Transcriptional gene silencing requires dedicated interaction between HP1 protein Chp2 and chromatin remodeler Mit1

Karoline Leopold,1 Alessandro Stirpe,1 and Thomas Schalch1,2

1Department of Molecular Biology, Faculty of Science, Sciences III, University of Geneva, CH-1211 Geneva 4, Switzerland; 2Leicester Institute for Structural and Chemical Biology, Department of Molecular and Cell Biology, University of Leicester, Leicester LE1 9HN, United Kingdom

Heterochromatin protein 1 (HP1) proteins are key factors of eukaryotic heterochromatin that coordinate chromatin compaction and transcriptional gene silencing. Through their multivalency they act as adaptors between histone H3 Lys9 di/trimethyl marks in chromatin and effector complexes that bind to the HP1 chromoshadow domain. Most organisms encode for multiple HP1 isoforms and the molecular mechanisms that underpin their diverse functions in genome regulation remain poorly understood. In fission yeast, the two HP1 proteins Chp2 and Swi6 assume distinct roles and Chp2 is tightly associated with the nucleosome remodeling and deacetylation complex SHREC. Here we show that Chp2 directly engages the SHREC nucleosome remodeler subunit Mit1. The crystal structure of the interaction interface reveals an extraordinarily extensive and specific interaction between the chromoshadow domain of Chp2 and the N terminus of Mit1. The integrity of this interface is critical for high affinity binding and for heterochromatin formation. Comparison with Swi6 shows that the Chp2–Mit1 interface is highly selective and thereby provides the molecular basis for the functional specialization of an HP1 isoform.

[Keywords: heterochromatin; nucleosome remodeling; S. pombe; HP1 proteins; recruitment; X-ray crystallography; protein interactions]

Supplemental material is available for this article.

Received September 5, 2018; revised version accepted February 5, 2019.
Most organisms have multiple HP1 isoforms including mammals (HP1α, HP1β, and HP1γ), *Xenopus laevis* (HP1α and HP1γ), *Drosophila* [HP1, HP1b, and HP1c, *Rhino*, HP1E], *Caenorhabditis elegans* [HPL-1 and HPL-2], and fission yeast [Swi6 and Chp2] [Vermaak et al. 2005; Lomberk et al. 2006]. Despite their similar architecture and sequence they are remarkably different. For example, different HP1 isoforms associate with different sets of client proteins [Motamedi et al. 2008; Fischer et al. 2009; Vermaak et al. 2005; Job et al. 2016]. The different isoforms also often have nonoverlapping roles in genome regulation [Minic et al. 1999; Smothers and Henikoff 2001; Vacic et al. 2005], and it remains poorly understood how diversification of function is established since they bind the same histone mark and the same motif in client proteins. Here we investigate the mechanism underlying HP1 diversification in the fission yeast *Schizosaccharomyces pombe*, where the nonoverlapping functions of the two HP1 isoforms in heterochromatin formation is well established.

Gene silencing in the *S. pombe* system relies on an interplay between RNA interference (RNAi) and chromatin-associated processes [Martienssen and Maizel 2015]. Small RNAs are produced from pericentromeritic heterochromatin loci and guide the RNAi-induced transcriptional gene silencing complex (RITS) to nascent transcripts, which are subsequently degraded by the RNAi machinery [Volpe et al. 2002; Verdel et al. 2004; Shimada et al. 2016]. Concomitantly, RITS recruits the H3K9 methyltransferase Clr4 to heterochromatic loci [Zhang et al. 2008; Bayne et al. 2010], and deposition of the H3K9 methyl marks provides chromatin binding sites for the HP1 proteins Swi6 and Chp2. These proteins cannot complement each other’s function in the regulation of transcription at heterochromatic loci [Sadaie et al. 2008], and they are found to be part of nonoverlapping complexes [Motamedi et al. 2008; Fischer et al. 2009]. While Swi6 is expressed at relatively high levels, Chp2’s expression levels are much lower [Sadaie et al. 2008]. In contrast to Swi6, which binds a wide range of client proteins, Chp2 is predominantly associated with the Snf2/HDAC repressor complex (SHREC) that belongs to the family of nucleosome remodeling and deacetylation complexes (NuRDs).

SHREC consists of the nucleosome remodeler Mit1, the histone deacetylase Clr3 and the MBD-like protein Clr2, which are all connected by the scaffolding subunit Clr1 [Sugiyama et al. 2007; Motamedi et al. 2008; Fischer et al. 2009; Job et al. 2016]. The previous work has also established a close functional and biochemical relationship between Chp2 and the Mit1 remodeler, which recruits the SHREC complex to heterochromatin. However, how Chp2 achieves the specific recruitment of the SHREC complex is not understood.

Here we show that Chp2 binds the N terminus of Mit1, and that an extensive interface between Chp2 and Mit1 provides a high-affinity interaction that is required for recruitment of Mit1 to heterochromatin and silencing of gene expression. Our structural and functional analysis provides insight into how an isoform-specific HP1 complex forms and how it contributes to heterochromatin function.

### Results

**Chp2 interacts with the N terminus of Mit1 to repress transcription**

Chp2 recruits the SHREC complex to elicit transcriptional gene silencing in heterochromatic regions [Motamedi et al. 2008; Fischer et al. 2009]. These mass spectrometry experiments suggested that the interaction occurs between Chp2 and either the SHREC scaffold Clr1 or the chromatin remodeler Mit1. To identify the molecular interface that underlies Chp2-mediated recruitment of SHREC we chose insect-cell based coexpression of Chp2 with Mit1 or Clr1. Because pull-down experiments of Chp2 with fragments of the Clr1 N terminus did not yield a complex, we focused on the chromatin remodeler Mit1 and its individual domains (Fig. 1A). Coexpression of a tertiary complex consisting of Chp2, Mit1, and Clr1’s Mit1 interaction domain (MID) yielded a stable biochemical entity corresponding to the remodeler module of SHREC [Supplemental Fig. S1A]. By testing various domain deletions of Mit1 for complex formation we found that the N terminus of Mit1, which does not harbor any known domains, was necessary and sufficient for mediating the Chp2 interaction in the heterologous insect cell system. Deletion of residues 61–200, which are predicted to be unstructured, lead to loss of the interaction, while the mutation of two LxVxL motifs in this region [Mit1V210F, Mit1V272F] [Supplemental Fig. S1A, lane N2F] has no effect on complex formation. These results suggested that Mit1 and Chp2 might deviate from a canonical PxVxL-mediated interaction. To confirm and further characterize complex formation we subjected Mit1 and Chp2 to a yeast-two-hybrid assay [Supplemental Fig. S1B]. The yeast-two-hybrid results showed that the Mit1 N terminus and the Chp2CSD are each necessary and sufficient for interaction. Under stringent selection on quadruple drop-out medium (QDO) Chp2 full length protein is required to sustain growth. This raises the possibility that Mit1 interacts with the CSD as well as the CD, the hinge region or the N terminus of Chp2.

