Phosphorylation by Casein Kinase 2 Facilitates Psh1 Protein-assisted Degradation of Cse4 Protein*

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Background: Psh1 is an E3 ubiquitin ligase that controls CenH3/Cse4 levels through proteolysis in budding yeast. Here we report that Psh1 is phosphorylated by the Cka2 subunit of casein kinase 2 (CK2) to promote its E3 activity for Cse4. Deletion of CKA2 significantly stabilized Cse4. Consistent with phosphorylation promoting the activity of Psh1, Cse4 was stabilized in a Psh1 phosphodepleted mutant strain in which the major phosphorylation sites were changed to alanines. Phosphorylation of Psh1 did not control Psh1-Cse4 or Psh1-Ubc3(E2) interactions. Although Cse4 was highly stabilized in a cka2Δ strain, mislocalization of Cse4 was mild, suggesting that Cse4 misincorporation was prevented by the intact Psh1-Cse4 association. Supporting this idea, Psh1 was also stabilized in a cka2Δ strain. Collectively our data suggest that phosphorylation is crucial in Psh1-assisted control of Cse4 levels and that the Psh1-Cse4 association itself functions to prevent Cse4 misincorporation.

Results: Psh1 is phosphorylated in vivo by CK2, and its E3 ligase activity is promoted.

Conclusion: Phosphorylation is crucial in Psh1-assisted control of Cse4 levels, and the Psh1-Cse4 association itself functions to prevent Cse4 misincorporation.

Significance: This work reports a previously unknown function of CK2 in CenH3/Cse4 regulation.

Cse4 is the centromeric histone H3 variant in budding yeast. Psh1 is an E3 ubiquitin ligase that controls Cse4 levels through proteolysis. Here we report that Psh1 is phosphorylated by the Cka2 subunit of casein kinase 2 (CK2) to promote its E3 activity for Cse4. Deletion of CKA2 significantly stabilized Cse4. Consistent with phosphorylation promoting the activity of Psh1, Cse4 was stabilized in a Psh1 phosphodepleted mutant strain in which the major phosphorylation sites were changed to alanines. Phosphorylation of Psh1 did not control Psh1-Cse4 or Psh1-Ubc3(E2) interactions. Although Cse4 was highly stabilized in a cka2Δ strain, mislocalization of Cse4 was mild, suggesting that Cse4 misincorporation was prevented by the intact Psh1-Cse4 association. Supporting this idea, Psh1 was also stabilized in a cka2Δ strain. Collectively our data suggest that phosphorylation is crucial in Psh1-assisted control of Cse4 levels and that the Psh1-Cse4 association itself functions to prevent Cse4 misincorporation.

Faithful chromosome segregation during cell division is crucial for the equal distribution of genetic material. Centromeric chromatin specifies the site of assembly of the kinetochore, a massive, evolutionarily conserved protein complex that attaches chromosomes to microtubules for segregation. In contrast to the complex regional centromeres found in higher eukaryotic organisms, budding yeast has a simple point centromere, as found in Drosophila. CenH3, the CenH3 in budding yeast, is the centromeric histone H3 variant, collectively known as CenH3 and in budding yeast as Cse4, replaces histone H3 in centromeric nucleosomes. Exclusive incorporation of CenH3 (or Cse4 in budding yeast) at the centromere is the foundation for proper kinetochore assembly, microtubule attachment, and chromosome segregation. Therefore, regulating Cse4 levels and deposition is a critical process. An increased dosage of Cse4 leads to mislocalization, kinetochore defects, and chromosome loss in budding yeast (4, 5). High levels of the human CenH3, CENP-A, result in misincorporation at non-centromeric sites (6). Overexpression of CENP-A leads to chromosomal instability in pRb-depleted human cells (7) and also causes CENP-A mistargeting and aneuploidy in colorectal cancer cells (8). Therefore, levels of CenH3 need to be carefully regulated.

Several factors regulate CenH3 proteins. Ubiquitin-mediated proteolysis controls Cse4 protein levels in budding yeast (9) and centromere identifier (CID), the CenH3 in Drosophila (10). Psh1 is an E3 ubiquitin ligase for Cse4 in budding yeast (11, 12). Psh1 controls Cse4 levels via ubiquitylation and proteolysis, preventing mislocalization of Cse4. In Drosophila, a multi-subunit RING type E3 ubiquitin ligase, SCF⁵⁺⁺, regulates centromere identifier (CID) proteolysis (13). Scm3, a centromeric nucleosome assembly factor, protects Cse4 from degradation possibly through binding to the CENP-A targeting domain of Cse4. However, additional mechanisms may contribute to Cse4 stability and deposition. Post-translational modifications (PTMs) of the enzymes involved in the ubiquitin-proteasome pathway for Cse4 could play a role in Cse4 regulation.

Phosphorylation can regulate ubiquitylation-coupled proteolysis at several levels. Both E3 ubiquitin ligase activity and E2 ubiquitin conjugating activity can be regulated via phosphorylation (14–16). In this study, we report the first evidence that phosphorylation regulates Psh1-assisted degradation of Cse4. Psh1 was phosphorylated primarily by the Cka2 subunit of casein kinase 2 (CK2). Our results suggest that phosphorylation of Psh1 by CK2 promotes its E3 activity toward Cse4. Although deletion of CKA2 stabilized Cse4 more than deletion of PSH1, ectopic incorporation of Cse4 occurred at a lower level, suggesting that Psh1-Cse4 association itself prevents Cse4 misincorporation. Altogether, this study further advances our knowledge of how Cse4 is regulated in budding yeast.

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‡This article contains supplemental Table S1.

1The abbreviations used are: CEN, centromeric DNA; CID, centromeric histone H3 variant; CENP-A, centromere protein A; SCF, Skp1-Cdc53/Cullin-F box; PTM, post-translational modification; TAP, tandem affinity purification tag; IP, immunoprecipitation; CHX, cycloheximide; Ubᵦ, Cse4, polyubiquitylated Cse4; SD, synthetic defined.

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EXPERIMENTAL PROCEDURES

Yeast Strains—All yeast strains are listed in supplementary Table S1.

Psh1-TAP and Dsn1-FLAG Purifications—The Psh1-TAP purification method was adapted from Puig et al. (17). The protocol for kinetochore purification using Dsn1-FLAG was adopted from Akiyoshi et al. (18).

Whole Cell Extract Co-immunoprecipitation (Co-IP)—Cell lysates were prepared in lysis buffer (50 mM Tris (pH 7.5), 150 mM NaCl, 0.1% Nonidet P-40, 1 mM DTT, 10% glycerol, and protease inhibitors). Protein concentration was determined using the Bradford assay. Cell lysates were diluted with dilution/wash buffer (50 mM Tris (pH 7.5), 150 mM NaCl, and 0.1% Nonidet P-40) and incubated with the antibody overnight at 4 °C. Prewashed Protein G Dynabeads were added and incubated for 2 h at 4 °C. The beads were washed three times with dilution/wash buffer, and proteins were eluted with SDS buffer (10 mM Tris (pH 7.5), 1 mM EDTA, and 1% SDS). Immunoprecipitates were subjected to SDS-PAGE and Western blotting. Some co-IPs were performed using antibody-conjugated beads.

