of a psh1\(\textit{2}\) GALCSE4 strain. Growth assays of wild type (MSY559), psh1∆ (YMB11346), HHT1/hhf1-10 (MSY535), HHT1/hhf1-20 (MSY534), psh1∆ HHT1/hhf1-10 (YMB11347), and psh1∆ HHT1/hhf1-20 (YMB11348) with empty vector (pMB433) or expressing GAL1-6His-3HA-CSE4 (pMB1458). Cells were plated in five-fold serial dilutions on selective media plates containing either glucose (2% final concentration) or raffinose/galactose (2% final concentration each). Plates were incubated at 30°C for three to five days. Three independent transformants were tested and a representative image is shown. C. Mutations in the histone fold domain of histone H4 decrease levels of sumoylated Cse4. The levels of sumovlated Cse4 were determined using lysates from HHT1/HHF1 (MSY559), HHT1/hhf1-10 (MSY535), and HHT1/hhf1-20 (MSY534) strains in the $hht2\Delta/hhf2\Delta$ background, transformed with pGAL-8His-HA-CSE4 (pMB1345), as described in Figure 5A. Arrows indicate the three high molecular weight bands that represent sumoylated Cse4. Asterisk indicates nonspecific sumoylated proteins that bind to beads. D. Quantification of the relative levels of sumoylated Cse4 in hhf1 strains. Levels of sumoylated Cse4 were normalized to nonmodified Cse4 probed against HA in the pull-down samples and levels in the *HHT1/HHF1* strain were set to 1. Error bars indicate average deviation from the mean from two biological replicates.

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Figure 7. Cse4 mutants defective in the Cse4-H4 interaction do not cause SDL in a $psh1\Delta$ GALCSE4 strain and exhibit defects in Cse4 sumoylation. A. Schematic of CSE4. Displayed is a cartoon of the CSE4 gene highlighting mutations in the histone fold domain (HFD, red). The HFD is expanded under the representation of CSE4. Below the gene schematic is a key describing the symbol that represents a specific mutant cse4 allele and the residues mutated. **B.** Cse4-H4 assembly mutants in Cse4 do not cause SDL in a psh1\(\textit{D}\) GALCSE4 strain. Growth assays of a psh1\(\triangle \text{ (YMB8995)}\) strain transformed with pGAL1-8His-HA-Cse4 (pMB1344), pGAL1-8His-HA-cse4-102 (pMB1984), pGAL1-8His-HA-cse4-107^{MB} (pMB1985), pGAL1-8His-HA-cse4-108 (pMB1986), pGAL1-8His-HA-cse4-110 (pMB1987), or pGAL1-8His-HA-cse4-111 (pMB1988). Cells were plated in five-fold serial dilutions on selective media plates containing either glucose (2% final concentration) or raffinose/galactose (2% final concentration each). Plates were incubated at 30°C for three to five days. Three independent transformants were tested and a representative image is shown. C. The Y193A mutation in Cse4 does not cause SDL in a psh1\(\triangle \) GALCSE4 strain. Growth assays of a psh1\(\triangle \) (YMB9034) strain transformed with empty vector (pYES2), pGAL1-8His-HA-cse4^{Y193A} (pMB1766), or pGAL1-8His-HAcse4Y193F (pMB1787). Five-fold serial dilutions of the indicated strains were plated on glucose (2% final concentration)- or galactose (2% final concentration)-containing medium selective for the plasmid. The plates were incubated at 30°C for 3days. **D. The cse4D217A/E mutants do not** cause SDL in a psh1\(\Delta\) strain. Growth assays of a psh1\(\Delta\) (YMB9034) strain transformed with empty vector (pYES2), pGAL1-8His-HA-cse4D217A (pMB1910), or pGAL1-8His-HA-

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cse4D217E (pMB1920). Strains were assayed as described in (C). E. Cse4 sumovlation levels are decreased in Cse4-H4 assembly mutants. Levels of sumoylated Cse4 were assayed in a wild type strain (BY4741) transformed with empty vector (pYES2), pGAL1-8His-HA-CSE4 (pMB1345), pGAL1-8His-HA-cse4^{16KR} (pMB1344), pGAL1-8His-HA-cse4-107^{MB} (pMB1985), pGAL1-8His-HA-cse4-108 (pMB1986), pGAL1-8His-HA-cse4-102 (pMB1984), pGAL1-8His-HA-cse4-111 (pMB1988), pGAL1-8His-HA-cse4-110 (pMB1987), pGAL1-8His-HA-cse4Y193A (pMB1766), pGAL1-8His-HA-cse4Y193F (pMB1787), pGAL1-8His-HA-cse4D217A (pMB1910), or pGAL1-8His-HA-cse4D217E (pMB1920). Arrows indicate the three high molecular weight bands that represent sumoylated Cse4. Asterisk indicates nonspecific sumoylated proteins that bind to beads. F. Quantification of the relative levels of sumoylated Cse4 in cse4 mutants. Levels of sumoylated Cse4 in arbitrary density units were normalized to non-modified Cse4 probed against HA in the pull-down samples. Statistical significance from at least three biological repeats was assessed by one-way ANOVA (p-value < 0.0001) followed by Tukey post-test (all pairwise comparisons of means). Error bars indicate standard deviation from the mean.

