State changes of the HORMA protein ASY1 are mediated by an interplay between its closure motif and PCH2

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ABSTRACT

HORMA domain-containing proteins (HORMADs) play an essential role in meiosis in many organisms. The meiotic HORMADs, including yeast Hop1, mouse HORMAD1 and HORMAD2, and Arabidopsis ASY1, assemble along chromosomes at early prophase and the closure motif at their C-termini has been hypothesized to be instrumental for this step by promoting HORMAD oligomerization. In late prophase, ASY1 and its homologs are progressively removed from synapsed chromosomes promoting chromosome synopsis and recombination. The conserved AAA+ ATPase PCH2/TRIP13 has been intensively studied for its role in removing HORMADs from synapsed chromosomes. In contrast, not much is known about how HORMADs are loaded onto chromosomes. Here, we reveal that the PCH2-mediated dissociation of the HORMA domain of ASY1 from its closure motif is important for the nuclear targeting and subsequent chromosomal loading of ASY1. This indicates that the promotion of ASY1 to an 'unlocked' state is a prerequisite for its nuclear localization and chromosomal assembly. Likewise, we find that the closure motif is also necessary for the removal of ASY1 by PCH2 later in prophase. Our work results in a unified new model for PCH2 and HORMADs function in meiosis and suggests a mechanism to contribute to unidirectionality in meiosis.

INTRODUCTION

A fundamental process for sexual reproduction is meiosis during which one round of DNA replication is followed by two consecutive nuclear divisions resulting in the reduction of the chromosome set of a cell by half. Importantly, genetic information from maternal and paternal progenitors is re-shuffled during meiosis through homologous recombination, thereby generating genetic variation and thus contributing to the diversity of life.

Recombination requires the formation of a meiosis-specific proteinaceous structure called chromosome axis that spans along the entire length of the chromosomes. The chromosome axis is believed to organize sister chromatids into a loop-array configuration and thus facilitates interactions of homologous chromosomes (1–3). The chromosome axis consists of cohesin complexes encompassing sister chromatids, and other meiosis-specific proteins including the meiotic HORMA domain-containing protein (HORMADs) family (ASY1 in Arabidopsis, Hop1 in yeast; HORMAD1/2 in mouse); the coiled-coil domain-containing ‘linker’ proteins (ASY3 in Arabidopsis, Red1 in yeast; SYCP2 in mouse); and the small coiled-coil protein family proteins (ASY4 in Arabidopsis, the mammalian SYCP3/SCP3 homolog) (4–11). Current models suggest that the coiled-coil proteins form filamentous complexes that form the core of the chromosome axis. These filaments are thought to localize on chromosomes through binding to cohesin complexes and thereby organize the loop-array structure of meiotic chromosomes (12).

The meiotic HORMADs are involved in many key meiotic events, i.e. double-strand break (DSB) formation, synaptonemal complex assembly and crossover formation (4,5,13,14). These proteins are characterized by an N-terminal HORMA domain (for Hop1, Rev7 and MAD2), a conserved domain which interacts with a short sequence motif termed ‘closure motif’ (15,16). The HORMA domain-to-closure motif interaction is thought to anchor ASY1 and other HORMADs to the axis via interaction with the closure motif of ASY3 and its orthologs (9,12,17).

All meiotic HORMADs studied, including Hop1, HORMAD1/2 and ASY1 also contain a closure motif at their own C-terminus, and because of this, these HORMADs tend to fold into a HORMA domain-closure motif bound state called self-‘closed’ state (12,15,17,18). This self-closed state of HORMADs is expected to block the binding of their HORMA domain with the closure motifs.
of the linker proteins such as ASY3, raising the question of how this interaction is formed \textit{in vivo} (15,17,18). It has been proposed that the conformational conversion from a self-closed to the ‘unlocked’ state might be mediated by the conserved AAA+ ATPase PCH2/TRIP13 protein (15,17,19). However, there is no experimental evidence to support this up to now.

The meiotic HORMADs undergo a highly dynamic assembly and disassembly process, which is essential for the homologous chromosome synapsis and recombination. The removal of meiotic HORMADs from chromosome axes at late prophase is catalyzed by PCH2/TRIP13 and has been extensively studied in different organisms including budding yeast, mouse and Arabidopsis (20–22). However, how the chromosomal assembly of meiotic HORMADs is controlled is much less clear. Recently, we have shown that the position and origin (the sequence \textit{per se}) of the closure motif are not essential for its functional interplay with PCH2.

\textbf{MATERIALS AND METHODS}

\textbf{Plant materials}

The Arabidopsis thaliana accession Columbia (Col-0) was used as wild-type reference throughout this research. The T-DNA insertion lines SALK\textsubscript{046272} (as1\textsubscript{1-4}) (23) and SALK\textsubscript{031464} (pch2\textsubscript{2-2}) (20) were obtained from the T-DNA mutant collection at the Salk Institute Genomics Analysis Laboratory (SIGnAL, http://signal.salk.edu/cgi-bin/tdnaexpress) via NASC (http://arabidopsis.info/). The \textit{PROASY1:ASY1:GFP}, \textit{PROASY1:ASY1:RFP} and \textit{PROZYP1B:ZYP1B:GFP} reporters were described previously (18,24). All plants were grown in growth chambers at 21°C under a 16 h light/8 h dark cycle.