Proteasome Inhibition— Cultures grown to midlog phase in appropriate media were treated with MG132 (100 μM) or DMSO for 2–3 h. Cells were pelleted, washed with PBS, frozen in liquid N2, and stored in −80 °C.

Polyubiquitylated Protein Pulldown—Cell lysates were prepared in lysis buffer (same buffer used in co-IP with 15 mM N-ethylmaleimide), and protein concentration was measured using the Bradford assay. Polyubiquitin affinity resin (Thermo, 89899) (30 μl of a 25% slurry) was added to 2 mg of total proteins diluted 1:1 with TBS and incubated for 3 h or overnight at 4 °C. The beads were washed three times with wash buffer (lysis buffer: TBS, 1:9). Beads were boiled with 30 μl of SDS sample buffer, and samples were subjected to SDS-PAGE and Western blotting.

SDS-PAGE/Phos-tag-SDS-PAGE and Western Blotting—SDS-PAGE was performed using 4–12% precast gels (Invitrogen) or homemade 6% gels. For Phos-tag-SDS-PAGE, gels were prepared including acrylamide-pendant Phos-tag ligand (AAL-107). After running the gels, proteins were transferred onto PVDF membrane (Millipore, Immobilon-P), and Western blotting was performed according to a standard protocol.

Antibodies/Conjugated Beads—The antibodies used are as follows: anti-Myc (Covance, MMS150P), anti-Myc-HRP (Roche Applied Science, 11814150001), anti-HA (Covance, PRB101P, Roche Applied Science, 11867430001), anti-HA-HRP (Roche Applied Science, 12013819001), anti-ubiquitin (Covance, MMS257P), anti-Cse4 (polyclonal rabbit antibody against recombinant Cse4), anti-Pgk1 (Invitrogen, 459250), anti-FLAG (Sigma, F3165), anti-TAP (Open Biosystems, CAB1001), and anti-FLAG beads (Sigma, F2426).

Protein Stability Assay—Cells were grown to midlog phase in appropriate media. If Gal induction was needed, galactose was added to a final concentration of 4%, and cells were incubated for another 2 h. Cycloheximide (Sigma) was added to a final concentration of 10 μg/ml, and culture fractions were harvested at different time points. Cell lysates were prepared in lysis buffer, and protein concentration was determined using the Bradford assay. Proteins were analyzed by SDS-PAGE and Western blotting using the same amount of total proteins from each time point.

In Vitro Kinase Assay— Purified FLAG-His6-Psh1 (1 μg) was mixed with CK2 (1000 units) and 200 μM γ-labeled ATP (600 μCi/μmol) in a 50-μl final volume of CK2 reaction buffer (20 mM Tris-HCl, 50 mM KCl, and 10 mM MgCl2 (pH 7.5)). The reaction mixture was incubated at 30 °C for 2.5 h and boiled with SDS sample buffer. Proteins were separated using a 4–12% gel, and the gel was exposed to autoradiography film.

ChIP-Quantitative PCR— Cultures grown to an A600 of ~0.7 were fixed in 1% formaldehyde for 15 min and then quenched with glycine (final concentration, 0.125 M) for 5 min. Cells were lysed for 1 h at 4 °C in FA-Lysis SDS buffer (50 mM Hapes (pH 7.5), 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 0.2% SDS, and protease inhibitors). The lysate was sonicated using a Biorupter for 20 min (30 s on/off) at medium intensity. Protein concentration was determined, and samples were normalized before proceeding with the ChIP. 20% of total chromatin extract was used for IP using 2 μl of anti-Cse4 antibody and incubated at 4 °C overnight. 25 μl of Protein G Dynabeads (Invitrogen, 100-04D) prewashed in FA-Lysis buffer (50 mM Hapes (pH 7.5), 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate) were added and incubated at 4 °C for 3 h. Beads were washed at room temperature with the following sequence of buffers: FA-Lysis buffer, FA-Lysis buffer with 1 mM NaCl, FA-Lysis buffer with 500 mM NaCl, TEL buffer (0.25 mM LiCl, 10 mM Tris-HCl (pH 8.0), 1 mM EDTA, 1% Nonidet P-40, and 1% sodium deoxycholate), and twice with 1× Tris-EDTA. Chromatin fragments were eluted twice using 200 μl of Elution Buffer (1% SDS, 250 mM NaCl, and 1× Tris-EDTA) at 65 °C with agitation. Two elutions were combined, treated with Proteinase K for 1 h at 55 °C, and incubated overnight at 65 °C. DNA was extracted with phenol-chloroform and precipitated using 100% ethanol. 2% of the total chromatin extract was processed for the input sample. Cse4 ChIPs were performed in triplicate with two no-antibody controls for each strain. Quantitative PCR was performed in triplicate for each sample using a Quanta Biosciences Perfecta SYBR Green FastMix.

Recombinant Protein Expression and Purification—The method for FLAG-His6-Psh1 expression and purification has been published (11). Recombinant CK2 was purchased from New England Biolabs (catalog number P6010L).

RESULTS

Psh1 Is Phosphorylated in Vivo—How Psh1 E3 ligase activity is controlled is unknown, but we suspected that PTMs played a role. To investigate whether PTMs could regulate Psh1 activity, we affinity-purified Psh1-TAP and analyzed samples using mass spectrometry (MS) for PTMs. We were able to identify multiple phosphorylation sites on Psh1 (Fig. 1A). The Psh1-TAP purification is enriched mainly for the soluble pool of Psh1. To specifically examine centromere-associated Psh1, we purified kinetochore proteins using Dsn1-FLAG, a subunit of the kinetochore MIND complex (18), and analyzed PTMs of the Psh1 in these samples by MS. We identified the same phosphorylation sites in centromere-associated Psh1 as found in Psh1-TAP
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samples as well as two additional sites (Fig. 1A). Major phosphorylation sites with >20% of modified/total spectra are indicated in Fig. 1B relative to the major protein domains in Psh1. To further confirm the phosphorylation of Psh1, we used an epitope-tagged Psh1-HA strain (Fig. 1C). Psh1-HA was immunoprecipitated from whole cell extracts, and eluates were subjected to shrimp alkaline phosphatase treatment to remove phosphate groups from phosphorylated proteins. Samples were analyzed by Phos-tag-SDS-PAGE and Western blotting with anti-HA antibodies. On a Phos-tag gel, phosphorylated counterparts. We detected a faster migrating band indicated by an asterisk could be a dimer of Psh1. F, the Cka2 catalytic subunit of CK2 is necessary for phosphorylation of Psh1. Strains expressing Psh1-HA were constructed bearing deletions in CKA1 or CKA2. After Phos-tag-SDS-PAGE (7.5% gel with 10 μM Phos-tag) of cell lysates, Western blotting was carried out using anti-HA antibodies. The differences in band intensities in C and F could be due to the inefficient transfer of phosphorylated proteins from the Phos-tag gel to the membrane due to high affinity toward the Phos-tag. G, the Cka2 regulatory subunits Ckb1 and Ckb2 pull down Psh1. Affinity-tagged strains (Psh1-HA/Ckb1-FLAG and Psh1-HA/Ckb2-FLAG) were used in co-IP. A strain lacking the tag on Ckb1 and Ckb2 was used as the control. In anti-FLAG co-IP using cell lysates, both Ckb1 and Ckb2 pull down Psh1. Ckb1 shows a much stronger interaction with Psh1 than Psh1-HA as revealed by a more intense band. H, Ckb1 binds to Psh1 via its RING finger domain. Strains were constructed expressing the affinity-tagged proteins Psh1-HA and Ckb1-FLAG. A co-IP using anti-FLAG antibody from whole cell lysates shows that the RING finger domain of Psh1 is necessary for Ckb1 interaction. A control IP was performed from a strain expressing Psh1-HA but lacking a FLAG tag on Ckb1. P-Psh1, phosphorylated Psh1.