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TABLE 1. Candidate double mutant strains with the indicated mutant allele combined with *psh1∆ GALCSE4* were generated and used for secondary validation using growth assays. Indicated is the allele analyzed, systematic name, gene name, standard name, visual scoring from the primary screen for growth score (from one to four) and colony score (from one to four), and suppression of SDL (Y: SDL was suppressed; N: SDL was not suppressed; Partial: SDL was partially suppressed).

Allele	Systematic Name	Gene Name	Standard Name	Growth Score	Colony Score	SDL Suppression
Non-essentia	ıl	•		•		
hhf1∆	YBR009C	HHF1	Histone H4	3	4	Y
hhf2∆	YNL030W	HHF2	Histone H4	3	4	Y
ies2∆	YNL215W	IES2	Ino Eighty Subunit	2	3	N
arp8∆	YOR141C	ARP8	Actin-Related Protein	3	4	Y
swc5∆	YBR231C	SWC5	SWr Complex	1	4	N
eaf1∆	YDR359C	EAF1	Esa1p-Associated Factor	2	3	Partial
eap1∆	YKL204W	EAP1	EIF4E-Associated Protein	2	4	Y
cse2∆	YNR010W	CSE2	Chromosome SEgregation	2	3	Partial
cse2∆_tsa	YNR010W	CSE2	Chromosome SEgregation	3	3	Y
mrm2∆	YGL136C	MRM2	Mitochondrial rRNA Methyl transferase	2	3	N
hap3∆	YBL021C	HAP3	Heme Activator Protein	3	4	Partial
hap5∆	YOR358W	HAP5	Heme Activator Protein	3	4	Y
rpl6b∆	YLR448W	RPL6B	Ribosomal Protein of the Large subunit	2	3	Y
rad4∆	YER162C	RAD4	RADiation sensitive	2	3	Partial
rad14∆	YMR201C	RAD14	RADiation sensitive	2	2	Y
Essential						
act1-132	YFL039C	ACT1	ACTin	3	4	N
mob1-5001	YIL106W	MOB1	Mps One Binder	4	4	Y
tbf1-5001	YPL128C	TBF1	TTAGGG repeat-Binding Factor	3	4	Y
csl4-5001	YNL232W	CSL4	Cep1 Synthetic Lethal	4	4	Y
pop4-5001	YBR257W	POP4	Processing Of Precursor RNAs	4	4	Y
orc1-5001	YML065W	ORC1	Origin Recognition Complex	4	4	Y
orc6-5001	YHR118C	ORC6	Origin Recognition Complex	4	4	Y
cft2-1	YLR115W	CFT2	Cleavage Factor Two	3	4	Partial
cft2-5001	YLR115W	CFT2	Cleavage Factor Two	4	4	Y
clp1-5001	YOR250C	CLP1	CLeavage/Polyadenylation factor Ia subunit	4	3	Y
ipa1-5001	YJR141W	IPA1	Important for cleavage and PolyAdenylation	3	4	Y
hrp1-1	YOL123W	HRP1	Heterogenous nuclear RibonucleoProtein	2	4	Y
rpb5-5001	YBR154C	RPB5	RNA Polymerase B	4	4	Y
rpc17-5001	YJL011C	RPC17	RNA Polymerase C	4	4	Y
pol31-5001	YJR006W	POL31	POLymerase	4	4	Y
srp54-5001	YPR088C	SRP54	Signal Recognition Particle 54-kD subunit	4	4	Y
dbp6-5001	YNR038W	DBP6	Dead Box Protein	4	4	Y

dbp9-5001	YLR276C	DBP9	Dead Box Protein	4	4	Y
yef3-f650s	YLR249W	YEF3	Yeast Elongation Factor	2	4	Y
cdc5-1	YMR001C	CDC5	Cell Division Cycle	2	4	Y
cdc31-1	YOR257W	CDC31	Cell Division Cycle	2	4	Y
hrr25-5001	YPL204W	HRR25	HO and Radiation Repair	3	4	Y
ost2-5001	YOR103C	OST2	OligoSaccharylTransferase	4	4	Y

TABLE 2. Summary of the SDL growth phenotypes of mutants that exhibit SDL with GALCSE4 and combined with $hhf1\Delta$ or $hhf2\Delta$. Shown is the protein function, relevant strain genotype, and growth with GALCSE4. Wild type growth is indicated as ++; SDL as --- and extent of suppression (++ or +++).