\textbf{Plasmid construction and plant transformation}

To generate the \textit{ASY1\textsubscript{Δclosure}:GFP} reporter, the PCR for the closure motif deletion was performed with the primers (gASY1\textsubscript{1–570aa-R} and mGFP-F) using the entry clone of \textit{PROASY1:ASY1:GFP} as a template and then re-ligated, producing the \textit{PROASY1:ASY1\textsubscript{1–570}:GFP} construct. For creating the \textit{ASY1-NLS:GFP} reporter, the PCR fragments were obtained with the primer pair (gASY1\textsubscript{R} and mGFP-F) using \textit{PROASY1:ASY1:GFP} as a template and then re-ligated, producing the \textit{PROASY1:ASY1-NLS:GFP} construct. For creating the \textit{asy1\textsubscript{1-300}::GFP} and \textit{asy1\textsubscript{1-200}::GFP} reporters, PCR fragments were amplified with primer pairs (gASY1\textsubscript{1-11aa-F} and gASY1\textsubscript{-intron1-R or ASY1-intron2-F} and gASY1\textsubscript{-intron1-R}) and subsequently self-ligated, generating the \textit{PROASY1::ASY1\textsubscript{Δ2-10}:GFP\textsubscript{pENTR2B} and PROASY1::ASY1\textsubscript{Δ2-20}:GFP\textsubscript{pENTR2B}} constructs. For creating the \textit{ASY1\textsubscript{Δclosure}:GFP\textsubscript{pENTR2B}, the backbone of the construct was obtained with primer pairs (ASY1\textsubscript{T142V-G} and mGFP-SmaI-R) using \textit{PROASY1:ASY1\textsubscript{1–570}:GFP\textsubscript{pENTR2B}} as a template, and the closure motif insert (ASY1\textsubscript{571–596aa-F} and ASY1\textsubscript{571–596aa-R}) was retrieved by annealing of two synthesized long primers (ASY1\textsubscript{571–596aa-F} and ASY1\textsubscript{571–596aa-R}). Next, the \textit{PROASY1:ASY1\textsubscript{1–570}:GFP\textsubscript{pENTR2B}} construct was generated by integration of the backbone and the insert using a SLICE reaction. For creating the \textit{ASY1\textsubscript{Δclosure}:ASY1\textsubscript{1-500}:NLS:GFP} reporter, two types of PCR fragments with 25bp overlapping sequence in both ends, i.e. ASY1 backbone and ASY3 1–50aa insert, were amplified with two primer pairs (NLS-SmaI-F and gASY1\textsubscript{1–570aa-R}, ASY3-SLICE-F and ASY3-SLICE-50aa-R) using \textit{PROASY1:ASY1:GFP\textsubscript{pENTR2B}} or \textit{ASY3/pDONR223} (generated in (24)) as a template. Subsequently, a SLICE reaction was performed by mixing the ASY1 backbone with ASY3 1–50aa fragments, producing the \textit{PROASY1:ASY1\textsubscript{1–570}:ASY3\textsubscript{1-50aa}:NLS:GFP\textsubscript{pENTR2B}} entry construct. For creating the \textit{ASY1\textsubscript{T142V-T184G}:GFP} reporter, a PCR-based mutagenesis was performed using with primer pair (ASY1\textsubscript{T142V-G} and mGFP-SmaI-R) using \textit{PROASY1:ASY1\textsubscript{1–570}:GFP\textsubscript{pENTR2B}} as a template and the resulting fragments were then re-ligated.

Next, all these resulting expression cassettes were integrated into the destination vector \textit{pGWB501/pGWB601} by the gateway LR reaction. All constructs were transformed into \textit{Arabidopsis thaliana} plants by floral dipping.

For constructs of the yeast two-hybrid assays, the \textit{ASY1\textsubscript{571–596}:AD, ASY1\textsubscript{1-300-BD} and ASY3\textsubscript{-FL-AD}} constructs were generated previously (18). To generate \textit{ASY1\textsubscript{1-300}/AD\textsubscript{1-300}:BD} and \textit{ASY1\textsubscript{1-300}/AD\textsubscript{1-200}:BD}, a PCR-based fragment deletion was performed with two primer pairs (ASY1\textsubscript{1-11aa-F} and ATG-attL1-R2, ASY1\textsubscript{21aa-F} and ATG-attL1-R2) using \textit{ASY1\textsubscript{1-300}/pDONR223} as a template and the resulting fragments were then re-ligated producing the entry clones. To create \textit{ASY3\textsubscript{200}/AD\textsubscript{200} and ASY3\textsubscript{200}/FL-AD}, the coding sequences of the respective fragments were amplified by PCR with primers flanked by attB recombination sites (ASY3-attB1-F and ASY3-200aa-attB2-R, ASY3-201aa-attB1-F and ASY3-attB2-R) and subcloned into \textit{pDONR223} vector by gateway BP reactions. All these entry clones were subsequently integrated into...
the pGADT7-GW or pGBK7-GW vectors by gateway LR reactions. Primers used for generating all constructs mentioned above are shown in Supplemental Table S1.

Microscopy

Images of pollen staining were taken using an Axiophot microscope (Zeiss). For the protein localization analyses in male meioocytes, young anthers harboring the relevant reporters were dissected and imaged directly using the Leica TCS SP8 inverted confocal microscope. The meiotic stages for live cell imaging were determined by combining the criteria of the chromosome morphology, nucleolus position, and cell shape (25).

Yeast two-hybrid assay

Yeast two-hybrid assays were performed according to the manual of Matchmaker Gold Yeast two-hybrid system (Clontech). The relevant combinations of constructs were co-transformed into yeast strain AH109 using the polyethylene glycol/lithium acetate method as described in the manual. Yeast cells expressing the relevant proteins were dotted on the plates of double (-Leu-Trp), triple (-Leu-Trp-His) and quadruple (-Leu-Trp-His-Ade) synthetic dropout medium to test the protein-protein interactions.

Cytological analysis

Chromosome spread analyses were performed as described previously (18). In brief, fresh flower buds were fixed in fixation solution containing 75% ethanol and 25% acetic acid for 48 h at 4°C. After two times of washing with 75% ethanol, the fixed flower buds were stored in 75% ethanol at 4°C. To perform chromosome spread, flower buds were first digested in the enzyme solution (10 mM citrate buffer containing 1.5% cellulose, 1.5% pectolyase and 1.5% cytohelicase) for 3 h at 37°C. Subsequently, single flowers were transferred on a glass slide followed by a fine smashing with a bended needle in the enzyme solution. The spreading step was performed on a 46°C hotplate after adding 10 µl of 45% acetic acid. Subsequently, the slide was rinsed with ice-cold ethanol/acetic acid (3:1) solution and mounted with DAPI solution (Vector Laboratories).

Immunolocalization analyses was performed according to (24). Briefly, fresh flower buds were first sorted by size and then intact anthers likely at meiotic stage were collected and macerated in 10 µl enzyme solution (0.4% cytohelicase and 1% polyvinilpyrrolidone) on the Poly-Prep slides (Sigma) and macerated in 10°C containing 1.5% cellulose, 1.5% pectolyase and 1.5% cytohelicase for 3 h at 37°C. Subsequently, single flowers were transferred onto a glass slide followed by a gentle stirring and drying at room temperature for 2–3 h. Afterwards, the slides were washed three times with PBST buffer (PBS with 1% Triton X-100), and were then blocked in PBST containing 1% BSA for 1 h at 37°C in the moisture chamber. Then the slides were incubated with anti-GFP (Takara 632381/JL-8)) (1:100 dilution) and/or anti-ZYP1 (1:500 dilution) antibodies at 4°C for 48 h. Next, following three times of washing (10 min each) in PBST, the slides were incubated with fluorescein-conjugated secondary antibodies for 24 h incubation at 4°C in a moisture chamber. After three times of washing, the DNA was counterstained with anti-fade DAPI solution (Vector Laboratories). Images were captured using the Leica SP8 laser scanning microscopy.