FIGURE 1. Psh1 is phosphorylated in vivo by CK2. A, PTM analysis using MS5 after affinity purification of Psh1-TAP and Dsn1-FLAG reveals phosphorylation sites of Psh1. The combined percentages of modified spectra/total spectra of the two purifications are graphed. The number of modified peptides/total peptides detected in Psh1-TAP (*) and Dsn1-FLAG (Δ) samples are indicated above each bar. CK2 sites predicted by NetPhosK are in black. B, the major phosphorylation sites of Psh1 are shown relative to the three defined domains of a tripartite motif protein: RING finger motif, Cys4 zinc finger motif, and an acidic stretch. Lys-303 is the autoubiquitylation site of Psh1 identified by in vitro ubiquitylation and MS. C, Psh1-HA was immunoprecipitated from cell lysates using anti-HA, and eluates were subjected to shrimp alkaline phosphatase (SAP) treatment. Samples were analyzed by Phos-tag-SDS-PAGE (12% gel with 10 μM Phos-tag) and Western blotting with anti-HA antibodies. D, Psh1-TAP purification brings down all four subunits of CK2 as detected by MS. The normalized spectral abundance factor (NSAF) shows the estimated relative protein abundance calculated as described previously (38). E, Psh1 can be phosphorylated in vitro using recombinant CK2. Recombinant proteins and γ-labeled ATP were used to carry out a kinase assay in vitro. An autoradiograph following SDS-PAGE is shown. The band indicated by * can be phosphorylated. F, the Cka2 catalytic subunit of CK2 is necessary for phosphorylation of Psh1. Strains expressing Psh1-HA were constructed bearing deletions in CKA1 or CKA2. After Phos-tag-SDS-PAGE (7.5% gel with 10 μM Phos-tag) of cell lysates, Western blotting was carried out using anti-HA antibodies. The differences in band intensities in C and F could be due to the inefficient transfer of phosphorylated proteins from the Phos-tag gel to the membrane due to high affinity toward the Phos-tag. G, the Cka2 regulatory subunits Ckb1 and Ckb2 pull down Psh1. Affinity-tagged strains (Psh1-HA/Ckb1-FLAG and Psh1-HA/Ckb2-FLAG) were used in co-IP. A strain lacking the tag on Ckb1 and Ckb2 was used as the control. In anti-FLAG co-IP using cell lysates, both Ckb1 and Ckb2 pull down Psh1. Ckb1 shows a much stronger interaction with Psh1 than Psh1-HA as revealed by a more intense band. H, Ckb1 binds to Psh1 via its RING finger domain. Strains were constructed expressing the affinity-tagged proteins Psh1-HA and Ckb1-FLAG. A co-IP using anti-FLAG antibody from whole cell lysates shows that the RING finger domain of Psh1 is necessary for Ckb1 interaction. A control IP was performed from a strain expressing Psh1-HA but lacking a FLAG tag on Ckb1. P-Psh1, phosphorylated Psh1.

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Psh1-HA band in the shrimp alkaline phosphatase-treated sample compared with the untreated sample, further confirming in vivo phosphorylation of Psh1.

CK2 Is a Kinase for Psh1—CK2 is a highly conserved serine/threonine kinase in eukaryotic organisms. This versatile kinase controls many important cellular processes such as signal transduction, transcription, translation, metabolism, and cell cycle progression. Hundreds of CK2 substrates have been identified so far (19). CK2 is a heterotetrameric enzyme composed of two catalytic and two regulatory subunits. In budding yeast, the catalytic subunits are Cka1 and Cka2, and the regulatory subunits are Ckb1 and Ckb2. Most of the Psh1 phosphorylation sites identified by MS are predicted to be CK2 sites by NetPhosK, a program for prediction of potential kinases. These potential CK2 sites are depicted in black bars in Fig. 1A. In addition, all four subunits of CK2 co-purified with Psh1-TAP (Fig. 1D), further suggesting CK2 as a potential kinase for Psh1.

To determine whether CK2 is a kinase for Psh1, we performed a kinase assay in vitro utilizing recombinant Psh1, CK2, and radiolabeled ATP (Fig. 1E). The autoradiograph clearly shows phosphorylated Psh1 in the presence of both CK2 and ATP, suggesting that Psh1 is a substrate for CK2. Recombinant Psh1 used in this assay was purified using a baculovirus expression system in insect cells. The lower intensity phosphorylated Psh1 band observed with the control sample without CK2 (Fig. 1E, lane 4) is possibly due to some kinase activity that co-purified with Psh1.

To identify which subunit of CK2 is responsible for phosphorylating Psh1, we deleted each catalytic subunit (ckaΔ or cka2Δ) in the strain expressing Psh1-HA. Single deletions were made because double deletion of the two catalytic subunits is reported to be lethal (20). Cell lysates of the mutants were compared with WT using Phos-tag-SDS-PAGE and Western blotting with anti-HA antibodies (Fig. 1F). Psh1-HA in the cka1Δ strain migrated similarly to WT, whereas in the cka2Δ strain it showed a faster migrating, lesser or non-phosphorylated band, suggesting that Cka2 is primarily responsible for Psh1 phosphorylation.

To identify the subunits of CK2 that are physically interacting with Psh1, epitope-tagged strains with Psh1-HA/Ckb1-FLAG and Psh1-HA/Ckb2-FLAG were used to perform co-IP and Western blotting (Fig. 1G). We were able to detect both Ckb1 and Ckb2 interacting with Psh1. However, the interaction between Psh1 and Ckb1 was much stronger. These data demonstrate that CK2 is a kinase for Psh1 with catalysis and interaction primarily mediated by Cka2 and Ckb1, respectively.

Psh1 contains a RING domain, a type of zinc finger that can bind two zinc cations, and a zinc finger domain (Fig. 1B). The RING domain is essential for the Psh1-Cse4 interaction, and a mutation in the RING domain stabilizes Cse4 (11). We tested the RING and zinc finger mutants for interaction with Ckb1 of CK2 and Ubc3/Cdc34, the E2 ubiquitin-conjugating enzyme for Psh1 (see comment in Hewawasam et al. (11)). Co-IP showed that the Ckb1-Psh1 interaction required the RING finger domain of Psh1 (Fig. 1H). The RING domain appears to be required for Psh1 to interact with both Cse4 and Ckb1. However, neither the RING nor zinc finger domain was required for the Psh1-Ubc3 interaction (data not shown). This suggests that the interaction between Psh1 and Ubc3 is facilitated by an unidentified region of Psh1.