To investigate the role of the Mit1 N terminus under physiological conditions we endogenously tagged Mit1 with a C-terminal 13-myc tag and deleted the N-terminal 200 amino acids [mit1ΔN] [Supplemental Fig. S1C]. Coimmunoprecipitation experiments in a 6xFlag-Chp2 background showed complete loss of Chp2 interaction in the mit1ΔN mutant [Fig. 1B, Supplemental Fig. S1D]. We therefore conclude that Chp2 forms a complex with Mit1 by binding to its N-terminal domain.

We next subjected the mit1ΔN strain to comparative growth assays in an *otr1R:ura4* background that allows assessment of heterochromatin integrity in the pericentromeric region of chromosome 1 in order to test the functional relevance of the Chp2–Mit1 interaction. In a *ura4Δ* background the expression levels of the *ura4* marker genes can be monitored by increased growth of cells on medium lacking uracil. In contrast, medium containing the drug 5-fluorouracil (FOA) inhibits growth of cells that fail to silence *ura4*, because its gene product converts FOA into...
fluorodeoxyuridine, which is toxic to the cells. Similar to a complete deletion of mit1+, deletion of the Mit1 N terminus leads to loss of silencing of the otr1R::ura4 reporter (Fig. 1C). Thus, full length Mit1 is critical for heterochromatin formation. To analyze the function of the Mit1 N terminus at endogenous loci we measured centromeric and telomeric transcript levels by RT-qPCR and compared them to wild-type cells (Fig. 1D). We found five to 15-fold elevated levels of centromeric transcripts in the mit1ΔN mutant when compared to wild type. This is similar to the changes observed at the otr1R::ura4 reporter (Fig. 1C). Even though the myc-tagged wild-type allele shows a small defect at the otr1R::ura4 reporter it is fully functional at endogenous loci. As expected from previous observations (Sugiyama et al. 2007; Motamedi et al. 2008), the centromeric transcript changes for mit1 mutants are moderate when compared to clr4Δ (Supplemental Fig. S1E). In our experiments telomeric transcripts show a stronger relative change in silencing between wild-type and mutant loci than the centromeric repeats, which might be due to the extremely low levels of transcripts in wild-type cells and consequently large fold-changes upon loss of silencing. Nevertheless, at all loci heterochromatic transcript levels in mit1ΔN cells are very similar to the ones observed for chp2Δ or mit1Δ. These results suggest that an intact physical connection between Chp2 and Mit1 is essential for mediating their function in heterochromatin formation.

Crystal structure of the Chp2–Mit1 interface reveals extensive interaction

To understand how Chp2 recognizes the N terminus of Mit1 we decided to determine the atomic structure of the interaction interface. Limited proteolysis of a complex consisting of the Chp2CSD and the N-terminal 300 residues of Mit1 revealed a protected Mit1 fragment corresponding to residues 1–81, which we will refer to as the Chp2 interaction interface (CII) (Fig. 2A; Supplemental Fig. S2A). This Mit1 fragment formed a stable complex with Chp2 (Supplemental Fig. S2B), which readily crystallized and allowed us to determine the structure by molecular replacement at a resolution of 1.6 Å (Table 1). Electron density is well defined for the two CSD copies of Chp2, and we observed an equally well defined density for residues 8–81 of the Mit1CII (Supplemental Fig. S2C).

The structure of this minimal Chp2–Mit1 complex revealed a 2:1 stoichiometry with one Mit1 molecule bound to one Chp2CSD dimer (Fig. 2B). Mit1CII occupies the canonical peptide-binding groove at the CSD dimerization interface and, intriguingly, it wraps around...
one of the Chp2CSDs and engages an extensive interaction interface. This structure allows for the unambiguous identification of residues 9–13 of Mit1 in the CSD dimerization groove. These residues correspond to the sequence CKIVV with I11 occupying the central position on the symmetry axis of the CSD dimer [position 0 when numbered according to Thiru et al. 2004] [Fig. 2C]. The central I11 as well as C9 at position −2 and V13 at position +2 are buried inside hydrophobic pockets at the bottom of the binding groove, while K and V at positions ±1 are exposed to the solvent. The binding groove is lined by two symmetric sets of residues comprising Y372, Y373, H376, I377, and F379 from both Chp2 protomers. The central I11 is a large hydrophobic side chain, and when compared to HP1-CAF1 [Thiru et al. 2004], a canonical HP1-PxVxL complex, Chp2 accommodates the extra bulk of the isoleucine by a side chain rotation of Y373 such that the aromatic face of the tyrosine residue packs against the methyl groups of I11 [Supplemental Fig. S2D]. The outer pocket where the side chains of C9 and V13 are found is formed by Y372 and H376 from one protomer and F379 and I377 from the other protomer. This pocket is relatively spacious and easily accommodates the cysteine and valine residues, but without strong complementarity or formation of specific hydrogen bonds. Thus, the structure suggests that the Chp2–Mit1 equivalent of the HP1–PxVxL interaction is a CkIvV motif that binds in a well-defined manner to the Chp2CSD dimer interface due to spacious hydrophobic binding pockets and the hydrogen bonds established by β-sheet formation.

**Figure 2.** Crystal structure of the Chp2–Mit1 complex reveals extensive interface. (A) Schematic of the minimal complex between the N-terminal 81 residues of Mit1 and the CSDs of Chp2 based on limited proteolysis [Supplemental Fig. S1A]. (B) Surface and cartoon representation of the Chp2–Mit1 crystal structure. Eye symbols with letters indicate viewing angles for corresponding details panels C–F. (C–F) Close-up views of the Chp2–Mit1 crystal structure colored as in (B). (C) Binding of the CkIvV motif to the groove formed by the Chp2CSD dimer. (D) Hydrophobic interactions of the linker region of Mit1 with the surface of Chp2CSD2. (E) Hydrogen bonding network between Mit1 linker domain and Chp2CSD2. (F) Water-mediated Mit1CDL–Chp2CSD2 interaction interface. (G) Superposition of the Mit1CDL domain with the CD of Chp1 bound to a H3K9 trimethyl peptide [PDBID: 5G7L, RMSD = 1.38 Å].

The Chp2–Mit interface includes a cryptic Mit1 domain with CD fold

The crystal structure reveals that the interface between Mit1 and Chp2 includes extensive interactions that go far beyond the classical PxVxL binding groove at the CSD dimer interface. These additional interactions are provided by residues downstream of the CkIvV motif, which are bound to the surface of Chp2CSD2 [Fig. 2B]. The first stretch of residues, which we refer to as the linker region, assumes an extended configuration and corresponds to Mit1 positions 15–31. It contains a short 310
helix formed by a hydrophobic stretch between residues 18–22. The three leucines on this helix all fit complementary pockets in the Chp2 surface [Fig. 2D]. The structure also shows that the V20 of the previously tested LxVxL motifs [Supplemental Fig. S1A] is on the surface and can accommodate a mutation to phenylalanine comfortably (the same is true for V72). After these hydrophobic interactions follows a series of polar and acidic residues, which form a hydrogen-bonding network with residues H356 and D357 of Chp2 [Fig. 2E, Supplemental Fig. S2E,F]. Chp2H356 engages the side chain of Mit1E29 in a bifurcated hydrogen bond and Chp2D357 establishes one hydrogen bond with Mit1Y25 and one with Mit1T30. Due to these hydrophobic and polar interactions the entire Mit1 linker region is tightly associated with the Chp2 surface and is therefore expected to play a significant role in mediating the interaction between Chp2 and Mit1.