RESULTS

The closure motif is not required for chromosomal association of ASY1

Previous experiments suggested that the closure motif is involved in the chromosome association of ASY1 (17,18,26). To test this, we introduced the previously generated deletion construct of ASY1 (PROASY1:ASY1Δ571-596:GFP, designated ASY1Δclosure::GFP) in which the closure motif comprising the last 25 amino acids (aa) of ASY1 (596 aa in total) is removed from a functional reporter construct (PROASY1:ASY1::GFP, called ASY1::GFP) (18), into asyl mutants (Figure 1A).

In asyl mutants harboring the full length reporter, ASY1::GFP accumulated in the nuclei of male meioocytes during meiotic prophase I, as reported previously (18) (Figure 1B, Table 1). In contrast, ASY1Δclosure::GFP was found to be present only in the cytoplasm and no signal was detected in the nucleus, suggesting that the closure motif plays a role in nuclear targeting of ASY1 (Figure 1B, Table 1). To test this possibility, we generated plants containing a construct, in which only the closure motif sequence fused with GFP driven by the ASY1 promoter (PROASY1:ASY1Δ571-596::GFP, called closure::GFP) was present (Figure 1A). Unexpectedly, we did not detect any signal of the closure::GFP in male meioocytes (Figure 1C), indicating that for the proper expression in meiocytes sequence information from the introns of ASY1 are necessary. Nonetheless, closure::GFP was found to be expressed in the epidermal cells of the connective tissue of anthers, where it specially localized in nuclei, corroborating that the closure motif of ASY1 functions as a nuclear localization signal (NLS) (Figure 1C, CLOSE-UP).

To test whether the closure motif is required for chromosomal association of ASY1 after nuclear import, we next generated a separation-of-function allele, in which the closure motif was substituted with the NLS sequence of the SV40 Large T-Antigen (PKKKRKV) (PROASY1:ASY1Δ571-596_NLS::GFP, called ASY1Δclosure::NLS:GFP) (Figure 1A). Notably, ASY1Δclosure::NLS::GFP was exclusively present in the nuclei of male meioocytes in asyl mutants suggesting that the closure motif indeed functions in nuclear targeting of ASY1 (Figure 1B, Table 1).

To further dissect the importance of the closure motif for the function of ASY1, we first performed immunofluorescence experiments using an antibody against GFP and compared in detail the localization of ASY1::GFP and ASY1Δclosure::NLS::GFP in male meioocytes of asyl mutants. To this end, we found that ASY1Δclosure::NLS::GFP showed an indistinguishable chromosomal association from
Figure 1. Closure motif of ASY1 functions as a nuclear localization signal (NLS) and is not required for its chromosomal localization. (A) Schematic of different ASY1 versions and ASY3 used in this study. (B) Localization of ASY1:GFP, ASY1Δclosure:GFP, and ASY1Δclosure-NLS:GFP in the male meiocytes of asy1 mutants using confocal microscopy. Bar: 10 μm. (C) Localization of closure:GFP in the anther of wildtype using confocal microscopy. The CLOSE-UP shows the magnification of the connective tissue highlighted by the red rectangle. Bar: 10 μm. Arrow indicates the position of male meiocytes. (D) Localization of ASY1:GFP and ASY1Δclosure-NLS:GFP at different prophase stages in male meiocytes of asy1 mutants. Bar: 5 μm.

Table 1. Summary of the subcellular localization of different versions of ASY1 used in this study in different plant backgrounds. N/A denotes not analyzed.

| Constructs                          | Background       | Source            | Subcellular localization | Functionality |
|-------------------------------------|------------------|-------------------|--------------------------|--------------|
| ASY1:GFP                            | WT and asy1      | From (18)         | Nucleus                  | Functional   |
| ASY1:GFP                            | pch2             | From (18)         | Cytoplasm and nucleus    | Functional   |
| ASY1:GFP                            | PCH2E283Q (pch2) | This study        | Cytoplasm and nucleus    | Functional   |
| ASY1Δclosure:GFP                     | asy1             | This study        | Cytoplasm                | Non-functional |
| ASY1Δclosure-NLS:GFP                | asy1             | This study        | Nucleus                  | Non-functional |
| ASY1Δ100:GFP                         | WT and asy1      | This study        | Nucleus                  | Functional   |
| ASY1Δ200:GFP                         | WT               | This study        | Nucleus                  | N/A          |
| ASY1Δ142V/184D:GFP                  | WT               | This study        | Nucleus                  | Non-functional |
| ASY1Δclosure,GFP:closure             | WT and asy1      | This study        | Nucleus                  | Functional   |
| ASY1Δclosure,GFP:closure             | pch2             | This study        | Cytoplasm and nucleus    | Functional   |
| ASY1Δclosure-ASY31-50-NLS:GFP       | pch2             | This study        | Cytoplasm and nucleus    | N/A          |
ASY1-GFP forming a thread-like signal along all chromosomes in leptotene (Supplementary Figure S1A). This was also confirmed by the localization patterns of ASY1-GFP and ASY1\textsuperscript{-\textit{closure-NLS}}:GFP in live cells of \textit{asy1} mutants by confocal microscopy (Figure 1D). Taken together, these results suggest that the closure motif of ASY1 is not essential for its chromosomal association.

Deletion of the closure motif dominantly interferes with synopsis

Given the apparently correct localization on chromosomes, we asked whether ASY1\textsuperscript{-\textit{closure-NLS}}:GFP was also functional. However, whereas ASY1-GFP fully rescued the fertility defects of \textit{asy1} mutants, ASY1\textsuperscript{-\textit{closure-NLS}}:GFP did not rescue the \textit{asy1} mutants, as revealed by short silique (7.57 ± 0.96 seeds in \textit{asy1} mutants versus 8.0 ± 0.82 in ASY1\textsuperscript{-\textit{closure-NLS}}:GFP (asy1), \(P = 0.39\)), high pollen abortion and defective chromosome segregation (Supplementary Figure S2). This suggests that the closure motif is important for ASY1 function beyond promoting its nuclear localization. Notably, ASY1\textsuperscript{-\textit{closure-NLS}}:GFP functioned in a dominant negative manner since we also observed apparent fertility defects in 27 out of 35 T1 plants when this construct was introduced into a wild-type background (called ASY1\textsuperscript{-\textit{closure-NLS}}:GFP/WT) (Supplementary Figure S2A and B).

To exclude that this loss of ASY1 functionality was due to the added NLS signal, we created a full-length version of ASY1 with this NLS sequence (PRO\textsubscript{ASY1}-ASY1\textsuperscript{-\textit{NLS}}:GFP, called ASY1\textsuperscript{-\textit{NLS}}:GFP) (Figure 1A) and introduced this construct into wild-type and \textit{asy1} mutant plants. Most of the wild-type T1 transformants (35 out of 40) expressing the ASY1\textsuperscript{-\textit{NLS}}:GFP (called ASY1\textsuperscript{-\textit{NLS}}:GFP/WT) were fully fertile, and the fertility of \textit{asy1} mutant plants harboring this construct was fully restored (Supplementary Figure S2A and B), suggesting that ASY1\textsuperscript{-\textit{NLS}}:GFP was a functional protein.