CK2 Is Important for Efficient Regulation of Cellular Cse4 Levels—PSh1 is not an essential gene, and a simple deletion of PSH1 does not cause a defect in a minichromosome loss assay (11). Deletion of PSH1 resulted in slow growth when Cse4 was overexpressed from the GAL promoter on a 2-μm plasmid. Cse4 overexpression in a psh1Δ strain alters kinetochore function as revealed by a delay in destruction of anaphase inhibitor Pds1 (12). If CK2 is important for Psh1-mediated Cse4 regulation, then overexpression of Cse4 in a CK2 mutant might be expected to show a growth phenotype similar to PSH1 deletion, and Cse4 protein levels should be subsequently stabilized. To test this, we overexpressed Cse4 in catalytic subunit deletion mutants cka1Δ and cka2Δ (Fig. 2A). We observed slow growth with cka2Δ but not with cka1Δ (Fig. 2A, compare cka1Δ + CSE4 and cka2Δ + CSE4 with psh1Δ + CSE4), consistent with our observation that Cka2 was primarily responsible for phosphorylation of Psh1. Cse4 overexpression in the double mutant psh1Δ/cka2Δ showed a growth phenotype similar to that of either single mutant, suggesting the Psh1 and Cka2 function in the same pathway to control Cse4 levels.

We then examined the Cse4 protein levels in these strains using a protein stability assay (Fig. 2B). After Gal induction of Cse4 overexpression, cycloheximide (CHX) was added to inhibit protein translation, and the level of Cse4 in cell lysates was determined at different time points using Western blotting with anti-Cse4 antibody. We observed highly stabilized Cse4 levels in a ckaΔ mutant compared with a WT strain, and the half-life of Cse4 (Cse4 t1/2) was about 3 times longer in a cka2Δ strain (Fig. 2B, left panel). Cse4 was more stabilized by cka2Δ than psh1Δ. We next measured the stability of endogenous levels of Cse4-Myc (Fig. 2C). Cse4-Myc was also stabilized in a cka2Δ strain compared with a WT strain. These results suggest that phosphorylation of Psh1 by CK2 may promote its E3 activity for Cse4. Because the two stability assays in Fig. 2B were performed separately, the difference in Cse4 stability in WT strain between the left and right panels could be due to fluctuations in the experimental conditions. However, the two stability assays with Cse4 overexpression resulted in an average Cse4 t1/2 of 12 min for WT strain, which is comparable with a Cse4 t1/2 of WT strain (12.7 min) with endogenous Cse4 levels in Fig. 2C. Conversely, in cka2Δ strain, Cse4 t1/2 was elevated with Cse4 overexpression (Fig. 2, compare B, left panel, with C; 29.0 versus 18.2 min, respectively). Assuming that the experimental conditions are comparable in Fig. 2, B, left panel, and C, a high level of soluble Cse4 and inefficient Cse4 degradation in the absence of Cka2 may elevate Cse4 t1/2 in the cka2Δ strain with Cse4 overexpression.

We next examined ubiquitlated forms of Cse4 in WT and cka2Δ strains with both endogenous Cse4 and overexpression (Fig. 3). Cells expressing Cse4-Myc at endogenous levels were grown to midlog phase and treated with MG132 for 2 h to inhibit proteasome function. Pdr5, a plasma membrane transporter, is deleted from these strains to improve MG132 uptake. The efficiency of proteasome inhibition was confirmed by high accumulation of polyubiquitlated proteins in MG132-treated cells compared with DMSO control samples (Fig. 3A, left panel,
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A

|          | SD-His | SGal-His |
|----------|--------|----------|
| WT+EV    | ![Image](image1.png) | ![Image](image2.png) |
| WT+CSE4  | ![Image](image3.png) | ![Image](image4.png) |
| psh1Δ+EV | ![Image](image5.png) | ![Image](image6.png) |
| psh1Δ+CSE4 | ![Image](image7.png) | ![Image](image8.png) |
| cka1Δ+EV | ![Image](image9.png) | ![Image](image10.png) |
| cka1Δ+CSE4 | ![Image](image11.png) | ![Image](image12.png) |
| cka2Δ+EV | ![Image](image13.png) | ![Image](image14.png) |
| cka2Δ+CSE4 | ![Image](image15.png) | ![Image](image16.png) |
| psh1Δ+cka2Δ+EV | ![Image](image17.png) | ![Image](image18.png) |
| psh1Δ+cka2Δ+CSE4 | ![Image](image19.png) | ![Image](image20.png) |

B

|          | Before gal | After CHX |
|----------|------------|-----------|
|          | Before CHX | 15 min | 30 min | 45 min | 2.5 hrs |
|          | Before CHX | 15 min | 30 min | 45 min | 2.5 hrs |
|          | Before CHX | 15 min | 30 min | 45 min | 2.5 hrs |
|          | Before CHX | 15 min | 30 min | 45 min | 2.5 hrs |
|          | Before CHX | 15 min | 30 min | 45 min | 2.5 hrs |
|          | Before CHX | 15 min | 30 min | 45 min | 2.5 hrs |

C

|          | Before CHX | After CHX |
|----------|------------|-----------|
|          | Before gal | 15 min | 30 min | 45 min | 1.5 hrs |
|          | Before gal | 15 min | 30 min | 45 min | 1.5 hrs |
|          | Before gal | 15 min | 30 min | 45 min | 1.5 hrs |
|          | Before gal | 15 min | 30 min | 45 min | 1.5 hrs |
|          | Before gal | 15 min | 30 min | 45 min | 1.5 hrs |
|          | Before gal | 15 min | 30 min | 45 min | 1.5 hrs |

FIGURE 2. The catalytic subunit Cka2 of CK2 is important for efficient regulation of Cse4 levels. A, Cse4 was overexpressed from the GAL promoter on a 2-μm plasmid in WT, psh1Δ, cka1Δ, cka2Δ, and psh1Δcka2Δ strains. EV indicates an empty vector control as a point of comparison. 10-Fold serial dilutions of overnight cultures were plated to either SD-His or Gal-His medium. All strains grow similarly on SD-His but psh1Δ, cka2Δ, and psh1Δcka2Δ strains overexpressing Cse4 grow poorly. B, deletion of CKA2 stabilizes Cse4. WT, psh1Δ, and cka2Δ strains from A were used to perform a Cse4 ubiquitylation assay. After Gal induction of Cse4 overexpression, CHX was added, and cells were collected at the time points indicated for analysis by Western blotting using an anti-Cse4 antibody. C, Cka2 regulates endogenous levels of Cse4. Cse4-Myc was expressed from the native Cse4 promoter. Levels of Cse4-Myc were measured in cell lysates at the time points indicated following CHX treatment in WT and cka2Δ strains. In B and C, Pgk1 was the loading control. The same amount of total protein was loaded per lane. Graphs show quantifications of the Cse4 bands normalized to Pgk1. Cse4 t1/2 was calculated by fitting normalized Cse4 band intensity data to a first-order decay function as explained in Belle et al. (39).

anti-ubiquitin Western blot). Anti-Myc Western blotting showed similar Cse4-Myc levels in the lysates (Fig. 3A, left panel). Polyubiquitylated proteins were pulled down from MG132-treated samples using polyubiquitin affinity resin, and eluates were probed with anti-ubiquitin and anti-Myc antibodies (Fig. 3A, right panel). Pulldown samples probed with ubiquitin showed that the efficiency of enrichment of polyubiquitylated proteins was nearly the same. A psh1Δ strain was included as a control. We did not observe an obvious difference in polyubiquitylated Cse4-Myc (Ubα, Cse4-Myc) levels among WT, cka2Δ, and psh1Δ strains when Cse4-Myc was expressed at an endogenous level.