Table 1. Crystallographic table

| Parameter | Native |
|-----------|--------|
| Wavelength (Å) | 1.000 |
| Resolution range (Å) | 46.50–1.60 (1.66–1.60) |
| Space group | P31,21 |
| Unit cell parameters (Å, °) | a = 58.1, b = 58.1, c = 139.5, α = β = 90, γ = 120 |
| Total reflections | 704923 (62573) |
| Unique reflections | 36819 (3611) |
| Multiplicity | 19.1 (17.3) |
| Completeness (%) | 99.9 (99.9) |
| Mean I/σ(I) | 17.6 (3.9) |
| Wilson B factor (Å²) | 21.8 |
| Rmerge (%) | 11.2 (79.4) |
| Rfree (%) | 11.5 (81.0) |
| CC (work) | 0.97 (0.90) |
| CC (free) | 0.97 (0.81) |
| Number of reflections in test set | 1893 (189) |
| Resolution range | 46.50–1.60 (1.66–1.60) |
| Total number of reflections | 36801 (3612) |
| Number of protein residues | 207 |
| Macromolecule | 1715 |
| Ligands | 16 |
| Solvent | 151 |
| Number of protein residues | 207 |
| RMSD, bonds (Å) | 0.010 |
| RMSD, angles (°) | 1.00 |
| Ramachandran favored (%) | 98.5 |
| Ramachandran outliers (%) | 0.0 |
| Ramachandran allowed (%) | 1.5 |
| Clash score | 1.7 |
| Average B factor (Å²) | 32.2 |
| Macromolecule | 31.9 |
| Ligands | 33.0 |
| Solvent | 35.3 |

*Statistics for the highest-resolution shell are shown in parentheses.

The linker region is followed by a folded domain, easily recognizable as a CD fold with close resemblance to the Chp1 CD [Fig. 2G; Schalch et al. 2009]. The domain assumes the typical arrangement of a warped three-stranded β sheet that packs against a C-terminal α-helix. However, the Mit1 CD-like (CDL) domain lacks an aromatic cage and the binding groove for a methyl-lysine histone peptide. Instead, the peptide binding groove of the domain is occupied by Chp2CSD2 [Fig. 2F,G], with Chp2I360 occupying the space of the H3K9 trimethyl group when compared with Chp1.

The contacts between Chp2 and the Mit1CDL domain extend the interface established by the hydrogen bonding network in the linker region [Fig. S2F]. Key residues in the CDL are Mit1K62 and Mit1Y63. The Mit1K62 sidechain is involved in a hydrogen bond to the backbone oxygen of Chp2N339 and reciprocally the side chain group of Chp2K341 hydrogen-bonds to the backbone oxygen of Mit1K62. Mit1Y63 is involved in a water-mediated hydrogen-bonding network that connects it to backbone atoms of Chp2N358, Chp2I359, and Chp2L342 [Fig. 2F]. Furthermore, we found hydrophobic contacts between Chp2F319 and Mit1A49 at the edge of the CDL, as well as in the hydrophobic core of the Mit1CDL–Chp2CD interface that is formed by Mit1F47, Mit1V51, Mit1A53, Chp2I360, and Chp2I359 [Supplemental Fig. S2F].

The Chp2–Mit1 structure reveals an extensive interface that buries 3354 Å² of surface area involving the CkIvV motif, linker region and CDL domain [CkIvV motif: 1465 Å², Linker: 1131 Å², CDL: 750 Å²]. Each of them provides a significant number of hydrogen bonds and hydrophobic interactions. In contrast to the CkIvV motif, which displays little side-chain specificity, the Mit1 linker and CDL domain engage in numerous hydrogen-bonding interactions. The structure predicts that all three parts are required for full binding affinity and that linker and CDL provide the extra specificity of Mit1 for the Chp2CSD.

The extended Chp2–Mit1 interface is required for high affinity interaction

To characterize the biophysical properties of the Chp2–Mit1 interaction we subjected the complex to analysis by isothermal calorimetry (ITC) measurements. Figure 3A shows that combination of wild-type Mit1CDL and Chp2CD results in a exothermic binding reaction, revealing a tight interaction between Chp2 and Mit1 with a Kd of 2.6 nM [Table 2]. As Chp2 is predicted to bind canonical PxVxL motifs [Thiru et al. 2004] we tested binding of a mutant Mit1 with engineered PkVvL motif instead of CkIvV, and obtained a Kd of 8.2 nM. These data show that despite its nonconsensus sequence the CkIvV motif binds at least equally well to the Chp2CD dimer as a canonical PxVxL motif.

We decided to mutate the central isoleucine to arginine to disrupt CkIvV binding to the dimerization groove and assess its contribution the Chp2–Mit1 interaction. The Mit1I11R mutation in the CkIvV motif lead to a strong

GENES & DEVELOPMENT 5
reduction in observable binding energy (Table 2, Fig. 3A) and resulted in a \( K_d \) of 21.5 \( \mu \text{M} \), three to four orders of magnitude higher than the intact interface. A smaller drop in affinity to 2.2 \( \mu \text{M} \) is observed for the removal of the CDL (Table 2, Fig. 3A). Thus, CDL as well as CktvV-mediated interactions make critically important contributions to the Chp2–Mit1 interaction interface. We conclude that the extensive contact surface observed in the crystal

![Figure 3](image_url)

**Figure 3.** Extended interaction between Chp2 and Mit1 is required for heterochromatin silencing. (A) ITC heat rates and fits for titration of Chp2CSD with StrepSumo-tagged Mit1CII wild-type and indicated mutant proteins. Open circles represent StrepSumo-tag only control. See Table 2 for fitted parameters. (B) Comparative growth assays for Mit1 mutants in an \( \text{otr1R::ura4}^+ \) background with the corresponding transcript levels of the \( \text{otr1R::ura4} \) reporter displayed as horizontal bars measured by quantitative real-time RT-PCR. RT-qPCR levels for wild type, \( \text{mit1-13myc} \), \( \Delta \text{chp2} \), and \( \Delta \text{mit1} \) from Figure 1D are shown for comparison. (C) Changes in steady-state transcript levels in mutant strains relative to wild-type cells were determined by RT-qPCR for centromeric \( \text{dg/dh} \) repeats and \( \text{tlh1} \) transcripts at telomeres. \( \text{mit1A} \) from Figure 1D is shown for comparison. (D) Chromatin immunoprecipitation (ChIP) for indicated mutant strains against Mit1-13myc, the H3K14ac histone mark and for RNA polymerase II. \( \text{act1} \) was used as internal standard for all measurements and standard errors were calculated from three independent biological experiments.
structure is required for providing high-affinity binding between Chp2 and Mit1.