Next, we compared the ASY1\textsuperscript{-\textit{NLS}}:GFP/WT and the ASY1\textsuperscript{-\textit{closure-NLS}}:GFP/WT plants in detail. Both ASY1\textsuperscript{-\textit{NLS}}:GFP variants showed a tight chromosomal association at early stage of meiotic prophase I (Supplementary Figure S1B). Subsequent chromosome spread analyses revealed that, consistent with the high level of fertility, ASY1\textsuperscript{-\textit{NLS}}:GFP/WT plants followed a regular meiotic path in which five paired bivalents were aligned on the metaphase I plate and then segregated equally at the first and second meiotic divisions, resulting in the formation of daughter cells with an equal number of chromosomes in the tetrad stage (Figure 2A). While no obvious differences were observed until zygotene compared to the ASY1\textsuperscript{-\textit{closure-NLS}}:GFP/WT plants (\(n = 98\)) (Figure 2A). Instead, only a partial synopsis was observed in ASY1\textsuperscript{-\textit{closure-NLS}}:GFP/WT plants. A defective synopsis in ASY1\textsuperscript{-\textit{closure-NLS}}:GFP/WT plants was further confirmed by the detection of ZYP1 in immunofluorescence experiments (Figure 2B). Defects in synopsis are likely the reason for the frequent observation of univalents at metaphase I (76%, \(n = 17\)) followed by unbalanced chromosome segregation at the first and second meiotic division (Figure 2A).

This is likely the reason for the high level of pollen abortion and reduced fertility (Supplementary Figure S2A and B). These results suggest that ASY1 without the closure motif dominantly interferes with the progression of synopsis.

The closure motif is required for the removal of ASY1 from chromosomes

The defective synopsis in ASY1\textsuperscript{-\textit{closure-NLS}}:GFP/WT plants together with the tight association of ASY1\textsuperscript{-\textit{closure-NLS}}:GFP with the chromosomes at late prophase I (Supplementary Figure S1B), is reminiscent of the phenotype of \textit{pch2} mutants in which the removal of ASY1 from chromosomes is compromised resulting in synopsis defects (20). Thus, we wondered whether the deletion of the closure motif might lead to a defective removal of ASY1\textsuperscript{-\textit{closure-NLS}}:GFP from synapsed chromosomes and hinder the progression of synopsis.

To answer this, we checked the presence of ASY1\textsuperscript{-\textit{closure-NLS}}:GFP at synapsed chromosomes in male meiocytes of wildtype by the co-immunolocalization analysis of ASY1 and ZYP1 using antibodies against GFP and ZYP1. The ASY1\textsuperscript{-\textit{NLS}}:GFP/WT plants were used as a control (Figure 3). (Please note that, in contrast to the previous experiments, the ASY1 signal is depicted in Figure 3 in magenta while the ZYP1 signal is shown in green.) While ASY1\textsuperscript{-\textit{NLS}}:GFP in the wild-type background was removed from the synapsed regions as normal ASY1-GFP (Figure 3A), the ASY1\textsuperscript{-\textit{closure-NLS}}:GFP signal was still not reduced at late prophase I, and its signal intensity at synapsed chromosomes appeared even stronger than that in non-synapsed regions (Figure 3B). This suggests that the presence of the closure motif is required for the efficient depletion of ASY1 when chromosomes become synapsed.

The defective synopsis in ASY1\textsuperscript{-\textit{NLS}}:GFP/WT plants might also be due to a competition between a non-functional ASY1\textsuperscript{-\textit{closure-NLS}}:GFP and the wild-type version of ASY1 for chromosomal binding in a non-preferential manner that would reduce the abundance of the functional ASY1 on chromosomes. If this were the case, the chromosomal removal of the wild-type version of ASY1 would also be likely affected by the presence of ASY1\textsuperscript{-\textit{closure-NLS}}:GFP. To test this hypothesis, we introduced ASY1\textsuperscript{-\textit{closure-NLS}} without the GFP tag into wild-type plants expressing a full-length version of the ASY1 functional reporter (PRO\textsubscript{ASY1}-ASY1:RFP, called ASY1:RFP) together with a ZYP1b reporter (PRO\textsubscript{ZYP1b}-ZYP1b:GFP, called ZYP1b:GFP), generated previously (18,24). Compared to the control plants without ASY1\textsuperscript{-\textit{closure-NLS}}, ASY1:RFP appeared to persist at synapsed regions in the plants expressing ASY1\textsuperscript{-\textit{closure-NLS}}, and the signal intensity of ASY1:RFP at synapsed chromosomes appeared even stronger than that in the non-synapsed parts reflecting the presence of ASY1\textsuperscript{-\textit{closure-NLS}}:GFP in the wildtype (Figure 3B and C). Thus, the non-functional ASY1\textsuperscript{-\textit{closure-NLS}}:GFP and the wild-type ASY1 version likely compete with each other along the chromosome axis, interfering with chromosome synopsis.

Taken together, we conclude that the closure motif of ASY1 is required for its efficient removal from the synapsed chromosomes.
Figure 2. Deletion of the closure motif dominantly interferes with synapsis. (A) Chromosome spread analysis of male meiocytes at different meiotic stages in wild-type (WT) plants harboring ASY1-NLS:GFP or ASY1Δclosure-NLS:GFP construct. Bar: 10 μm. (B) Immunolocalization of ZYP1 at different meiotic prophase I stages in wild-type plants harboring ASY1-NLS:GFP or ASY1Δclosure-NLS:GFP construct. Bar: 5 μm.

An interplay between the closure motif and PCH2 controls the nuclear targeting and chromosomal association of ASY1

Previously, we have reported that PCH2 is essential for the efficient nuclear targeting of ASY1, as seen by the accumulation of ASY1:GFP in the cytoplasm of male meiocytes of pch2 mutants (Figure 4A) (18). In contrast, we found that ASY1Δclosure-NLS:GFP in pch2 mutants localized exclusively to the nuclei of male meiocytes (Figure 4B, Table 1). To explore the epistatic relationship between the closure motif and PCH2 for the nuclear targeting of ASY1, we introduced the ASY1-NLS:GFP into pch2 mutants creating a situation in which we had two opposing forces: the promotion of nuclear entry by the NLS versus the cytoplasmic retention of the full-length ASY1 protein caused by the absence of PCH2.