The Cse4 stability assay in Fig. 2B suggests that a reduction in the Ubα,Cse4 levels between WT and cka2Δ might be more obvious if Cse4 was overexpressed. Therefore, we examined Ubα,Cse4 levels in the strains in Fig. 2B (Fig. 3B). After Gal induction of Cse4, cells were treated with MG132. The method described in Liu et al. (21) was used to inhibit proteasome function using MG132. The efficiency of proteasome inhibition was confirmed by high accumulation of polyubiquitylated proteins.
in MG132-treated cells compared with DMSO control samples (Fig. 3B, left panel, anti-ubiquitin Western blot). Anti-Cse4 Western blotting showed similar Cse4 levels in the lysates (Fig. 3B, left panel). Polyubiquitylated proteins were pulled down as explained before, and samples were analyzed by Western blotting (Fig. 3B, right panel). The efficiency of enrichment of polyubiquitylated proteins in MG132-treated samples was very similar, and the signal was reduced in DMSO control samples. Ub$_{n}$-Cse4 was detected only in MG132-treated samples, confirming that these bands are polyubiquitylated Cse4 normally targeted for proteasomal degradation. Under our experimental conditions, we could not observe an obvious difference in Ub$_{n}$-Cse4 levels between WT and psh1Δ strains, suggesting that differences in ubiquitylation may be more difficult to detect than differences in protein stability. As expected, however, we detected a dramatic reduction in Ub$_{n}$-Cse4 levels in a cka2Δ strain compared with WT. These results further suggest that CK2 promotes ubiquitylation and degradation of Cse4.

Cse4 is only partially stabilized in a psh1Δ strain, indicating the presence of additional factors controlling Cse4 levels in vivo (11, 12). Interestingly, Cse4 was more stabilized by cka2Δ than psh1Δ (Fig. 2B), suggesting that CK2 may control factors in addition to Psh1. Ubc3/Cdc34, the E2 ubiquitin-conjugating enzyme for Psh1, is regulated by CK2 phosphorylation (15, 16). CK2 phosphorylation within the N-terminal catalytic domain of Ubc3 upregulates its ubiquitin charging activity (16). Therefore, in a cka2Δ mutant, ubiquitin charging of Ubc3 might be inefficient, causing extra stabilization of Cse4 levels. Recently, Fpr3 has been reported as another regulator of Cse4 stability (22). Fpr3 is also phosphorylated by CK2 (23). Moreover, there are several reports implicating CK2 in regulation of 26 S proteasome subunits via phosphorylation (24, 25). These reports, along with data from our studies, suggest that CK2 could regulate Cse4 levels via phosphorylation of Psh1, E2/Ubc3, other regulators of Cse4, and the proteasome. We continued to explore the role of CK2 phosphorylation in Psh1 activity in particular.

**FIGURE 3. Deletion of CKA2 reduces cellular levels of polyubiquitylated Cse4.** A, levels of Ub$_{n}$-Cse4-Myc were measured in WT, psh1Δ, and cka2Δ strains with a deletion in PDR5. Cse4-Myc was expressed from its native promoter. An untagged strain was used as a control. Cultures were grown to midlog phase and treated with either MG132 (100 μM) or DMSO for 3.5 h. Anti-ubiquitin and anti-Myc Western blots of cell lysates confirmed the efficiency of proteasome inhibition by MG132 and similar Cse4 levels, respectively (left panel). Polyubiquitylated proteins were pulled down from 2 mg of total proteins (from MG132-treated samples) using polyubiquitin affinity resin. Final elutions were probed with anti-ubiquitin and anti-Myc antibodies after SDS-PAGE (right panel). No obvious difference was detected in Ub$_{n}$-Cse4-Myc levels among WT, psh1Δ, and cka2Δ strains. A nonspecific band is marked with an asterisk. B, Ub$_{n}$-Cse4 levels are reduced in the cka2Δ strain. WT, psh1Δ, and cka2Δ strains from Fig. 2A were used to perform a Cse4 ubiquitylation assay. EV indicates the empty vector control. After Gal induction of Cse4 for 2 h, cells were treated with either MG132 (100 μM) or DMSO for another 2 h. Anti-ubiquitin and anti-Cse4 Western blots of cell lysates confirmed the efficiency of proteasome inhibition by MG132 and similar Cse4 levels, respectively (left panel). Polyubiquitylated proteins were pulled down, and final elutions were analyzed as in A (right panel).
**Phosphorylation of Psh1 Promotes Destruction of Cse4**—Because CK2 can potentially control ubiquitin-mediated proteolysis of Cse4 at multiple levels, stabilization of Cse4 in a cka2Δ strain may not directly reflect how Psh1 phosphorylation by CK2 affects the stability of Cse4. Therefore, we wanted to examine how Psh1 phosphorylation affects Cse4 using phosphomimic and phosphodepleted Psh1 mutants. We identified 10 major in vivo phosphorylation sites of Psh1 using MS (Fig. 1, A and B). All these residues are predicted to be potential CK2 sites by NetPhosK. In the phosphomimic Psh1 mutant (Psh1-S6D), six of these phosphorylation sites, identified using Psh1-TAP purification, were changed to aspartic acids (S143D, S191D, S229D, S230D, S275D, and S344D) to mimic phosphate groups. In the phosphodepleted Psh1 mutant (Psh1-S8A/T2A), all 10 identified phosphorylation sites were changed to alanine (S143A, S191A, S229A, S230A, S275A, T310A, S341A, S344A, T382A, and S403A) to prevent phosphorylation of Psh1. WT or Psh1 mutants were ectopically expressed from a plasmid in a psh1Δ background. Psh1 protein levels expressed from the plasmid were very similar as confirmed using an HA affinity-tagged version of these plasmids (Fig. 4A). However, compared with endogenous Psh1 levels, ectopic expression from the plasmid gave about 3 times more Psh1 (Fig. 4B).