Protein Function	Relevant Strain Genotype	Growth with GALCSE4
	WT	++
Histone H4	hhf1∆	+++
	hhf2∆	+++
Histone H2A	hta1∆	++
	hta2∆	++
Histone H3	hht1∆	++
	hht2∆	++
E3 Ubiquitin Ligase	psh1∆	
	psh1∆ hhf1∆	+++
	psh1∆ hhf2∆	+++
	$psh1\Delta \ hhf1\Delta + HHF1$	
	$psh1\Delta hhf2\Delta + HHF2$	
	psh1∆ hta1∆	
	psh1∆ hta2∆	
	psh1∆ hht1∆	
	psh1∆ hht2∆	
SUMO-Targeted	slx5∆	
Ubiquitin Ligase	slx5∆ hhf1∆	+++
	slx5∆ hhf2∆	+++
Ubiquitin Binding	doa1∆	
	doa1∆ hhf1∆	+++
	doa1∆ hhf2∆	+++

HIR Nucleosome	hir2∆	
Binding Complex	hir2∆ hhf1∆	-
	hir2∆ hhf2∆	++
F-box of the SCF	cdc4-1	
Complex	cdc4-1 hhf1∆	+++
	cdc4-1 hhf2∆	+++
Dbf4-Dependent	cdc7-4	
Kinase	cdc7-4 hhf1∆	++
	cdc7-4 hhf2∆	++

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CITATIONS Allshire, R. C., and G. H. Karpen, 2008 Epigenetic regulation of centromeric chromatin: old dogs, new tricks? Nat Rev Genet 9: 923-937. Amato, A., T. Schillaci, L. Lentini and A. Di Leonardo, 2009 CENPA overexpression promotes genome instability in pRb-depleted human cells. Mol Cancer 8: 119. Athwal, R. K., M. P. Walkiewicz, S. Baek, S. Fu, M. Bui et al., 2015 CENP-A nucleosomes localize to transcription factor hotspots and subtelomeric sites in human cancer cells. Epigenetics Chromatin 8: 2. Au, W. C., M. J. Crisp, S. Z. DeLuca, O. J. Rando and M. A. Basrai, 2008 Altered dosage and mislocalization of histone H3 and Cse4p lead to chromosome loss in Saccharomyces cerevisiae. Genetics 179: 263-275. Au, W. C., A. R. Dawson, D. W. Rawson, S. B. Taylor, R. E. Baker et al., 2013 A Novel Role of the N-Terminus of Budding Yeast Histone H3 Variant Cse4 in Ubiquitin-Mediated Proteolysis. Genetics 194: 513-518. Au, W. C., T. Zhang, P. K. Mishra, J. R. Eisenstatt, R. L. Walker et al., 2020 Skp, Cullin, F-box (SCF)-Met30 and SCF-Cdc4-Mediated Proteolysis of CENP-A Prevents Mislocalization of CENP-A for Chromosomal Stability in Budding Yeast. PLoS Genet 16: e1008597. Biggins, S., 2013 The Composition, Functions, and Regulation of the Budding Yeast Kinetochore. Genetics 194: 817-846. Boltengagen, M., A. Huang, A. Boltengagen, L. Trixl, H. Lindner et al., 2016 A novel role for the histone acetyltransferase Hat1 in the CENP-A/CID assembly pathway in Drosophila

melanogaster. Nucleic Acids Res 44: 2145-2159.