We found a higher fraction of ASY1-NLS:GFP present in the nucleus of pch2 mutants compared to that of ASY1:GFP in pch2 (Figure 4A, C and D), indicating that the NLS sequence used promotes the import of a small part of ASY1-NLS:GFP into the nucleus in the pch2 mutant background. However, it is very clear that the addition of the NLS cannot fully rescue the nuclear targeting of the full-length ASY1 version in pch2 mutants and a considerable amount of ASY1-NLS:GFP was still detected in the cytoplasm of male meiocytes (Figure 4C). Thus, the presence of the NLS sequence is not sufficient to restore full nuclear localization of ASY1 in pch2 mutants.

At the same time, we noticed that ASY1-NLS:GFP showed a diffuse presence in the whole nucleus in pch2 mutants at early prophase while it was normally assembled on chromosomes in the wildtype (Figure 4C). Therefore, we reasoned that the diffuse appearance of ASY1-NLS:GFP in the nucleus of pch2 mutants is due to a fraction of ASY1-NLS:GFP proteins that are imported into the nucleus aided by the NLS, but cannot properly localize onto the chromosomes as PCH2 is absent. Thus, we conclude that in addition to the well-known role of PCH2 for ASY1 removal at late prophase, it is also indispensable for the normal chromosomal assembly of ASY1 at early meiosis. Notably, the ASY1-NLS:GFP proteins without the closure motif, i.e. ASY1Δclosure-NLS:GFP, displayed a clear chromosomal association in pch2 mutants (Figure 4B).

Taken together, these results suggest that the nuclear targeting defects of ASY1 in pch2 mutants result from the presence of the closure motif, i.e. that there is a functional interplay between the closure motif and PCH2 for the efficient nuclear localization of ASY1. In addition, a PCH2-closure motif interaction seems relevant for the chromosomal association of ASY1 at early prophase.
Figure 3. The closure motif is required for the PCH2-mediated removal of ASY1 from synapsed chromosomes. (A and B) Co-immunolocalization of ASY1-NLS::GFP (A) or ASY1 Δ closure-NLS::GFP (B) with ZYP1 in male meiocytes of wild-type plants using antibody against GFP and ZYP1. Note that for increased visibility, the brightness of the magnified panel in (B) was enhanced by 40%. (C) Co-localization of ASY1-RFP with ZYP1b::GFP in male meiocytes of wildtype (WT) and ASY1 Δ closure-NLS/WT plants using confocal microscopy. All blue and white arrows indicate the non-synapsed and synapsed regions, respectively. Bar: 5 μm.

PCH2 controls the nuclear targeting of ASY1 through regulating the HORMA domain-to-closure motif interaction

PCH2 is thought to antagonize the chromosomal association of the meiotic HORMADs, including ASY1, by driving a conformational conversion of the HORMAD proteins and regulating their interaction with the binding partners, i.e. ASY3 in Arabidopsis (15,17,19). In addition, we previously found that ASY1, at least when being expressed in yeast cells, tends to fold into a self-closed state through binding of its closure motif to its HORMA domain (18). These observations, together with the complete restoration of the nuclear localization of ASY1 without the closure motif in pch2 mutants when a NLS (ASY1 Δ closure-NLS::GFP) was added (Figure 4B, Table 1), led us to hypothesize that the interaction of the HORMA domain with the closure motif might account for the cytoplasmic retention of ASY1 when PCH2 is absent. Thus, the efficient nuclear targeting of ASY1 might require the dissociation of the HORMA
domain from the closure motif (a conformation called ‘unlocked’ state in this study) mediated by PCH2.

To test this hypothesis, we first analyzed whether the PCH2-mediated conformational change of ASY1 is necessary for its efficient nuclear targeting. The disordered N-termini of meiotic HORMADs have been found to make transient contact with PCH2, an interaction by which PCH2 partially unfolds the protein to allow for the disengagement of the HORMA domain from its interacting sequence, i.e. the closure motif (17,19,27). To this end, we generated a reporter version of ASY1 lacking the first 2–10 aa (*ProASY1*:ASY1*Δ*2–10::GFP, called *ASY1Δ*10::GFP) (Figure 1A), which theoretically should disturb the interaction of PCH2 with ASY1, and then checked the localization of ASY1*Δ*10::GFP in the wild-type background in the presence of PCH2. In support of the idea that the N-terminus of ASY1 is the action site of PCH2, the localization pattern of ASY1*Δ*10::GFP at different prophase I stages in the wild-type phenocopied that of ASY1::GFP in *pch2* mutants (Figures 4A and 5A, Table 1). This is consistent with the con-
Figure 5. The PCH2-mediated dissociation of the closure motif of ASY1 from the HORMA domain is required for its efficient nuclear targeting. (A) Localization of ASY1^Δ10:GFP and ASY1^Δ20:GFP at different meiotic stages in male meiocytes of wildtype (WT) using confocal microscopy. Bar: 10 μm. Red rectangles highlight the areas of the close-ups. (B) Yeast two-hybrid assays for the interaction of the closure motif (ASY1^571-596) with ASY1 HORMA domain (ASY1^1-300) and the domains having either the first 9 (ASY1^1-300(Δ10)) or 19 (ASY1^1-300(Δ20)) amino acid deletion at the N-terminus. (C) Yeast two-hybrid interaction assays of the closure motif with ASY1 HORMA domain and the domain harboring two non-phosphorylatable mutations ASY1^1-300(ΔT142V,ΔT184G)). (D) Localization of ASY1^T142V;T184G:GFP at late G2 and early prophase stages in male meiocytes of pch2 mutants using confocal microscopy. Bar: 10 μm. Red rectangles highlight the areas of the close-ups.

Conclusion that the PCH2-mediated conformational change of ASY1 is required for its efficient nuclear targeting. Moreover, we constructed a mutant version of PCH2 in which the glutamic acid (E283 of AT4G24710.1) of the conserved Walker B motif involved in ATP hydrolysis (28,29) was substituted with a glutamine (called PCH2^E283Q) in a functional genomic construct of PCH2 generated previously (18). Notably, we found that PCH2^E283Q introduced into pch2 mutants could not rescue the cytoplasmic retention of ASY1:GFP (Supplementary Figure S3), indicating that the ATP hydrolyzing activity of PCH2 is required for the conformational conversion of ASY1.

If ASY1 in an unlocked state goes into the nucleus more efficiently than in a self-closed state, and if the cytoplasmic retention of ASY1^Δ10:GFP in the wildtype is attributed to the tight binding of the HORMA domain to the closure motif, a disturbance of this binding should recover the nuclear targeting of ASY1^Δ10:GFP in the wildtype as well as ASY1:GFP in pch2 mutants. To test this idea, we generated different versions of ASY1 in which the binding of the
HORMA domain to the closure motif is either abolished or largely compromised.