If phosphorylation of Psh1 is important for Cse4 regulation, then overexpression of Cse4 in the phosphodepleted Psh1 mutant might be expected to show a growth phenotype similar to that of PSH1 and CKA2 deletion, and Cse4 protein levels should be stabilized. To test this, we overexpressed Cse4 in the presence of either Psh1-S6D or Psh1-S8A/T2A mutant (Fig. 4C). As expected, we observed slow growth with the Psh1-S8A/T2A strain. We next examined Cse4 stability in these strains. Cse4 was destabilized in the Psh1-S6D strain (Fig. 4D) and was dramatically stabilized in the Psh1-S8A/T2A strain (Fig. 4E). We further performed a co-IP in the presence of affinity-tagged Psh1-S6D-HA or Psh1-S8A/T2A-HA overexpressing Cse4 (Fig. 4F). Both Psh1-S6D-HA and Psh1-S8A/T2A-HA interacted with Cse4. This confirms that the stabilization of Cse4 in the presence of Psh1-S8A/T2A mutant is not the result of poor association of Cse4 with the mutant but more likely due to inefficient E3 ligase activity of the mutant. These results suggest that phosphorylation of Psh1 by CK2 may promote the E3 ubiquitin ligase activity of Psh1 for Cse4. Further confirming our hypothesis, we did not observe substantial Cse4 stabilization in a CKA2 deletion strain relative to WT in the presence of Psh1-S6D (Fig. 4G). The phosphomimetic mutant of Psh1 bypassed the effect of CKA2 deletion, and Cse4 was efficiently targeted for degradation (compare Fig. 4G with Fig. 2B, left panel). We did not observe an obvious Cse4 stabilization in a strain expressing the Psh1-S6A mutant, which is the phosphodepleted counterpart of Psh1-S6D (data not shown). Because the Psh1-S6A mutant still contains four major phosphorylation sites, lack of Cse4 stabilization in Psh1-S6A may suggest that Thr-310, Ser-344, Thr-382, and Ser-403 are some of the most important phosphorylation sites for activation of Psh1. These phosphorylation sites were detected mainly in centromere/kinetochore-bound Psh1 in the Dsn1-FLAG purification.

Next, we looked at Ub_{n}-Cse4 levels in the strains in Fig. 4, D and E, using the method explained earlier, polyubiquitylated protein pulldown and Western blotting (Fig. 5). Although we could not detect a difference in Ub_{n}-Cse4 in the strain expressing Psh1-S6D compared with a strain with wild type Psh1, an obvious reduction in Ub_{n}-Cse4 levels was observed between WT and the Psh1-S8A/T2A mutant (Fig. 5B). These results further support the idea that CK2 phosphorylation of Psh1 activates its E3 ubiquitin ligase activity for Cse4.

**Phosphorylation of Psh1 by CK2 May Control Ubiquitin Transfer onto Cse4**—During Cse4 ubiquitylation, Psh1 should act as an adaptor by recruiting Cse4, the substrate, and the ubiquitin-charged E2, Ub−Ubc3, thereby facilitating ubiquitin transfer from Ubc3 to Cse4. Phosphorylation may control interactions of Psh1 with Cse4 and/or Ub−Ubc3, ubiquitin transfer from Ub−Ubc3 to Cse4, or all of these steps. We have reported previously that Psh1 interacts with Cse4 and localizes to centromeres/kinetochores (11). In an affinity purification of kinetochore-associated proteins using Dsn1-FLAG followed by MS analysis, we detected Psh1 in both WT and cka2Δ strains (Fig. 6A). Therefore, phosphorylation of Psh1 by CK2 may not be important for Psh1 interaction with Cse4 and centromere localization. Supporting this, we observed no change in Cse4 association with the Psh1-S6D-HA or Psh1-S8A/T2A-HA mutants (Fig. 4F).

To further investigate how CK2 phosphorylation affects the interaction of Psh1 with Cse4, we performed a co-IP (Fig. 6B). Psh1-HA pulled Cse4-Myc down to a similar level in both WT and cka2Δ, confirming that phosphorylation by CK2 is not necessary for Psh1-Cse4 interaction. In another co-IP, we observed comparable levels of Psh1-HA coming down with Ubc3-TAP in both WT and cka2Δ backgrounds, demonstrating that phosphorylation by CK2 is not necessary for Psh1-Ubc3 interaction (Fig. 6C). Therefore we reasoned that phosphorylation does not control the physical interactions between Psh1 and Cse4 or Psh1 and Ubc3 but rather that the ubiquitin transfer from Ub−Ubc3 to Cse4 is controlled by phosphorylation. Supporting this idea, we detected reduced levels of Ub_{n}-Cse4 in both a cka2Δ strain and the strain expressing phosphodepleted Psh1-S8A/T2A (Figs. 3B and 5). Ubiquitin charging of Ubc3 has been reported to be positively regulated by CK2 phosphorylation of Ubc3. Therefore, reduced levels of Ub_{n}-Cse4 in the cka2Δ strain could be due to the collective effect of poor ubiquitin charging on Ubc3 and inefficient ubiquitin transfer from Ub−Ubc3 to Cse4. Reduced levels of Ub_{n}-Cse4 in the Psh1-S8A/T2A strain could reflect inefficient ubiquitin transfer from Ub−Ubc3 to Cse4.

Recent work suggests a role for protein subunits of the Skp1-Cdc53/Cullin-F box (SCF) complex in centromere/kinetochore regulation. For example, a subunit of the Mis18 complex, which regulates CENP-A localization in human cells, is targeted for proteolysis by SCF (26). SCF is an evolutionarily conserved multisubunit E3 ubiquitin ligase (27). Interestingly, Ubc3, the E2 ubiquitin-conjugating enzyme for Psh1, functions as the E2 for SCF E3 ligases (28). This raises the question whether Psh1 functions as a subunit of the SCF complex to facilitate Cse4 proteolysis. The stability of centromere identifier (CID), Drosophila CenH3, is regulated by Fpa, which is an F box protein subunit of the SCF^{TM} E3 ligase complex (13). We tested whether Psh1 can interact with Cdc53, an invariant core sub-
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A  pMOBY-Psh1

B  Endogenous Psh1-HA (WT) expressed under the native promoter. Cell lysates probed with anti-HA antibody show an about 3 times higher Psh1-HA level after ectopic expression. Similar Psh1-HA levels are detected in phosphodepleted (Psh1-S8A/T2A-HA) and phosphomimic mutants (Psh1-S6D-HA) ectopically expressed on a plasmid. Co-immunoprecipitation using anti-Cse4 antibody pulls down Psh1-HA from WT, Psh1-S6D-HA, or Psh1-S8A/T2A-HA strain. A control IP was performed without antibody.