**Disturbing the Chp2–Mit1 interaction abolishes transcriptional gene silencing**

To test the functional significance of an intact Chp2–Mit1 interface we replaced the endogenous copy of Mit1 with Mit1 mutant versions targeting the CkIvV motif, the linker region and the CDL domain. Subjecting these strains to growth assays on selective medium revealed that disruptive mutations in any part of the Chp2–Mit1 interface lead to alleviated gene silencing comparable to the deletion of Chp2 or to the deletion of the Mit1 N terminus [Fig. 3B; Supplemental Fig. S3]. Furthermore, we observed that heterochromatic transcript levels in selected mutants increased to similar levels as observed in mit1ΔN (Fig. 3B,C). These results further highlight the importance of an intact interface between Chp2 and Mit1 for appropriate gene silencing in heterochromatic regions.

The silencing assays, together with the structural and biophysical data, suggest that recruitment of Mit1 to heterochromatin is severely impaired when the Chp2–Mit1 interface is missing or partially disrupted. We therefore used chromatin immunoprecipitation [ChIP] at dh repeats against mit1-13myc wild-type and mit1-13myc interface mutants to test Mit1 recruitment to heterochromatin directly [Fig. 3D]. These experiments show a complete loss of Mit1 upon deletion of the complete Mit1CII or upon disruption of the CkIvV motif by the I11R mutation, demonstrating that Mit1 association with heterochromatin sequences relies critically on the intact Chp2–Mit1 interaction. ChIP against the histone H3K14 acetyl mark and against RNA polymerase II reveals that loss of Mit1 correlates with the accumulation of H3K14ac and increased RNA polymerase II occupancy. Thus, disrupting the Chp2–Mit1 interface leads to the same loss of transcriptional gene silencing at heterochromatic repeats as observed for SHREC mutants [Sugiyama et al. 2007; Motamedi et al. 2008].

**Swi6 binds the Mit1 CkIvV motif, but lacks further interaction surface**

Several lines of evidence have implicated Swi6 in SHREC recruitment, but the mechanisms of this recruitment have remained elusive [Yamada et al. 2005; Sadaie et al. 2008; Fischer et al. 2009]. Since Chp2 shows strong affinity for the PxVxL substitution it is conceivable that reversely Swi6 is able to bind the CkIvV motif. Indeed, we obtained a binding curve with a $K_d$ of 28 µM for Swi6CSD binding to wild type Mit1CII [Table 2; Fig. 4A]. Binding to the Mit1 PxVxL substitution mutant shows a tighter interaction with a $K_d$ of 15 µM, while the Mit1 I11R mutation completely abolished the interaction and thereby corroborates the CkIvV motif as the site of interaction with Swi6. These observations reveal a complex interplay between the two HP1 proteins for Mit1, where a high-affinity interaction provides specificity for Chp2, while Swi6 has the potential to bind to the same region with low affinity.

The reason for Swi6’s inability to bind Mit1 with high affinity can be rationalized by comparing the Chp2CSD dimer in the Chp2–Mit1 complex structure with the crystal structure of the free Swi6CSD dimer [Cowieson et al. 2000]. They superimpose with a RMSD of 0.85 Å for Ca atoms and confirm the expected very close structural similarity [Supplemental Fig. S4A]. Nevertheless, a striking difference between the two structures is observed in the folding of the N terminus where Chp2 shows seven structured residues more than Swi6, which form a loop that packs against the C-terminal β-strand and the following a-helix and thereby provide significantly more bulk to the Chp2 CDS [Fig. 4B]. We also observe a molecule of hexanediol that is deeply buried by the Chp2 N terminus, and we speculate that it substitutes for one or two N-terminal residues by which our CSD construct is too short. The particular folding of the N terminus is likely to be independent of Mit1 binding, since it is identical between both CSDs even though only one is bound to Mit1 [Supplemental Fig. S4B]. It is however noteworthy that the trajectory of the N-terminal loop differs around residue Q320 by a shift

### Table 2. ITC data

| Cell          | Syringe       | ΔH [kcal mol⁻¹] | ΔS [cal mol⁻¹ K⁻¹] | $K_a$ [M⁻¹] | $K_d$ [M⁻¹] | N   | Figure |
|---------------|---------------|-----------------|--------------------|-------------|-------------|-----|--------|
| Chp2CSD dimer | Mit1CIIb,c     | −15.7 ± 0.1     | −13.8              | 3.9 ± 0.6 × 10⁸ | 2.6 ± 0.4 × 10⁻⁹ | 1.1 | 3⁴     |
| Chp2CSD dimer | Mit1CII PkVvlb,c | −10.2 ± 0.1  | 2.5                | 1.2 ± 0.2 × 10⁸ | 8.2 ± 1.6 × 10⁻⁹ | 0.9 | 3⁴     |
| Mit1CII I11R | Chp2CSD dimer | −11.5 ± 3.9     | −17.3              | 4.7 ± 0.9 × 10⁴ | 21.5 ± 3.9 × 10⁻⁶ | 0.5 | 3⁴     |
| Mit1CII ΔCDL | Chp2CSD dimer | −4.0 ± 0.3      | −4.0               | 45.5 ± 16 × 10⁴ | 2.2 ± 0.8 × 10⁻⁶ | 0.8 | 3⁵     |
| StrepSumo tag | Chp2CSD dimer | NA              | NA                 | NA          | NA          | NA  | NA     |
| Mit1CIIb      | Swi6CSD dimer | −8.0 ± 0.8      | −6.3               | 3.5 ± 0.3 × 10⁴ | 28.4 ± 2.4 × 10⁻⁶ | 0.8 | 4⁴     |
| Mit1CII PkVvlb | Swi6CSD dimer | −4.2 ± 0.2      | 7.8                | 6.5 ± 0.6 × 10⁴ | 15.3 ± 1.4 × 10⁻⁶ | 1.3 | 4⁴     |
| Mit1CII I11R | Swi6CSD dimer | NA              | NA                 | NA          | NA          | NA  | NA     |

*Untagged, residues 314–380, dimer concentration.

*bStrep-sumo tagged.

*cFor Chp2CSD vs. Mit1CII and Mit1CII PkVvl, Mit1 was injected from the syringe. For the other experiments, Chp2CSD or Swi6CSD was injected from the syringe due to practical limitations in obtaining the high Mit1CII concentrations that are required to measure the weaker interactions.

*dMeasured on VP-ITC (Microcal).

*eMeasured on NanoITC (TA Instruments).

fUntagged, residues 261–328.
of up to 1.5 Å, which is probably due to the methyl-π interaction of Chp2F319 with Mit1A49 (Supplemental Fig. S4B). The Chp2 N terminus thus provides a unique binding surface for Mit1 that contributes to extension of the interaction interface when compared with Swi6.

Superposition of the surfaces of Chp2 and Swi6 also shows that the short helix in the linker region of Mit1 around residues 18–24 would collide with the surface of Swi6 [Fig. 4B]. Furthermore, this surface exposes a hydrophobic patch in Chp2, while Swi6 forms an intramolecular salt bridge in the equivalent position, which disfavors accommodation of the hydrophobic stretch of Mit1 residues from L18 to L22. When analyzed in detail it is clear that the Swi6 sequence differs from Chp2 in several other key positions that are important for binding Mit1. These differences are conserved between Chp2 and Swi6 proteins of the Schizosaccharomyces family [Supplemental Fig. S4C]. Conversely, the N-terminal domain architecture of Mit1 proteins is also conserved, particularly in key hydrogen bonding positions like Y25 [Supplemental Fig. S4D], supporting the notion that the Chp2–Mit1 interaction is conserved across the Schizosaccharomyces clade. Despite the differences between the Chp2 and Swi6 proteins, the canonical binding groove formed by the CSD dimerization interface of Swi6 can easily accommodate the CkIvV motif of Mit1 [Fig. 4C], which is entirely consistent with the ITC data.