First, considering that the N-terminal HORMA domain (15-228 amino acid for ASY1) is well-known to function as an intact unit (15,30,31), we wondered whether an additional short deletion at the N-terminus of ASY1Δ15-228:GFP would disrupt the interaction of the HORMA domain with the closure motif and thus, recover its nuclear localization in the wild-type background. To test this, we created a version of ASY1 missing the first 2–20 aa (ΔPROASY1:ASY1Δ2-20:GFP, called ASY1Δ20:GFP) (Figure 1A). Subsequently, yeast two-hybrid assays were performed for checking the binding of the HORMA domain-to-closure motif. Indeed, while ASY1Δ1-300Δ20 showed a strong interaction with the closure motif (ASY1Δ1-300), which is indistinguishable from the ASY1Δ1-300, the binding of the HORMA domain-to-closure motif was largely compromised when ASY1Δ1-300Δ20 was used for the assay (Figure 3B). Consistent with this result and the idea that a disturbance of the HORMA domain-to-closure motif binding might recover the nuclear targeting defect of ASY1Δ1-300:GFP in the wildtype, we observed that ASY1Δ1-300:GFP was localized exclusively in the nucleus of wild-type plants at late G2 and early prophase I and only a weak signal in the cytoplasm was detected at late prophase I (Figure 5A).

Second, based on our previous finding that the mutation of two presumptive CDK phosphorylation sites in ASY1 (T142 and T184) to non-phosphorylatable residues compromises the HORMA domain-to-closure motif binding (18), we generated a new non-phosphorylatable version of ASY1 harboring mutations at these two sites (ΔPROASY1:ASY1Δ142V;T184G:GFP, called ASY1Δ142V;T184G:GFP) that almost completely abolishes the interaction of the HORMA domain-to-closure motif as shown by the yeast two-hybrid assay (Figure 5C). Subsequently, the ASY1Δ142V;T184G:GFP construct was introduced into pch2 mutants (Table 1). Interestingly, we found that ASY1Δ142V;T184G:GFP was mainly localized in the nucleus of male meiocytes in pch2 mutants and only a weak signal in the cytoplasm was observed (Figure 5D), lending further support to our hypothesis that ASY1 in an unlocked state can be imported into the nucleus more efficiently.

Takentogether,theseresultssuggestthatasalongasthere is a sequence in ASY1 that binds to its HORMA domain presumably causing ASY1 to adopt a closed state, PCH2 is needed for converting ASY1 into the unlocked state and subsequent nuclear localization.

**DISCUSSION**

Meiotic HORMADs including the budding yeast Hop1, mammalian HORMAD1 and HORMAD2, C. elegans HORMADs (HTP-1, HTP-2, HTP-3 and HIM3) and Arabidopsis ASY1 play a crucial role in meiotic recombination. Their chromosomal assembly was recently suggested to depend on at least two mechanisms, the initial recruitment by its binding partners such as ASY3 in Arabidopsis, its yeast homolog Red1 and mice homolog SYCP2, and the putative self-assembly (HORMAD oligomerization) mediated by the HORMA domain-to-closure motif interaction (9,12,15,17,22,26,32). While the localization dependence of meiotic HORMADs on its binding partners, e.g. ASY3, has been experimentally proven in many organisms, the self-assembly mechanism of HORMAD proteins has remained poorly characterized (15,17). The base of the self-oligomerization mechanism is the HORMA domain-to-closure motif interaction. Here, this mechanism has been challenged, at least in Arabidopsis, since ASY1 without the closure motif was found to localize on chromosomes normally (Figure 1D and S1). In support of this finding, the point mutation K593A in the closure motif of Hop1 largely abolished the interaction of the HORMA domain with the closure motif and had no impact on the chromosomal localization of Hop1 (13,17).

Instead, our study implicated the closure motif of ASY1 in its chromosomal removal mediated by PCH2, i.e. ASY1 lacking the closure motif was constantly associated with
chromosomes and did not dissociate from synapsed chromosomes (Figure 3B). Since PCH2 acts through the N-terminus of meiotic HORMADs but not the C-terminal closure motif (Figure 5A) (19), and since ASY1 in an unlocked state is a prerequisite for its binding with ASY3 (9,15,18), we propose that the closure motif ensures the success of ASY1 removal from chromosomes through forcing ASY1 in an self-closed state in which the HORMA domain is bound by the closure motif. We propose that this self-closed state prevents the re-binding of the dissociated ASY1 to the chromosome axis via interacting with ASY3 (Figure 7).

However, this raises the question of why PCH2 does not convert the nucleoplasmic ASY1 depleted from the axis to an unlocked state at late prophase, presumably causing ASY1 to re-bind to the axis by interacting with ASY3. This question is pressing since we reveal here that PCH2 mediates the chromosomal association of ASY1 at early prophase (see below, Figure 7). The answer to this question possibly lies in the dynamic localization pattern of PCH2. Previous work from us and others show that PCH2 is first diffusely present in the nucleoplasm of male meiocytes at early prophase when ASY1 needs to be assembled on the axis. In contrast, PCH2 specifically accumulates on synapsed chromosomes and largely decreases in the nucleoplasm when the synaptonemal complex is formed at late prophase (Figure 7) (18,20). However, what determines this localization pattern of PCH2 has still to be revealed.

PCH2/TRIP13 proteins have been extensively studied with regard to their function in mediating the dissociation of meiotic HORMADs from chromosomes at late prophase (20,22,29,33). In contrast, their function at early prophase was only recently recognized (18). Here, we show that PCH2 is also indispensable for the proper chromosomal assembly of ASY1 at early prophase in Arabidopsis, as seen by the defective chromosomal association of the ASY1-NLS:GFP proteins in pch2 mutants but not in the wildtype (Figure 4C). Interestingly, the dependency of ASY1 localization on PCH2 can be eliminated by deleting the closure motif of ASY1 (Figure 4B). This suggests a PCH2-mediated conversion of ASY1 from the self-closed state to a transient unlocked state at early prophase, allowing the binding of ASY1 to ASY3 (Figure 7).

However, it seems that there are two fractions of ASY1 proteins in Arabidopsis: one fraction requires PCH2 for its nuclear targeting and chromosomal association while the other one not. This is evidenced by the strong cytoplasmic retention of some ASY1 molecules in pch2 mutants versus the clear, albeit weaker than in the wildtype, chromosomal association of some ASY1 proteins (Figure 4A). Likely, there is an equilibrium between closed and unlocked ASY1 in which the closed ASY1 is efficiently converted to the unlocked state by PCH2 for nuclear import. When PCH2 is absent, the conformational conversion of ASY1 is blocked and thus, the bulk of ASY1 stays at the closed form failing to enter the nucleus but a little portion in unlocked state that can be imported into nucleus and localizes on chromosomes. However, we cannot fully exclude that there might be other factors regulating/facilitating the nuclear import and chromosomal assembly of the PCH2-independent ASY1 fraction. One possibility is that by chance ASY1 and ASY3 may form into a complex in the cytoplasm, which is imported into the nucleus together. However, this remains to be analyzed in the future.