C  Remaining Cse4 %

D  WT/S6DWT/S6D

E  With Cse4 overexpression

F  Cse4

G  With Cse4 overexpression

FIGURE 4. Phosphorylation of Psh1 promotes its E3 activity for Cse4. A, ectopic expression levels of HA-tagged WT, phosphomimic mutant (Psh1-S6D-HA), and phosphodepleted (Psh1-S8A/T2A-HA) Psh1 were compared. Cell lysates probed with anti-HA antibody show very similar Psh1-HA levels. Intensities of the Psh1-HA bands relative to WT protein are indicated. B, ectopic expression level of WT Psh1-HA was compared with endogenous Psh1-HA (WT) expressed under the native promoter. Cell lysates probed with anti-HA antibody show an about 3 times higher Psh1-HA level after ectopic expression. C, a growth assay was performed using phosphomimic and phosphodepleted Psh1 strains. In these strains, the endogenous copy of PSH1 is deleted, and Psh1-S6D or Psh1-S8A/T2A was ectopically expressed using a plasmid. Cse4 was overexpressed from the GAL promoter on a 2-μm plasmid. D, ectopic expression level of WT Psh1-HA was compared with endogenous Psh1-HA (WT) expressed under the native promoter. Cell lysates probed with anti-HA antibody show an about 3 times higher Psh1-HA level after ectopic expression. Similar Psh1-HA levels are detected in phosphodepleted (Psh1-S8A/T2A-HA) and phosphomimic mutants (Psh1-S6D-HA) ectopically expressed on a plasmid. Cse4 was overexpressed from the GAL promoter on a 2-μm plasmid. E, F, G, H, and I, anti-HA antibody was used for Western blotting using an anti-HA antibody. Cse4 was destabilized in the presence of the phosphomimic Psh1 mutant. E, the stability of Cse4 in the presence of phosphomimic Psh1 mutant (Psh1-S6D) was measured using a protein stability assay. After 2 h of Cse4 induction, CHX was added, and cells were collected at the time points indicated for analysis by Western blotting using an anti-Cse4 antibody. Cse4 was destabilized in the presence of the phosphomimic Psh1 mutant. F, phosphodepleted and phosphomimic mutants of Psh1 interact with Cse4. Cse4 was overexpressed from the GAL promoter on a 2-μm plasmid in the presence of affinity-tagged Psh1-S6D-HA or Psh1-S8A/T2A-HA. Co-immunoprecipitation using anti-Cse4 antibody pulls down Psh1-HA from WT, Psh1-S6D-HA, or Psh1-S8A/T2A-HA strain. A control IP was performed without antibody. G, phosphomimic Psh1 mutant can bypass the effect of Cka2 deletion on Cse4 stability. In these strains, the endogenous copy of PSH1 is deleted, and Psh1-S6D was ectopically expressed using a plasmid. The stability of Cse4 was measured using a protein stability assay as in D. In the presence of the phosphomimic Psh1 mutant, Cse4 stabilization in a cka2Δ strain relative to WT is very mild compared with the stability observed in a cka2Δ strain in the presence of wild type Psh1 (see Fig. 2B, left panel). H, D, E, and G, Pgk1 strain was the loading control. The same amount of total protein was loaded per lane. The graphs show quantification of the Cse4 bands normalized to Pgk1. Cse4 t1/2 was calculated as in Fig. 2. Ab, antibody.

unit of SCF, using a co-IP. We did not observe any interaction (data not shown), suggesting that Psh1/Ubc3 acts on Cse4 independently of other SCF components.

We previously reported an autoubiquitylation site, Lys-303 (indicated in Fig. 1B), on Psh1 identified by in vitro ubiquitylation followed by MS (11). This indicates that Psh1 levels could also be controlled through proteolysis. In vivo, Psh1 ubiquitylation could be mediated by itself and by one or more other E3 ligases. Phosphorylation of Psh1 by CK2 may regulate ubiquitylation and proteolysis of Psh1 as well. To test this, we per-
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FIGURE 5. Ub₇₃-Cse4 levels are reduced in the presence of the phosphodepleted Psh1 mutant. A, strains from Fig. 4 were used to measure polyubiquitylated Cse4 levels as in Fig. 3. A wild type strain without Gal induction (WT/No Gal) was used as a control. Anti-ubiquitin and anti-Cse4 Western blots of cell lysates confirmed the efficiency of proteasome inhibition by MG132 and similar Cse4 levels, respectively. B, polyubiquitylated proteins were pulled down, and final elutions were analyzed as in Fig. 3. An anti-Cse4 Western blot showed reduced levels of Ub₇₃-Cse4 in the presence of the phosphodepleted Psh1 mutant.

| Kinetochore purification using Dsn1-FLAG | WT | cka2Δ |
|-----------------------------------------|----|-------|
| Sequence coverage %                     | NSAF| NSAF  |
| Psh1                                    | 39.41| 0.0071|
|                                         | 33.5 | 0.0039|

FIGURE 6. The interaction of Psh1 with Cse4 and Ubc3 was unaffected by phosphorylation. A, Psh1 co-purifies with kinetochores in a cka2Δ strain. Purification of kinetochores via Dsn1-FLAG followed by MS analysis shows Psh1 in both WT and cka2Δ strains. The normalized spectral abundance factor (NSAF) shows the estimated relative Psh1 protein abundance. B, Psh1-Cse4 interaction was unaffected by deletion of CKA2. Strains expressing Cse4-Myc/ Psh1-HA were used for co-IP. A control IP was performed from a strain expressing Cse4-Myc but lacking Psh1-HA. Psh1-HA pulls down Cse4-Myc in lysates made from either WT or a cka2Δ strain. C, the interaction between Psh1 and Ubc3 was unaffected by deletion of CKA2. Strains expressing Ubc3-TAP/Psh1-HA were used for co-IP. A control IP was performed from a strain expressing Psh1-HA but lacking Ubc3-TAP. Ubc3-TAP pulls down Psh1-HA in lysates made from either WT or a cka2Δ strain.

formed a Psh1 stability assay using a strain expressing Psh1-HA as the sole copy from the endogenous promoter (Fig. 7A). We observed stabilized Psh1 levels with CKA2 deletion. We further examined Ub₇₃-Psh1 levels in vivo (Fig. 7B). If CK2 phosphorylation of Psh1 facilitates ubiquitylation of Psh1, then Ub₇₃-Psh1-HA should be diminished in a cka2Δ strain. Although Psh1 was stabilized in cka2Δ strain, we did not see a reduction in Ub₇₃-Psh1-HA (Fig. 7B, lower panel). Therefore, phosphorylation of Psh1 by CK2 may not control ubiquitylation of Psh1, or the difference may be beyond the detection limit of this assay such as we observed with Cse4 ubiquitylation in the psh1Δ strain.

Therefore, we examined Ub₇₃-Cse4 levels in vivo (Fig. 7B). If CK2 phosphorylation of Psh1 facilitates ubiquitylation of Psh1, then Ub₇₃-Psh1-HA should be diminished in a cka2Δ strain. Although Psh1 was stabilized in cka2Δ strain, we did not see a reduction in Ub₇₃-Psh1-HA (Fig. 7B, lower panel). Therefore, phosphorylation of Psh1 by CK2 may not control ubiquitylation of Psh1, or the difference may be beyond the detection limit of this assay such as we observed with Cse4 ubiquitylation in the psh1Δ strain. Supporting the idea that phosphorylation may not facilitate ubiquitylation and degradation of Psh1, we did not observe a stabilization of Psh1-S8A/T2A-HA or destabilization of Psh1-S6D-HA compared with Psh1 (Fig. 7, C and D). In fact, the phosphodepleted mutant, Psh1-S8A/T2A-HA, was destabilized compared with WT (Fig. 7C). Although this mutant could associate with Cse4 to a similar level as wild type Psh1 (Fig. 4F), it might be recognized as a defective protein due to compromised E3 ligase function and targeted for faster degradation by cellular mechanisms.