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Burrack, L. S., and J. Berman, 2012 Flexibility of centromere and kinetochore structures. Trends Genet 28: 204-212. Camahort, R., B. Li, L. Florens, S. K. Swanson, M. P. Washburn et al., 2007 Scm3 is essential to recruit the histone H3 variant Cse4 to centromeres and to maintain a functional kinetochore. Mol Cell 26: 853-865. Camahort, R., M. Shivaraju, M. Mattingly, B. Li, S. Nakanishi et al., 2009 Cse4 is part of an octameric nucleosome in budding yeast. Mol Cell 35: 794-805. Chen, C. C., M. L. Dechassa, E. Bettini, M. B. Ledoux, C. Belisario et al., 2014 CAL1 is the Drosophila CENP-A assembly factor. J Cell Biol 204: 313-329. Cheng, H., X. Bao, X. Gan, S. Luo and H. Rao, 2017 Multiple E3s promote the degradation of histone H3 variant Cse4. Sci Rep 7: 8565. Cheng, H., X. Bao and H. Rao, 2016 The F-box Protein Rcy1 Is Involved in the Degradation of Histone H3 Variant Cse4 and Genome Maintenance. J Biol Chem 291: 10372-10377. Chereji, R. V., J. Ocampo and D. J. Clark, 2017 MNase-Sensitive Complexes in Yeast: Nucleosomes and Non-histone Barriers. Mol Cell 65: 565-577 e563. Choy, J. S., R. Acuna, W. C. Au and M. A. Basrai, 2011 A role for histone H4K16 hypoacetylation in Saccharomyces cerevisiae kinetochore function. Genetics 189: 11-21. Choy, J. S., P. K. Mishra, W. C. Au and M. A. Basrai, 2012 Insights into assembly and regulation of centromeric chromatin in Saccharomyces cerevisiae. Biochim Biophys Acta 1819**:** 776-783. Ciftci-Yilmaz, S., W. C. Au, P. K. Mishra, J. R. Eisenstatt, J. Chang et al., 2018 A Genome-Wide Screen Reveals a Role for the HIR Histone Chaperone Complex in Preventing Mislocalization of Budding Yeast CENP-A. Genetics 210: 203-218.

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Cole, H. A., J. Ocampo, J. R. Iben, R. V. Chereji and D. J. Clark, 2014 Heavy transcription of yeast genes correlates with differential loss of histone H2B relative to H4 and queued RNA polymerases. Nucleic Acids Res 42: 12512-12522. Collins, K. A., S. Furuyama and S. Biggins, 2004 Proteolysis contributes to the exclusive centromere localization of the yeast Cse4/CENP-A histone H3 variant. Curr Biol 14: 1968-1972. Costanzo, M., B. VanderSluis, E. N. Koch, A. Baryshnikova, C. Pons et al., 2016 A global genetic interaction network maps a wiring diagram of cellular function. Science 353. Cross, S. L., and M. M. Smith, 1988 Comparison of the structure and cell cycle expression of mRNAs encoded by two histone H3-H4 loci in Saccharomyces cerevisiae. Mol Cell Biol 8**:** 945-954. Deyter, G. M., and S. Biggins, 2014 The FACT complex interacts with the E3 ubiquitin ligase Psh1 to prevent ectopic localization of CENP-A. Genes Dev 28: 1815-1826. Deyter, G. M., E. M. Hildebrand, A. D. Barber and S. Biggins, 2017 Histone H4 Facilitates the Proteolysis of the Budding Yeast CENP-ACse4 Centromeric Histone Variant. Genetics 205: 113-124. Eisenstatt, J. R., L. Boeckmann, W. C. Au, V. Garcia, L. Bursch et al., 2020 Dbf4-Dependent Kinase (DDK)-Mediated Proteolysis of CENP-A Prevents Mislocalization of CENP-A in Saccharomyces cerevisiae. G3 (Bethesda). Fillingham, J., P. Kainth, J.-P. Lambert, H. van Bakel, K. Tsui et al., 2009 Two-color cell array screen reveals interdependent roles for histone chaperones and a chromatin boundary regulator in histone gene repression. Mol Cell 35: 340-351.