Discussion

The unique structure of an HP1–client interface

The results presented here establish the mechanistic details of how the HP1 protein Chp2 interacts with the chromatin remodeler Mit1. The unique interaction between the two proteins combines binding in the canonical PxVxL binding groove at the dimerization interface of HP1 proteins with an extensive additional interaction interface created by wrapping the N-terminal domain of Mit1 around one of the Chp2CSDs. This elaborate
interaction results in a low nanomolar dissociation constant and renders the Chp2–Mit1 interface highly specific, thereby explaining the exclusive client selectivity of Chp2 observed in previous experiments (Motamedi et al. 2008; Fischer et al. 2009), and why Chp2 is a de facto constitutive subunit of SHREC. The dedicated interface thereby uncouples SHREC’s recruitment to heterochromatin from Swi6 and enables the cell to specifically regulate SHREC levels at heterochromatin loci through fine tuning expression levels of Chp2 [Fig. 4C].

The gain in specificity can be attributed to sequence differences in key surface residues that are involved in shape complementarity and hydrogen bonding networks between Chp2 and Mit1. The extended interaction interface is provided by the linker and the well folded CDL domain in Mit1, which has been missed so far by sequence comparison. It clearly resembles a CD and demonstrates a further use of this versatile fold in protein–protein interactions. The yeast-two-hybrid results suggest involvement of Chp2 domains other than the CSD. To our knowledge, the currently available evidence does not rule out the possibility of an even more extensive interface between Mit1 and Chp2.

With the CkIvV sequence bound to the CSD dimer interface the Chp2–Mit1 structure reveals a remarkably degenerate equivalent of the PxVxL motif that is not recognized by motif searches. Sequence comparison of Mit1 in Schizosaccharomyces species indicates that the cysteine residue in the −2 position is poorly conserved and that it can accommodate lysine and arginine (Supplemental Fig. S4D). The 0 and +2 positions on the other hand are confined to the hydrophobic side chains valine or isoleucine. This suggests that the motif in the Mit1 context corresponds to xIVxIV. Interestingly, Swi6 is also able to bind this motif, though with an affinity that is weaker than for bona fide Swi6 clients [Isaac et al. 2017] or the Mit1 PxVxL substitution.

Regulation of Mit1

Our findings are consistent with previous observations that Mit1 and Chp2 are closely associated both functionally and physically [Motamedi et al. 2008; Fischer et al. 2009; Job et al. 2016]. The identification of the physical interface between Chp2 and Mit1 fleshes out the connectivity of subunits in the SHREC complex and establishes the nucleosome remodeler Mit1 as the connecting subunit between Chp2 and Clr1, which interacts with the C-terminal domain of Mit1 [Fig. 4D]. We further demonstrate that the repressive function of Mit1 and Chp2 depends strongly on their high-affinity interaction, which serves to recruit Mit1 to heterochromatin loci. Mit1 also harbors a PHD domain that binds histone H3 and potentially activating marks [Creamer et al. 2014], and it also interacts with Clr1, which connects Mit1 and Chp2 to the HDAC module of SHREC [Job et al. 2016]. It is likely to be an equilibrium between these factors that controls Mit1’s activity in transcriptional gene silencing. The identification of the Chp2–Mit1 interface and its atomic structure provides critical information to elucidate the recruitment mechanisms of SHREC and to inform on mechanisms governing the family of NuRD complexes.

Role of Chp2 and Swi6 HP1 isoforms

The Chp2–Mit1 high-affinity interaction described here explains why Chp2 is almost exclusively found in complex with components of the SHREC complex [Motamedi et al. 2008]. On the other hand, several reports have shown that Swi6 is implicated in SHREC recruitment [Yamada et al. 2005; Sugiyama et al. 2007; Sadaie et al. 2008; Fischer et al. 2009]. Our finding that Swi6 can also bind to the Mit1 N terminus provides an explanation that reconciles these results. The high abundance of Swi6 makes it plausible that it significantly contributes to Mit1 and SHREC recruitment despite its lower affinity for Mit1. Recent results also suggest that Swi6 interacts with Clr3, the HDAC of the SHREC complex, which might further contribute toward SHREC recruitment [Isaac et al. 2017]. However, we show here that Swi6 alone cannot support Mit1 function in CDL mutants, and Swi6 overexpression only partially complements loss of Chp2 [Sadaie et al. 2008]. These observations suggest that the fission yeast heterochromatin system relies on the specific aspects of the Chp2–Mit1 interaction. One possibility is that the high-affinity interaction permits Chp2 to efficiently recruit a very low abundance complex like SHREC to the H3K9 methyl mark without raising HP1 protein concentrations to levels that would disrupt the equilibrium between HP1 proteins, H3K9 methyl marks and HP1 client proteins.

Sequence comparison suggests that the mechanism of the Chp2–Mit1 interaction is conserved in the Schizosaccharomyces family, and how HP1 proteins interact with nucleosome remodelers remains an interesting subject. Human HP1 proteins have been found to associate with the Mit1 homologue CHD4 [Vermeulen et al. 2010], and recent evidence has revealed that human HP1 and CHD4 form a complex with the DNA-binding factor ADNP to regulate lineage-specific gene expression in mouse embryonic stem cells [Ostapcuk et al. 2018]. The findings presented here provide key information to study fundamental principles of how HP1 proteins and the chromatin remodeling machinery interact to regulate gene expression.

Materials and methods

Pull-down experiments

All sequences were cloned into YFP-containing vectors of the Multibac system [Bieniossek et al. 2009] using Gibson cloning [Gibson et al. 2008]. Mit1 constructs were cloned into acceptor vectors as N-terminal fusions of OneStep-MBP tags or OneStep Sumostar tags, followed by a T7 tag [Novagen] and a TEV cleavage site. Chp2 and the Clr1MID construct (included to stabilize the Mit1 C terminus) [Job et al. 2016] were cloned into donor vectors as N-terminal fusions to HisSUMO followed by T7 tags. The HisSUMO tag is cleaved off in the cells during expression by endogenous SUMO proteases, yielding protein carrying only the T7 tag. Donor and acceptor vectors were combined by Cre recombination for coexpression as needed. Proteins were expressed in transfected
SI9 insect cells in adhesion culture in six well plates with $3 \times 10^6$ cells per well in 3 mL media (Amimed) at 27°C. Cells were harvested by aspirating the media when all cells were expressing YFP and frozen (~80°C) until further use. Cells were scraped from the plate surface under the addition of pull-down buffer (400 mM KCl, 100 mM Tris at pH 7.5, 2 mM Mg-acetate, 5 mM β-mercaptoethanol) supplemented with 0.1% NP40 and protease inhibitors, cell debris was removed by centrifugation and the supernatant was incubated with StrepTactin (IBA) beads for 1 h at 4°C with agitation. After incubation, unbound material was removed, the beads were washed five times with pull-down buffer and bound proteins were eluted with elution buffer (pull-down buffer + 5 mM desthiobiotin).