Cse4 Is Mislocalized in a cka2Δ Strain—Overexpression of Cse4 in a psh1Δ strain was toxic, and Cse4 was significantly mislocalized (Figs. 2A and 8A and Ref. 11). Because a cka2Δ strain showed high Cse4 levels, we predicted that Cse4 mislocalization would increase. We used Cse4 strains from Fig. 2A to examine Cse4 levels at non-centromeric regions. We followed Cse4 at the ribosomal DNA and PHO5 promoter using ChIP followed by quantitative PCR (Fig. 8A). Cse4 mislocalization to these regions was elevated in a cka2Δ strain compared with a WT strain. However, the mislocalization was significantly lower compared with a psh1Δ strain, although the overall level of Cse4 in the cka2Δ strain was higher than that observed in a psh1Δ strain. Cse4 mislocalization was increased with the psh1Δcka2Δ double deletion compared with the cka2Δ single deletion. Both Cse4 and Psh1 were significantly stabilized in a cka2Δ strain (Figs. 2B and 7A), and Psh1-Cse4 interaction was not disrupted by the deletion of CKA2 (Fig. 6B). Therefore, we suggest that the association of Cse4 with Psh1 may reduce non-
centromeric deposition of Cse4 in the cka2Δ strain. Psh1-Cse4 association could reduce the amount of Cse4 available to the histone chaperones responsible for non-centromeric deposition. When Psh1 is removed in a cka2Δ strain, more Cse4 becomes available for chaperones, leading to increased misincorporation of Cse4.

Scm3 is a centromeric nucleosome assembly factor (29) and is essential to recruit Cse4 to centromeres and to maintain a functional kinetochore (30). Growth of a Scm3 deletion or shut-off (Scm3off) strain can be rescued by overexpression of Cse4, and deletion of PSH1 improves rescue (11, 30). Under these conditions, some other histone chaperone must be responsible for the assembly of Cse4 nucleosomes. (For instance, the H3.3 chaperone death domain associated protein (DAXX) was reported to regulate ectopic localization of CENP-A in human cells (31)). If Psh1-Cse4 association prevents Cse4 availability for nucleosome assembly, then deletion of cka2Δ may not rescue the Scm3off strain. To test this, we performed a growth assay using strains containing Gal-inducible endogenous Scm3 and copper-inducible Cse4 (Fig. 8B). The level of Cse4 can be controlled with copper concentration. Growth of WT, psh1Δ, and cka2Δ strains was compared under Scm3on and Scm3off conditions. We observed no growth difference between the strains when Scm3 was on. However, when Scm3 was off in the
FIGURE 8. Non-centromeric localization of Cse4 is increased in a cka2Δ strain. A, ChIP was performed for Cse4 in strains overexpressing Cse4 from Fig. 2A. Quantitative PCR was used to detect Cse4 levels for the PHO5 promoter and ribosomal DNA. The y axis indicates arbitrary units representing the enrichment for each sequence from ChIP performed with and without antibody with respect to the signal for total chromatin for each sample. The error bars represent the standard deviation for ChIP performed in triplicate. The same pattern of Cse4 localization in the PHO5 promoter and ribosomal DNA was observed in a biological replicate, confirming the reproducibility of the results. B, deletion of CKA2 does not improve growth of Scm3off strain. Strains were constructed in which the only copy of Scm3 was under the control of the GAL promoter, allowing the expression of Scm3 to be controlled with glucose/galactose. Cse4 was under the control of a copper-inducible promoter on a 2-microvector. EV indicates empty vector. When Scm3 is expressed (Scm3off), no growth differences are observed in 10-fold serial dilutions. Lower levels of induction of Cse4 (SD-Ura + 0.1 mM Cu2+) rescue Scm3off when PSH1 is deleted (psh1Δ) as compared with WT. CKA2 deletion does not improve growth of Scm3off under the same conditions. Ab, antibody.

cka2Δ strain compared with the psh1Δ strain, rescue of growth by Cse4 overexpression was poor (Fig. 8B, Scm3off). This result is consistent with the notion that Cse4 availability for nucleosome assembly requires deletion of PSH1, and CKA2 deletion does not suffice. Together, our data suggest that Psh1-Seq4 physical association itself functions to prevent mislocalization of Cse4.

DISCUSSION

Here we report that phosphorylation by CK2 facilitates Psh1-assisted Cse4 degradation. Lack of complete stabilization of Cse4 following deletion of PSH1 suggested additional regulation of Cse4 stability (11). We demonstrate that phosphorylation events by CK2 provide an additional mechanism for regulating Cse4 stability. Based on our findings, we propose a model for how Cse4 is controlled by Psh1 and CK2. In wild type cells (Fig. 9A), CK2 phosphorylates Psh1, activating the E3 ubiquitin ligase function of Psh1 toward Cse4. CK2 phosphorylation of Ubc3 also improves ubiquitin transfer from Ub~Ubc3 to Cse4. Activation of Psh1 promotes efficient removal of the soluble pool of Cse4 and may also improve the removal of misincorporated Cse4. This regulation is especially important when Cse4 is overexpressed. In a psh1Δ strain (Fig. 9B), lack of Psh1-mediated proteolysis results in stabilization and misincorporation of Cse4. The histone chaperones responsible for the misincorporation of Cse4 remain to be identified. In a cka2Δ strain (Fig. 9C), the lack of Psh1 and Ubc3 phosphorylation dramatically stabilizes Cse4. However, the reduced Psh1/Ubc3 activity is not problematic because Psh1 is also stabilized and can act as an effective sink for excess Cse4, preventing Cse4 misincorporation.

We suggest that Psh1/CK2 may help to control both the centromeric and non-centromeric pools of Cse4. Two recent studies show that Pat1 localizes to centromeres and maintains a pool of Cse4 in the vicinity of the kinetochore cluster (32, 33). Centromere/kinetochore-bound Psh1 and CK2 may help regulate this accessory Cse4 pool. Psh1 may also target non-centromeric Cse4 with the help of the facilitates chromatin transcriptions (FACT) complex, a chromatin disassembly/reassembly factor (12). Psh1 was first discovered associated with Spt16 and Pob3, components of the budding yeast FACT complex. All four subunits of CK2 co-purify with Spt16 along with Psh1 (34), and the Cka2 subunit of CK2 associates with chromosome arms as revealed by ChIP (35). This suggests that CK2 phosphorylation could regulate Psh1 targeting of misincorporated Cse4 as well.

Regulation of kinetochore proteins by kinases/phosphatases is well known. The evolutionarily conserved kinase Ipl1/Aurora B regulates kinetochore function through phosphorylation of kinetochore proteins (36). Opposing activities of Mps1 and PP1 regulate the phosphorylation level of kinetochore protein Spc105 (37). Evidence from previous reports suggests that CK2 might regulate kinetochore function. Studies using temperature-sensitive mutant cka1Δ cka2-8 demonstrate that Cka2 functions in cell cycle progression (35). cka1Δ CKA2 cells showed DNA content similar to wild type and normal mitotic spindle elongation and chromatid separation. However, cka1Δ cka2-8 cells had abnormal DNA profiles, short spindles, and...
problems with sister chromatid separation at non-permissive temperature, suggesting an important role of Cka2 in kinetochore function and chromosome segregation. Interestingly, Mif2 and Ndc10 are targets of CK2 (35). Although Lpl1 and CK2 show an antagonistic effect on Mif2 stability, the two kinases have a synergistic effect on Ndc10. By analogy, CK2 may play an antagonistic/synergistic role with another kinase for Psh1. The phosphorylation of Psh1 may be regulated by additional phosphatases or deubiquitylases. In any event, our results continue to implicate CK2 as an important kinase regulating kinetochore-associated proteins.

Our work reported here identifies a previously unknown function of CK2 in Cse4 regulation. Our study lays the foundation for future work on the role of CK2 in proteolysis of Cse4 in budding yeast and potentially of CenH3 variants or additional kinetochore-associated proteins.

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