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Foltz, D. R., L. E. Jansen, A. O. Bailey, J. R. Yates, 3rd, E. A. Bassett et al., 2009 Centromerespecific assembly of CENP-a nucleosomes is mediated by HJURP. Cell 137: 472-484. Fujita, Y., T. Hayashi, T. Kiyomitsu, Y. Toyoda, A. Kokubu et al., 2007 Priming of centromere for CENP-A recruitment by human hMis18alpha, hMis18beta, and M18BP1. Dev Cell 12**:** 17-30. Glowczewski, L., P. Yang, T. Kalashnikova, M. S. Santisteban and M. M. Smith, 2000 Histonehistone interactions and centromere function. Mol Cell Biol 20: 5700-5711. Heun, P., S. Erhardt, M. D. Blower, S. Weiss, A. D. Skora et al., 2006 Mislocalization of the Drosophila centromere-specific histone CID promotes formation of functional ectopic kinetochores. Dev Cell 10: 303-315. Hewawasam, G., M. Shivaraju, M. Mattingly, S. Venkatesh, S. Martin-Brown et al., 2010 Psh1 is an E3 ubiquitin ligase that targets the centromeric histone variant Cse4. Mol Cell 40: 444-454. Hewawasam, G. S., K. Dhatchinamoorthy, M. Mattingly, C. Seidel and J. L. Gerton, 2018 Chromatin assembly factor-1 (CAF-1) chaperone regulates Cse4 deposition into chromatin in budding yeast. Nucleic Acids Res 46: 4440-4455. Hewawasam, G. S., M. Mattingly, S. Venkatesh, Y. Zhang, L. Florens et al., 2014 Phosphorylation by casein kinase 2 facilitates Psh1 protein-assisted degradation of Cse4 protein. J Biol Chem 289: 29297-29309. Hildebrand, E. M., and S. Biggins, 2016 Regulation of Budding Yeast CENP-A levels Prevents Misincorporation at Promoter Nucleosomes and Transcriptional Defects. PLoS Genet 12: e1005930.

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Kastenmayer, J. P., L. Ni, A. Chu, L. E. Kitchen, W. C. Au et al., 2006 Functional genomics of genes with small open reading frames (sORFs) in S. cerevisiae. Genome Res 16: 365-373. Kitagawa, K., and P. Hieter, 2001 Evolutionary concervation between budding yeast and human kinetochores. Nature Reviews Molecular Cellular Biology 2: 678-687. Kurat, C. F., J. Recht, E. Radovani, T. Durbic, B. Andrews et al., 2014 Regulation of histone gene transcription in yeast. Cell Mol LIfe Sci 71: 599-613. Lacoste, N., A. Woolfe, H. Tachiwana, A. V. Garea, T. Barth et al., 2014 Mislocalization of the centromeric histone variant CenH3/CENP-A in human cells depends on the chaperone DAXX. Mol Cell 53: 631-644. Li, Y., Z. Zhu, S. Zhang, D. Yu, H. Yu et al., 2011 ShRNA-targeted centromere protein A inhibits hepatocellular carcinoma growth. PLoS One 6: e17794. Maddox, P. S., K. D. Corbett and A. Desai, 2012 Structure, assembly and reading of centromeric chromatin. Curr Opin Genet Dev 22: 139-147. McGovern, S. L., Y. Qi, L. Pusztai, W. F. Symmans and T. A. Buchholz, 2012 Centromere protein-A, an essential centromere protein, is a prognostic marker for relapse in estrogen receptor-positive breast cancer. Breast Cancer Res 14: R72. McKinley, K. L., and I. M. Cheeseman, 2016 The molecular basis for centromere identity and function. Nat Rev Mol Cell Biol 17: 16-29. Mishra, P. K., W. C. Au, J. S. Choy, P. H. Kuich, R. E. Baker et al., 2011 Misregulation of Scm3p/HJURP causes chromosome instability in Saccharomyces cerevisiae and human cells. PLoS Genet 7: e1002303.

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Mizuguchi, G., H. Xiao, J. Wisniewski, M. M. Smith and C. Wu, 2007 Nonhistone Scm3 and histones CenH3-H4 assemble the core of centromere-specific nucleosomes. Cell 129: 1153-1164. Moreno-Moreno, O., M. Torras-Llort and F. Azorin, 2006 Proteolysis restricts localization of CID, the centromere-specific histone H3 variant of Drosophila, to centromeres. Nucleic Acids Res 34: 6247-6255. Ohkuni, K., R. Abdulle and K. Kitagawa, 2014 Degradation of centromeric histone H3 variant Cse4 requires the Fpr3 peptidyl-prolyl Cis-Trans isomerase. Genetics 196: 1041-1045. Ohkuni, K., R. Levy-Myers, J. Warren, W. C. Au, Y. Takahashi et al., 2018 N-terminal Sumoylation of Centromeric Histone H3 Variant Cse4 Regulates Its Proteolysis To Prevent Mislocalization to Non-centromeric Chromatin. G3 (Bethesda) 8: 1215-1223. Ohkuni, K., E. Suva, W. C. Au, R. L. Walker, R. Levy-Myers et al., 2020 Deposition of Centromeric Histone H3 Variant CENP-A/Cse4 into Chromatin Is Facilitated by Its C-Terminal Sumoylation. Genetics 214: 839-854. Ohkuni, K., Y. Takahashi and M. A. Basrai, 2015 Protein purification technique that allows detection of sumoylation and ubiquitination of budding yeast kinetochore proteins Ndc10 and Ndc80. J Vis Exp: e52482. Ohkuni, K., Y. Takahashi, A. Fulp, J. Lawrimore, W. C. Au et al., 2016 SUMO-Targeted Ubiquitin Ligase (STUbL) Slx5 regulates proteolysis of centromeric histone H3 variant Cse4 and prevents its mislocalization to euchromatin. Mol Biol Cell. Pidoux, A. L., E. S. Choi, J. K. Abbott, X. Liu, A. Kagansky et al., 2009 Fission yeast Scm3: A CENP-A receptor required for integrity of subkinetochore chromatin. Mol Cell 33: 299-311.