**Western blotting**

Samples were run on Bis-Tris gels and transferred to nitrocellulose membrane [Biorad]. Proteins were detected by Western blotting using antibody against either tag or protein of interest, followed by incubation with secondary antibody labeled with the DyLight system and scanning with the Odyssey Imaging System [LI-COR].

**Yeast two-hybrid screening**

Chp2 constructs were cloned in-frame to the Gal4 binding domain into the pGBK7 vector, transformed into the Y2Hgold strain and selected on SD/-Trp/medium. Mit1 construct-containing pGAD7T vectors were transformed into the Y187 strain and transformed cells were selected on SD/-Leu medium. Appropriate strains were mated and selected on SD/-Trp/-Leu. To detect protein interactions, growth assays were performed at 30°C on SD/-Trp/-Leu/-His and SD/-Trp/-Leu/-His/-Ade medium.

**Protein expression and purification**

All sequences were cloned into vectors of the MultiColi system [Bieniossek et al. 2012] using Gibson cloning [Gibson et al. 2008]. All constructs for individual expression were cloned as N-terminal fusions to a Strep-SUMO-tag into the pACE1 vector, while for coexpression the chromo shadow domain of Chp2 was cloned without tag into pACE2. While for coexpression the chromo shadow domain of Chp2 was cloned without tag into pACE2, N-terminal fusions to a Strep-SUMO-tag into the pACE1 vector, and 0.5 mg/mL-Arabinose before incubation for 18 h at 18°C. Cells were grown to log phase at 37°C, cooled on ice, and induced with 0.4 mM IPTG. Pellets were resuspended in phosphate buffered saline (PBS) supplemented with protease inhibitors and frozen at ~80°C until further usage. For purification, cell pellets were thawed in purification buffer (400 mM KCl, 50 mM Tris at pH 7.5, 5 mM β-mercaptoethanol) supplemented with protease inhibitors and cells were ruptured using either sonication or the Emulsiflex homogenizer. Cell debris was removed by centrifugation and the supernatant was bound to a 5-mL Streptactin column (Qiagen). Cell debris was removed by centrifugation and the supernatant was bound to a 5-mL Streptactin column (Qiagen). After washing with purification buffer, tagged protein was eluted with elution buffer (pull-down buffer + 5 mM desthiobiotin).

**Yeast two-hybrid screening**

Chp2 constructs were cloned in-frame to the Gal4 binding domain into the pGBK7 vector, transformed into the Y2Hgold strain and selected on SD/-Trp/medium. Mit1 construct-containing pGAD7T vectors were transformed into the Y187 strain and transformed cells were selected on SD/-Leu medium. Appropriate strains were mated and selected on SD/-Trp/-Leu. To detect protein interactions, growth assays were performed at 30°C on SD/-Trp/-Leu/-His and SD/-Trp/-Leu/-His/-Ade medium.

**Protein expression and purification**

All sequences were cloned into vectors of the MultiColi system [Bieniossek et al. 2012] using Gibson cloning [Gibson et al. 2008]. All constructs for individual expression were cloned as N-terminal fusions to a Strep-SUMO-tag into the pACE1 vector, while for coexpression the chromo shadow domain of Chp2 was cloned without tag into pACE2. While for coexpression the chromo shadow domain of Chp2 was cloned without tag into pACE2, N-terminal fusions to a Strep-SUMO-tag into the pACE1 vector, and 0.5 mg/mL-Arabinose before incubation for 18 h at 18°C. Cells were grown to log phase at 37°C, cooled on ice, and induced with 0.4 mM IPTG. Pellets were resuspended in phosphate buffered saline (PBS) supplemented with protease inhibitors and frozen at ~80°C until further usage. For purification, cell pellets were thawed in purification buffer (400 mM KCl, 50 mM Tris at pH 7.5, 5 mM β-mercaptoethanol) supplemented with protease inhibitors and cells were ruptured using either sonication or the Emulsiflex homogenizer. Cell debris was removed by centrifugation and the supernatant was bound to a 5-mL Streptactin column (Qiagen). After washing with purification buffer, tagged protein was eluted with elution buffer (pull-down buffer + 5 mM desthiobiotin).

**Crystallization and structure processing**

Mit1(1-81)-Chp2(316-380) were grown at 18°C as 1-µL hanging drops in a 1:1 ratio of protein to reservoir solution [2.4 M sodium malonate, pH 7, 3% 1,6-hexanediol]. Crystals were flash frozen in liquid nitrogen and datasets were collected at beamline PXIII at the SLS, PSI Villigen. Data were processed with XDS [Kabsch 2010], scaled with Aimless [Evans and Murshudov 2013] and the structure was solved by molecular replacement using Swi6 as a model [PDB ID 1E0B] with Phaser [McCoy et al. 2007]. The model was built in Coot (Emsley and Cowtan 2004) and refined with Phenix [Emsley and Cowtan 2004, Adams et al. 2010], Pymol [Schrödinger] was used for the preparation of figures.

**ITC**

ITC experiments were performed at 25°C using a VP-ITC [Microcal] and a Nano ITC calorimeter [TA instruments]. All proteins used were dialyzed overnight against ITC buffer (400 mM NaCl, 50 mM Tris-HCl pH 7.5, 5 mM β-mercaptoethanol) prior to experiments. Ten microliters of Chp2 or Swi6 at 100 to 500 µM were injected in 200 sec time intervals into the cell holding Mit1 constructs at concentrations between 5 and 50 µM. After subtracting heat enthalpies for titrations of the respective proteins into buffer, the ITC data were analyzed with OriginLab [Figs. 3, 4] or Nano-Analyzer Data Analysis software [TA instruments] [Fig. 3, Mat1CILACDL]. Protein concentrations were determined by absorbance at 280 nm.

**Generation of S. pombe strains**

*S. pombe* strains were grown and manipulated as previously described [Job et al. 2016] and strains used in this study can be found in Supplemental Table S1. For crosses, tetrad dissection analysis was used, with plating on selective media and followed by PCR to check the obtained genetic backgrounds.

A strain in which endogenous Mit1 was replaced by the rpl42-natMX cassette *mit1::rpl42-natMX* was obtained by transforming wild-type yeast with a DNA fragment (generated by restriction digest of the respective plasmid) containing the rpl42-natMX cassette flanked on either side by 500 bp of respective sequence found in the 5′- and 3′-UTR of Mit1, making use of
homology recombination for insertion into the genome (Bähler et al. 1998; Fennessy et al. 2014). The strain was selected by rounds of positive and negative selection using the cycloheximide sensitivity conferred by rpl42Δ and the nourseothricin [clon-NAT] resistance conferred by natMX.

Strains carrying full-length Mit1 C-terminally tagged with 13 copies of the c-myc epitope [13myc] at the endogenous locus or truncations or mutations thereof were generated by replacing the rpl42-natMX cassette in the mit1::rpl42-natMX strain mentioned before with 13myc-tagged constructs of choice, by transformation with respective DNA fragments and homology recombination.

For silencing assays, strains were generated by crossing with an otr1::ura4+ reporter strain. Silencing of ura4+ renders cells auxotrophic for uracil and resistant to 5-fluoroorotic acid [5-FOA].