The necessity of PCH2 for both the chromosomal assembly and disassembly of ASY1 during meiosis largely resembles the roles of PCH2/TRIP13 orthologs in the spindle-assembly checkpoint (SAC), a mechanism ensuring that all chromosomes are attached to spindle fibers before anaphase and hence preventing aneuploidy (34–38). PCH2/TRIP13 proteins have been found to be required for not only the inactivation of SAC by disassembling the mitotic checkpoint complex (MCC) comprising BubR1, Bub3, CDC20 and the HORMA protein Mad2, but also for the establishment of the SAC by replenishing the opened Mad2 (O-Mad2) for MCC assembly (35,39–42). The findings here support a hypothesis that PCH2/TRIP13 proteins promote both the establishment and disassembly of the HORMAD signaling involved in different processes of both mitosis and meiosis in different organisms from yeast to animal and plant species. Our results support the finding that the defects in SAC activation in the absence of PCH2/TRIP13 are likely due to the deficiency of the conformational dynamics of the closed Mad2 (C-Mad2) to O-Mad2, thus abolishing MCC formation (19,42). Consistent with this assumption, the absence of PCH2/TRIP13 in C. elegans and human cells leads to the abnormal SAC activation during both mitosis and meiosis.

Figure 6. The position and origin of the closure motif is not required for its interplay with PCH2. (A) Localization of ASY1\textsuperscript{Δclosure}.GFP:closure at early prophase in male meiocytes of asy1 and pch2 mutants using confocal microscopy. Bar: 10 μm. (B) Localization of ASY1\textsuperscript{Δclosure},ASY3\textsuperscript{1-50}.NLS:GFP at early prophase in male meiocytes of pch2 mutants using confocal microscopy. Bar: 10 μm.
Figure 7. Model of the PCH2-mediated ASY1 dynamics. At early prophase, PCH2 converts the spontaneously self-closing ASY1 to a transient unlocked form by dissociating the closure motif of ASY1 from its HORMA domain allowing the nuclear import of ASY1. PCH2 is diffusely present in the nucleoplasm ensuring the formation of a pool of unlocked ASY1, which is then loaded onto chromosomes via the binding of the HORMA domain to the closure motif of ASY3 giving rise to closed and stably axis-bound form. In addition, PCH2 appears to play a role in a quality control system by releasing unstably/incorrectly bound ASY1 from the chromosome axis (18). Only one homologous chromosome is shown while the second one is indicated by cut DNA loops. At late prophase when homologous chromosomes synapse, PCH2 is recruited to the synaptonemal complex (SC) from the nucleoplasm. The SC-localized PCH2 removes ASY1 from ASY3 by disrupting the interaction of the closure motif of ASY3 with the HORMA domain of ASY1. The so released transient unlocked ASY1 immediately re-folds into a stable, closed form, thus preventing the re-binding of ASY1 to the chromosome axis.
reasoned that the inability of ASY1 without closure motif on the axis. required to dissect in depth the precise regulation of ASY1 synapsed chromosomes. Thus, further experiments are resis although later, ASY1 needs to be largely removed from function asional ASY1 could compromise early steps in synapsis. This has further demonstrated that PCH2 controls the nuclear targeting of ASY1 through dissociating the interaction between the HORMA domain and the closure motif (Figure 5). Thus, the unlocked state of ASY1 (the HORMA domain unbound by the closure motif) is a prerequisite for its nuclear import (Figure 7). Nonetheless, the underlying structural reasons remain to be understood.

The requirement of PCH2 for both the nuclear import and chromosomal assembly of ASY1 might explain the observation that no chromosomal association of PAIR2, the ASY1 homolog in rice, was detected in rice mutants of the $\textit{PCH2}$ ortholog $\textit{CRC1}$ (43). For instance, this could be due to the nuclear targeting defects of PAIR2 and/or the failure of PAIR2 to associate with the chromosome axis when CRC1 is absent. If our finding also holds true in $\textit{S. cerevisiae}$ and mammals, the failure to convert the closed into an unlocked state of Hop1 or HORMAD1/2 in $\textit{pch2}$/$\textit{trip13}$ mutants could explain the reduced number of DSBs and crossovers in these mutants resembling hop1 and homard1/2 mutants (4,5,13,33,44).

There are different, yet not mutually exclusive possibilities why ASY1 without the closure motif (ASY1$^{\text{closure-NLS:GFP}}$) dominantly interferes with synapsis. The first possibility concerns the defective elimination of ASY1 from the axis since several studies have revealed that removal of meiotic HORMADs including ASY1 is essential for achieving the full synapsis, leading to chromosome recombination and segregation defects in later stages (20,33). Thus, it seems that the compromised removal of ASY1$^{\text{closure-NLS:GFP}}$, presumably arising from the constant unlocked state, likely underlies the dominant effects on synapsis in Arabidopsis, i.e., the high abundance of ASY1 on synapsed chromosomes possibly impedes the progression of synapsis. Second, the apparently defective removal of wild-type ASY1 in the presence of ASY1$^{\text{closure-NLS:GFP}}$ being consistent with the previous scenario could also indicate a cross-talk/interdependency between these two ASY1 forms for their removal. The third possibility attributes to the likely reduced chromosomal loading of functional ASY1 due to its competition with the non-functional ASY1$^{\text{closure-NLS:GFP}}$. Possibly, this reduction of functional ASY1 could compromise early steps in synapsis although later, ASY1 needs to be largely removed from synapsed chromosomes. Thus, further experiments are required to dissect in depth the precise regulation of ASY1 dynamics on the axis.

Besides the defective removal from the chromosomes, we reasoned that the inability of ASY1 without closure motif (ASY1$^{\text{closure-NLS:GFP}}$) to complement, not even partially, the fertility and meiotic defects of $\textit{asy1}$ mutants (Supplementary Figure S2) might also result from the misregulation of the assembly/loading of components of the recombination machineries. Thus, it seems likely that the closure motif of ASY1 also plays additional roles in meiosis, e.g. promoting crossover formation. This idea is supported by the finding that hop1 cells of budding yeast harboring Hop1-K593A (a point mutation in the closure motif of Hop1) that localized to chromosomes normally, are defective in crossover formation due to the inability to properly activate the Mek1 kinase (13).

Moreover, we have tested the effects of the origin and position of the closure motif on its interplay with PCH2. Interestingly, we found that regardless of the origin and the distance between the closure motif and HORMA domain, as long as there is a HORMA-domain binding sequence in ASY1 that forces ASY1 to adopt a closed state, PCH2 is required for converting ASY1 into the unlocked state and subsequent the nuclear localization and chromosomal association (Figure 6). Thus, it seems that the presence of the closure motif mainly serves to confer ASY1 a structural plasticity essential for its functions in meiosis (Figure 7).