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Poch, O., and B. Winsor, 1997 Who's Who among the Saccharomyces cerevisiae Actin-Related Proteins? A Classification and Nomenclature Proposal for a Large Family. Yeast 13: 1053-1058. Prochasson, P., L. Florens, S. K. Swanson, M. P. Washburn and J. L. Workman, 2005 The HIR corepressor complex binds to nucleosomes generating a distinct protein/DNA complex resistant to remodeling by SWI/SNF. Genes Dev 19: 2534-2539. Ranjitkar, P., M. O. Press, X. Yi, R. Baker, M. J. MacCoss et al., 2010 An E3 ubiquitin ligase prevents ectopic localization of the centromeric histone H3 variant via the centromere targeting domain. Mol Cell 40: 455-464. Shen, X., G. Mizuguchi, A. Hamiche and C. Wu, 2000 A chromatin remodelling complex involved in transcription and DNA processing. Nature 406: 541-544. Shen, X., R. Ranallo, E. Choi and C. Wu, 2003 Inolvement of Actin-Related Proteins in ATP-Dependent Chromatin Remodeling. Mol Cell 12: 147-155. Shrestha, R. L., G. S. Ahn, M. I. Staples, K. M. Sathyan, T. S. Karpova et al., 2017 Mislocalization of centromeric histone H3 variant CENP-A contributes to chromosomal instability (CIN) in human cells. Oncotarget 8: 46781-46800. Shuaib, M., K. Ouararhni, S. Dimitrov and A. Hamiche, 2010 HJURP binds CENP-A via a highly conserved N-terminal domain and mediates its deposition at centromeres. Proc Natl Acad Sci U S A 107: 1349-1354. Smith, M. M., H. Yang, M. S. Santisteban, P. W. Boone, A. T. Goldstein et al., 1996 A Novel Histone H4 Mutant Defective in Nuclear Division and Mitotic Chromosome Transmission. Mol Cell Biol 16: 1017-1026.

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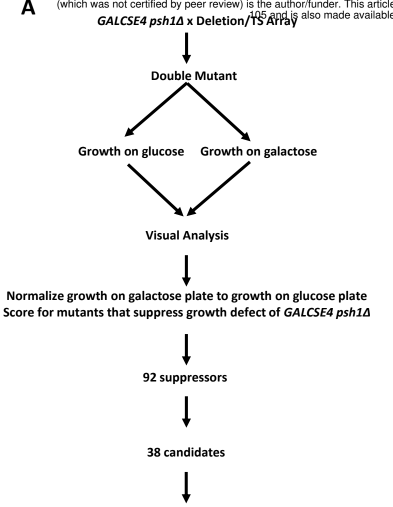
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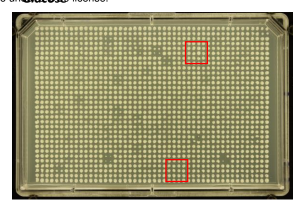
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1018