**Silencing assays S. pombe**

*otr1::ura4* reporter strains were grown overnight at 30°C in YES to a density of 5 × 10^6 cells/mL. Cells were pelleted by centrifugation and resuspended in water. Tenfold serial dilutions were spotted onto PMG + Adenine + Leucine + Uracl [nonselective], PMG + Adenine + Leucine –[ura], PMG + Adenine + Leucine + 100 mg/L Uracl + 2 mg/mL 5-FOA (FOA), with 1 × 10^6 cells in the highest density spots.

**Small-scale protein extraction from S. pombe**

Strains were grown in YES to a density of 5 × 10^6 cells/mL and 6 × 10^7 cells per strain were pelleted by centrifugation and resuspended in 300 µL 4x cOmplete, EDTA-free (Roche), 2 mM PMSF on ice. Samples were mixed with 300 µL cold NaOH (0.6 M), incubated on ice for 10 min and spun down. The supernatant was removed and the pellet was resuspended in 70 µL gel loading dye supplemented with 2x cOmplete, 1 mM PMSF and 4% β-mere before boiling for 3 min at 98°C.

**Coimmunoprecipitation from S. pombe**

Strains were grown in YES to a density of 5 × 10^6 cells/mL, pelleted by centrifugation, resuspended in cold PBS supplemented with 1x cOmplete, EDTA-free [Roche], 1 mM PMSF, and 10 mM DTT and frozen in liquid nitrogen. Then, 150 × 10^6 cells were thawed in lysis buffer [50 mM Heps, pH 7.5, 300 mM NaCl, 1 mM EDTA, 5 mM CHAPS, 10 mM DTT, 1 mM PMSF, 1x cOmplete [Roche]], added to glass beads and subjected to two 20 sec runs in a beadbeater for cell disruption. Debris was removed by centrifugation and the lysate was incubated with Myc-Trap_MA resin (ChromoTek) for 1 h with agitation. After washing, bound proteins were eluted by adding 30 µL gel loading dye +10 mM DTT and boiling for 5 min.

**ChIP**

Cells for ChIP were grown in 50 mL of YES medium to a density of 1.2 × 10^8 cells/mL. For RNA Polymerase II and H3K14ac, a dual-crosslinking approach was employed as previously described (Tian et al. 2012). Briefly, cells were incubated at 18°C for 2 h, pelleted, resuspended in 5 mL PBS and crosslinked at room temperature with 1.5 mM ethylene glycol bis-succinimidyl succinate (EGS, Thermo Scientific). After 30 min of incubation, cells were crosslinked in 1% formaldehyde. Cells were resuspended in ChIP Buffer (50 mM HEPES/KOH pH 7.6, 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% Na-Deoxycholate, 1 mM PMSF, 5 mM Na-Butyrate, 1x cOmplete [Roche]), and lysed by beadbeating. Chromatin was enriched by centrifugation and sonicated for 15 min (30 sec/30 sec ON/OFF) in a Bioruptor Pico. 50 µL of sheared soluble chromatin were diluted with 450 µL ChIP Buffer, mixed with 1 µg of antibody [H3K14ac [ab52946], RNA PolII [ab817], and Mpc mAb #2276] and incubated for 2 h, followed by 45 min incubation with Protein A/G magnetic beads. The bead/protein complex was washed three times with ChIP Buffer and once with 50 mM HEPES/KOH pH 7.6, 500 mM NaCl, 1 mM EDTA, once with 5 mM Tris-Cl pH 8, 250 mM LiCl, 0.5% Triton X-100, 0.5% Na-Deoxycholate, 0.05% Tween 20, and once with TE [10 mM Tris-Cl pH 8, 1 mM EDTA]. The protein-DNA complex was eluted in 50 mM HEPES/KOH pH 8, 10 mM EDTA, 1% SDS at 65°C for 15 min, and the crosslinking reversed overnight at 65°C. The samples were then treated with proteinase K and DNA was purified by phenol-chloroform extraction. qPCR was performed using primers given in Supplemental Table S2 and act1+ was used as internal control.

**RT-qPCR**

Strains were grown in YES to a density of 5 × 10^6 cells/mL, pelleted by centrifugation and washed with water. RNA isolation was performed with Trizol extraction followed by phenol/chloroform extraction and ethanol precipitation. DNA was then further removed by DnaseI treatment and phenol/chloroform extraction followed by ethanol precipitation. cDNA was generated with EvoScript Universal CDNA Master kit (Roche) and qPCR performed using SYBR Green I Master kit on LightCycler 480 instrument (Roche). qPCR primers used in this study are listed in Supplemental Table S2. Data were analyzed using the ΔΔCt method (Yuan et al. 2006).

### Accession codes

Coordinates and structure factors have been deposited in the Protein Data Bank under accession code 6FTO.

### Acknowledgments

We thank Yvan Piétre for technical help and Michael Hothorn and Ulrich Hofmann for help with ITC. We are grateful to Roman Ulm and Roman Podolec for help with yeast-two-hybrid. We thank Janet Partridge, Jun-ichi Nakayama, Robin Allshire, and Shiv Grewal for providing strains. We thank Chris Lima for supplying the Supmo expression vectors. We acknowledge the Swiss Light Source at the Paul Scherrer Institut, Villigen (SLS), Switzerland for provision of synchrotron radiation facilities and thank Vincent Olieric for assistance in using beam line PXIII. We thank the Functional Genomics Center Zurich for support with mass spectrometry analysis. We thank Michael Hothorn and Marc Bühler for helpful comments on the manuscript. This work was supported by the Swiss National Science Foundation SNF Professorship grants [PP00P3_139137, PP00P3_163760_1, PP00P5_172904 to T.S.], Fondation Ernest et Lucie Schmidheiny, Fonds Constantin Tepali and Société Académique de Genève (T.S.).

**Author contributions:** The paper was conceptualized by T.S.; methodology by K.L. and T.S.; investigation by K.L., A.S., and T.S.; writing the original draft by K.L. and T.S.; writing, reviewing, and editing by K.L., A.S., and T.S.; resources, T.S.; funding acquisition, T.S.; supervision, T.S.
References

Adams PD, Afonine PV, Bunkóczi G, Chen VB, Davis IW, Echols N, Headd JJ, Hung L-W, Kapral GJ, Grosse-Kunstleve RW, et al. 2010. PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr D Biol Crystallogr 66: 213–221. doi:10.1107/S0907444909052925

Azzaz AM, Vitalini MW, Thomas AS, Price JP, Blacketer MJ, Cryderman DE, Zirbel LN, Woodcock CL, Elcock AH, Wallrath LL, et al. 2014. Human heterochromatin protein 1α promotes nucleosomal associations that drive chromatin condensation. J Biol Chem 289: 6850–6861. doi:10.1074/jbc.M113.512137

Bähler J, Wu JQ, Longtine MS, Shah NG, McKenzie A III, Steever AB, Wach A, Philippen P, Pringle JR. 1998. Heterologous modules for efficient and versatile PCR-based gene targeting in Schizosaccharomyces pombe. Yeast 14: 943–951. <493.:AID-YEA2922.3.CO;2-Y