In this context, only the meiotic HORMADs but not the other types, e.g. Mad2 homologs, harbor the closure motifs in their own protein sequences (15). In addition, the accumulating evidence suggests that the PCH2/TRIP13-dependent conformational dynamics that allow the closure motif binding and dissociation seem to be an evolutionarily conserved mechanism of the HORMA protein signaling. Thus, we speculate that during evolution of the sexual reproduction, especially of meiosis, meiotic HORMADs might either combine the HORMA domain function and their binding regions into one protein sequence, or obtain their own closure motifs through a convergent evolution. Hence, the closure motif likely endows meiotic HORMADs with a more robust regulation with respect to the dynamic chromosomal assembly and disassembly, preventing meiotic defects and ensuring the genome stability over generations. Based on the universal existence of meiotic HORMADs and the closure motifs, together with the functional similarity of PCH2 orthologs, we postulate that the findings revealed here are also conserved in other sexually reproducing organisms.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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REFERENCES
1. Zickler, D. and Kleckner, N. (2015) Recombination, pairing, and synopsis of homologs during meiosis. Cold Spring Harb. Perspect. Biol., 7, a016626.
2. Lambing, C., Franklin,F.C.H. and Wang,C.-J.R. (2017) Understanding and manipulating meiotic recombination in plants. Plant Physiol., 173, 1530–1542.
3. Osman,K., Higgins,J.D., Sanchez-Moran,E., Armstrong,S.J. and Franklin,F.C.H. (2011) Pathways to meiotic recombination in Arabidopsis thaliana. New Phytol., 190, 523–544.
4. Daniel,K., Lange,J., Fu,J., Anastassiadis,K., Roig,I., Cooke,H.J., Stewart,A.F., Wassmann,K., Jasim,M. et al. (2011) Meiotic homologue alignment and its quality surveillance are controlled by mouse HORMAD1. Nature Publishing Group, 13, 599–610.
5. Wojtasz,L., Cloutier,J.M., Baumann,M., Daniel,K., Varga,J., Fu,J., Anastassiadis,K., Stewart,A.F., Remyani,Y., Turner,J.M.A. et al. (2012) Meiotic DNA double-strand breaks and chromosome asynapsis in mice are monitored by distinct HORMAD2-independent and -dependent mechanisms. Genes Dev., 26, 958–973.
6. Hollingsworth,N.M., Goetsch,L. and Byers,B. (1990) The HOP1 gene encodes a meiosis-specific component of yeast chromosomes. Cell, 61, 73–84.
7. Armstrong,S.J. (2002) Asy1, a protein required for meiotic chromosome synopsis, localizes to axis-associated chromatin in Arabidopsis and Brassica. J. Cell Sci., 115, 3645–3655.
8. Chambon,A., West,A., Vezon,D., Harashima,H., Stolze,S.C., Yen,T.J., Liu,S.T. and Hershko,A. (2014) Disassembly of the HORMAD2 checkpoint-silencing protein. Chromosoma, 123, 279–292.
9. Yen,T.J., Kim,D.H., Dereli,I., Rosenberg,S.C., Hagemann,G., Herzog,F., Töth,A., Cleveland,D.W. and Corbett,K.D. (2017) The AAA+ ATPase TRIP13 remodels HORMA domains through N-terminal engagement and unfolding. EMBO J., 36, 2419–2434.
10. Yang,C., Sofroni,K., Wijnker,E., Hamamura,Y., Carstens,L., Harashima,H., Stolze,S.C., Vezon,D., Chelysheva,L., Orban Nemeth,Z. et al. (2019) The Arabidopsis Cdk1/Cdk2 homolog CDKA1 controls chromosome axis assembly during plant meiosis. EMBO J., 38, 685–698.
11. Ye,Q., Kim,D.H., Dereli,I., Rosenberg,S.C., Hagemann,G., Herzog,F., Töth,A., Cleveland,D.W. and Corbett,K.D. (2017) The AAA+ ATPase TRIP13 remodels HORMA domains through N-terminal engagement and unfolding. EMBO J., 36, 2419–2434.
12. Yang,C., Sofroni,K., Wijnker,E., Hamamura,Y., Carstens,L., Harashima,H., Stolze,S.C., Vezon,D., Chelysheva,L., Orban Nemeth,Z. et al. (2019) The Arabidopsis Cdk1/Cdk2 homolog CDKA1 controls chromosome axis assembly during plant meiosis. EMBO J., 38, 685–698.
13. Ye,Q., Kim,D.H., Dereli,I., Rosenberg,S.C., Hagemann,G., Herzog,F., Töth,A., Cleveland,D.W. and Corbett,K.D. (2017) The AAA+ ATPase TRIP13 remodels HORMA domains through N-terminal engagement and unfolding. EMBO J., 36, 2419–2434.
mitotic checkpoint complexes by the joint action of the AAA-ATPase TRIP13 and p31comet. Proc. Natl Acad. Sci. U.S.A., 111, 12019–12024.

39. Ma, H. T. and Poon, R. Y. C. (2018) TRIP13 functions in the establishment of the spindle assembly checkpoint by replenishing O-MAD2. Cell Rep., 22, 1439–1450.

40. Eytan, E., Wang, K., Miniowitz-Shemtov, S., Sitry-Shevah, D., Kaisari, S., Yen, T. J., Liu, S. T. and Hershko, A. (2014) Disassembly of mitotic checkpoint complexes by the joint action of the AAA-ATPase TRIP13 and p31comet. Proc. Natl Acad. Sci. U.S.A., 111, 12019–12024.

41. Nelson, C. R., Hwang, T., Chen, P.-H. and Bhalla, N. (2015) TRIP13PCH-2 promotes Mad2 localization to unattached kinetochores in the spindle checkpoint response. J. Cell Biol., 211, 503–516.

42. Kim, D. H., Han, J. S., Ly, P., Ye, Q., McMahon, M. A., Myung, K., Corbett, K. D. and Cleveland, D. W. (2018) TRIP13 and APC15 drive mitotic exit by turnover of interphase- and unattached kinetochore-produced MCC. Nat. Commun., 9, 4354.

43. Miao, C., Tang, D., Zhang, H., Wang, M., Li, Y., Tang, S., Yu, H., Gu, M. and Cheng, Z. (2013) CENTRAL REGION COMPONENT1, a novel synaptonemal complex component, is essential for meiotic recombination initiation in rice. Plant Cell, 25, 2998–3009.

44. Joshi, N., Barot, A., Jamison, C. and Börner, G. V. (2009) Pch2 links chromosome axis remodeling at future crossover sites and crossover distribution during yeast meiosis. PLoS Genet., 5, e1000557.