Stoler, S., K. Rogers, S. Weitze, L. Morey, M. Fitzgerald-Hayes et al., 2007 Scm3, an essential Saccharomyces cerevisiae centromere protein required for G2/M progression and Cse4 localization. Proc Natl Acad Sci U S A 104: 10571-10576. Sun, X., P. L. Clermont, W. Jiao, C. D. Helgason, P. W. Gout et al., 2016 Elevated expression of the centromere protein-A(CENP-A)-encoding gene as a prognostic and predictive biomarker in human cancers. Int J Cancer 139: 899-907. Tomonaga, T., K. Matsushita, S. Yamaguchi, T. Oohashi, H. Shimada et al., 2003 Overexpression and mistargeting of centromere protein-A in human primary colorectal cancer. Cancer Res 63: 3511-3516. Tosi, A., C. Haas, F. Herzog, A. Gilmozzi, O. Berninghausen et al., 2013 Structure and Subunit Topology of the INO80 Chromatin Remodeler and Its Nucleosome Complex. Cell 154: 1207-1219. Verdaasdonk, J. S., and K. Bloom, 2011 Centromeres: unique chromatin structures that drive chromosome segregation. Nat Rev Mol Cell Biol 12: 320-332. Williams, J. S., T. Hayashi, M. Yanagida and P. Russell, 2009 Fission yeast Scm3 mediates stable assembly of Cnp1/CENP-A into centromeric chromatin. Mol Cell 33: 287-298. Zhang, W., J. H. Mao, W. Zhu, A. K. Jain, K. Liu et al., 2016 Centromere and kinetochore gene misexpression predicts cancer patient survival and response to radiotherapy and chemotherapy. Nat Commun 7: 12619. Zhou, Z., H. Feng, B.-R. Zhuou, R. Ghirlando, K. Hu et al., 2011 Structural basis for recognition of centromere histone variant CenH3 by the chaperone Scm3. Nature 472: 234-237.



Confirm SDL phenotype



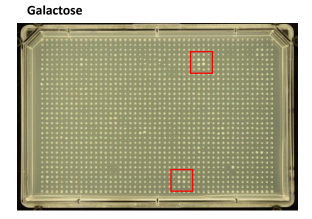
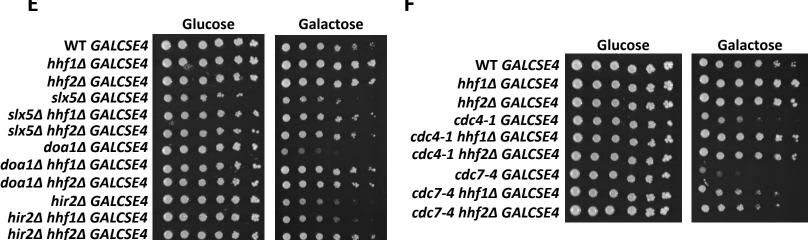


Figure 1



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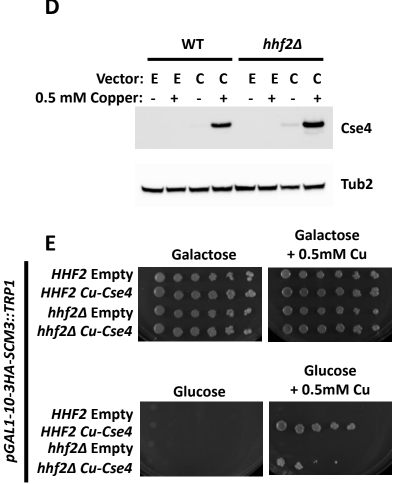