Bannister AJ, Zegerman P, Partridge JF, Miska EA, Thomas JO, Bayne EH, White SA, Kagansky A, Bijos DA, Sanchez-Pulido L, Bieniossek C, Nie Y, Frey D, Olieric N, Schaffitzel C, Collinson I, Bieniossek C, Imasaki T, Takagi Y, Berger I. 2012. MultiBac: expanding the research toolbox for multiprotein complexes. Trends Biochem Sci 37: 49–57. doi:10.1016/j.tibs.2011.10.005

Cowieson NP, Partridge JF, Allshire RC, McLaughlin PJ. 2000. Dimerisation of a chromo shadow domain and distinctions from the chromodomain as revealed by structural analysis. Curr Biol 10: 517–525. doi:10.1016/S0960-9822(00)00467-X

Creamer KM, Job G, Shanker S, Noller PH, Ponting CP, Rappsilber J, Romier C, Berger P, Richmond TJ, Steinmetz MO, et al. 2009. Automated unrestricted multigene recombineering for multigenic strains nucleosomes within heterochromatin to suppress transcription. Mol Cell Biol 34: 2046–2061. doi:10.1128/MCB.01609-13

Eisenberg JC, Elgin SCR. 2014. HP1α: a structural chromosomal protein regulating transcription. Trends Genet 30: 103–110. doi:10.1016/j.tig.2014.01.002

Emsley P, Cowtan K. 2004. Coot: model-building tools for molecular graphics. Acta Crystallogr D Biol Crystallogr 60: 2126–2132. doi:10.1107/S090744490419158

Evans PR, Murshudov GN. 2013. How good are my data and what is the resolution? Acta Crystallogr D Biol Crystallogr 69: 1204–1214. doi:10.1107/S0907444913090061

Fennessy D, Grallert A, Krupp A, Cokoja A, Bridge AJ, Petersen J, Patel A, Tallada LL, Boke E, Hodgson B, et al. 2014. Extending the Schizosaccharomyces pombe molecular genetic toolbox. PLoS One 9: e97683. doi:10.1371/journal.pone.0097683

Fischer T, Cui B, Dhakhshurnoorthy J, Zhou M, Rubin C, Zoccola M, Vecnstra TD, Grewal SIS. 2009. Diversity of HP1 proteins in heterochromatin assembly and functions in fission yeast. Proc Natl Acad Sci 106: 8998–9003. doi:10.1073/pnas.0813063106

Gibson DG, Benders GA, Axelrod KC, Zaveri J, Algire MA, Moadie M, Montague MG, Venter JC, Smith HO, Hutchison CA. 2008. One-step assembly in yeast of 25 overlapping DNA frag-
Nishihara K, Kanemori M, Yanagi H, Yura T. 2000. Overexpression of trigger factor prevents aggregation of recombinant proteins in Escherichia coli. *Appl Environ Microbiol* **66**:884–889. doi:10.1128/AEM.66.3.884-889.2000

Nishibuchi G, Machida S, Osakabe A, Murakoshi H, Hiragami-Hamada K, Nakagawa R, Fischle W, Nishimura Y, Kurumizaka H, Tagami H, et al. 2014. N-terminal phosphorylation of HP1a increases its nucleosome-binding specificity. *Nucleic Acids Res* **42**:12498–12511. doi:10.1093/nar/gku995

Ostapcuk V, Mohn F, Carl SH, Basters A, Hess D, Iesmantavicius V, Platero JS, Hartnett T, Eissenberg JC. 1995. Functional analysis of HP1 and CHD4 to control lineage-specifying genes. *EMBO J* **23**:489–499. doi:10.1083/jem.7600088

Shimada Y, Mohn F, Bühler M. 2016. The RNA-induced transcription of trigger factor prevents aggregation of recombinant proteins. *Nat Struct Mol Biol* **23**:2571–2580. doi:10.1038/sj.emboj.760098

Tian B, Yang J, Brasier AR. 2012. Two-step cross-linking for analysis of protein-chromatin interactions. *Methods Mol Biol* **809**:105–120. doi:10.1007/978-1-61779-376-9_7

Vakoc CR, Mandat SA, Olenchock BA, Blobel GA. 2005. Histone H3 lysine 9 methylation and HP1gamma are associated with transcription elongation through mammalian chromatin. *Mol Cell* **19**:381–391. doi:10.1016/j.molcel.2005.06.011

Verdel A, Jia S, Gerber S, Sugiyama T, Gygi S, Grewal SIS, Moazed D. 2004. RNAi-mediated targeting of heterochromatin by the RITS complex. *Science* **303**:672–676. doi:10.1126/science.1093686

Vermaak D, Henikoff S, Malik HS. 2005. Positive selection drives the evolution of rhino, a member of the heterochromatin protein 1 family in Drosophila. *PLoS Genet* **1**:96–108. doi:10.1371/journal.pgen.0010009

Vermeulen M, Eberl HC, Matarese F, Marks H, Denissov S, Butter F, Lee KK, Olsen JV, Hyman AA, Stunnenberg HG, et al. 2010. Quantitative interaction proteomics and genome-wide profiling of epigenetic histone marks and their readers. *Cell* **142**:967–980. doi:10.1016/j.cell.2010.08.020

Volpe TA, Kidner C, Hall IM, Teng G, Grewal SIS, Martienssen RA. 2002. Regulation of heterochromatic silencing and histone H3 lysine-9 methylation by RNAi. *Science* **297**:1833–1837. doi:10.1126/science.1074973

Yamada T, Fischle W, Sugiyama T, Allis CD, Grewal SIS. 2005. The nucleation and maintenance of heterochromatin by a histone deacetylase in fission yeast. *Mol Cell* **20**:173–185. doi:10.1016/j.molcel.2005.10.002

Ye Q, Callebaut I, Pechman A, Courvalin JC, Worman HJ. 1997. Domain-specific interactions of human HP1-type chromodomain proteins and inner nuclear membrane protein LBR. *J Biol Chem* **272**:14983–14989. doi:10.1074/jbc.272.23.14983

Yuan JS, Reed A, Chen F, Stewart CN Jr. 2006. Statistical analysis of real-time PCR data. *BMC Bioinformatics* **7**:85. doi:10.1186/1471-2105-7-85

Zhang K, Mosch K, Fischle W, Grewal SIS. 2008. Roles of the Clr4 methyltransferase complex in nucleation, spreading and maintenance of heterochromatin. *Nat Struct Mol Biol* **15**:381–388. doi:10.1038/nsmb.1406
Transcriptional gene silencing requires dedicated interaction between HP1 protein Chp2 and chromatin remodeler Mit1

Karoline Leopold, Alessandro Stirpe and Thomas Schalch

*Genes Dev.* published online February 26, 2019
Access the most recent version at doi:10.1101/gad.320440.118

### Supplemental Material

http://genesdev.cshlp.org/content/suppl/2019/02/26/gad.320440.118.DC1

Published online February 26, 2019 in advance of the full issue.

### Creative Commons License

This article, published in *Genes & Development*, is available under a Creative Commons License (Attribution 4.0 International), as described at http://creativecommons.org/licenses/by/4.0/.

### Email Alerting Service

Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article or click here.