Figure 3

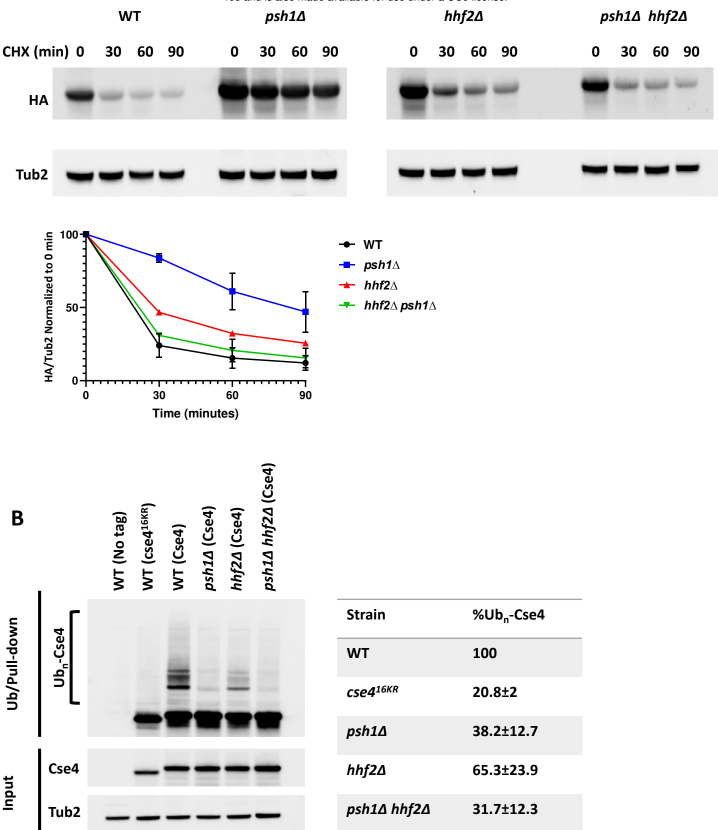
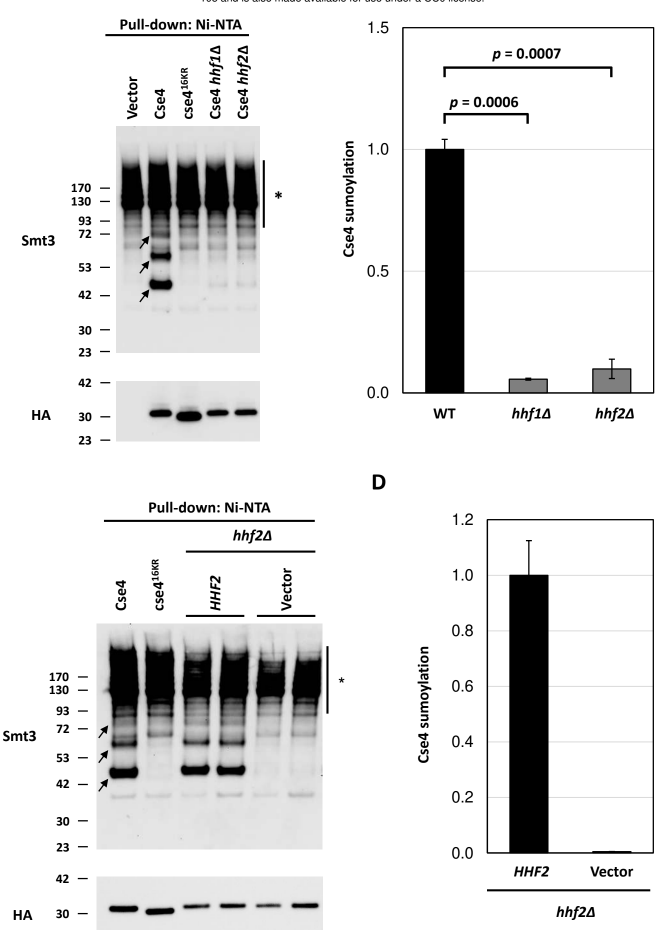


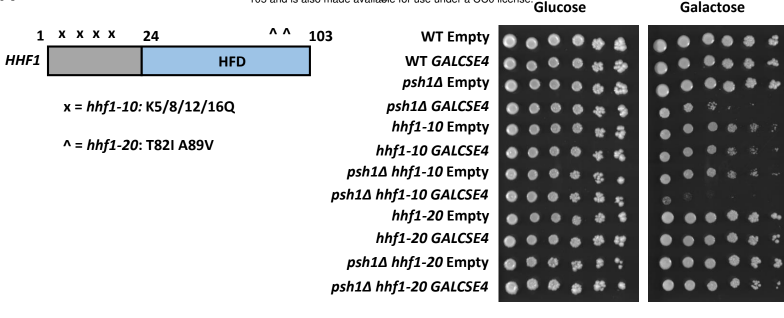
Figure 4



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Figure 5



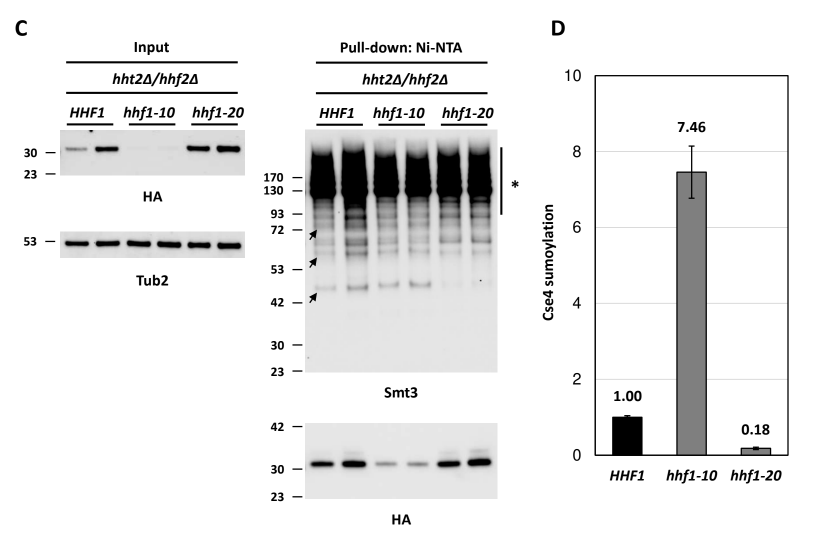


Figure 6

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Figure 7