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Mediator recruits the cohesin loader Scc2 to RNA Pol II-transcribed genes and promotes sister chromatid cohesion

Graphical abstract



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In brief

Mediator, Nipbl, and cohesin are hypothesized to connect enhancers and promoters of mammalian genes. Mattingly et al. reveal a role for Mediator in the recruitment of the Scc2 cohesin loader to RNA Pol II-transcribed genes and in sister chromatid cohesion in yeast, demonstrating evolutionary and functional conservation of these interactions.

Highlights

- Mediator and Scc2 display physical and genetic interactions
- Mediator recruits Scc2-Scc4 to RNA Pol II-transcribed genes
- Mediator-dependent recruitment of Scc2-Scc4 promotes sister chromatid cohesion
- Several Scc2-chromatin complex interactions are evolutionarily conserved



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Mediator recruits the cohesin loader Scc2 to RNA Pol II-transcribed genes and promotes sister chromatid cohesion

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https://doi.org/10.1016/j.cub.2022.05.019

SUMMARY

The ring-like cohesin complex plays an essential role in chromosome segregation, organization, and doublestrand break repair through its ability to bring two DNA double helices together. Scc2 (NIPBL in humans) together with Scc4 functions as the loader of cohesin onto chromosomes. Chromatin adapters such as the RSC complex facilitate the localization of the Scc2-Scc4 cohesin loader. Here, we identify a broad range of Scc2-chromatin protein interactions that are evolutionarily conserved and reveal a role for one complex, Mediator, in the recruitment of the cohesin loader. We identified budding yeast Med14, a subunit of the Mediator complex, as a high copy suppressor of poor growth in Scc2 mutant strains. Physical and genetic interactions between Scc2 and Mediator are functionally substantiated in direct recruitment and cohesion assays. Depletion of Med14 results in defective sister chromatid cohesion and the decreased binding of Scc2 at RNA Pol II-transcribed genes. Previous work has suggested that Mediator, Nipbl, and cohesin connect enhancers and promoters of active mammalian genes. Our studies suggest an evolutionarily conserved fundamental role for Mediator in the direct recruitment of Scc2 to RNA Pol II-transcribed genes.

INTRODUCTION

Cohesin is a vital component in the establishment of sister chromatid cohesion and functions in DNA double-strand break repair and genome organization through chromatin looping. This ringlike complex functions in these roles by the entrapment of one or more DNA helices. The four subunit cohesin complex consists of two structural maintenance of chromosomes (SMC) proteins (Smc1 and Smc3), the kleisin protein (Scc1), and the Huntingtin-elongation factor 3-protein phosphatase 2A-TOR1 (HEAT) repeat subunit Scc3. Smc1 and Smc3 form long coiled coils that dimerize via interactions between their hinge domains.¹⁻³ The N- and C-terminal ends of Smc1 and Smc3 form globular ATP binding heads bridged by Scc1 bound by Scc3.^{4–8} The capture of DNA by the cohesin ring requires ATP hydrolysis by the globular head domains of the SMC subunits.^{9,10} The loading of cohesin rings is dependent on a loading complex that comprises heterodimer Scc2-Scc4.¹¹ The human ortholog of SCC2, NIBPL, is mutated in the human developmental syndrome known as Cornelia de Lange syndrome (CdLS), with many of the mutations occurring in the evolutionarily conserved HEAT repeats.^{12,13} CdLS is characterized by limb malformation, facial dysmorphism, cognitive deficiencies, and structural defects in multiple organs. Sister chromatid cohesion during mitosis does not appear to be affected in CdLS patients. Instead, the misregulation of gene expression during development may underlie the clinical features. Much of what we understand about the function of the cohesin loader is based on studies performed in model organisms such as budding yeast.

Biochemical and structural studies have revealed insights into the cohesin loading mechanism. The Scc2-Scc4 loading complex is made up of two functional modules.^{14–17} The C-terminal end of Scc2 (Scc2C) forms a hook structure containing multiple contiguous HEAT repeats. Scc2C contacts the cohesin ring at numerous sites and is sufficient to load cohesin onto naked DNA *in vitro* but is not sufficient for *in vivo* loading.^{14,18} Scc4 binds to the N-terminal end of Scc2 forming the module crucial for localizing the loading complex onto chromatin.^{14,15} Scc4 bound to the N terminus of Scc2 does not bind DNA *in vitro*, suggesting complex localization is bridged via chromatin adaptors.

The recruitment of the loading complex and loading of cohesin is not sequence specific and instead depends on chromatin structure and chromatin remodelers. A common feature of Scc2 sites is nucleosome depletion.¹⁹ In budding yeast, the loading complex interacts with the chromatin remodeler remodels the structure of chromatin (RSC).^{19,20} RSC generates a nucleosome depleted region required for cohesin loading.²⁰ Additional Scc2 recruitment factors include the histone chaperone and remodeler

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Figure 1. Identification of essential domains in Scc2

(A) Schematic of Ashbya gossipii Scc2 domains aligned to S. cerevisiae Scc2. Point mutants are indicated in red and cyan bars represent regions of the alanine block scan.

(B) Plasmid shuffle assay of CdLS mutations at residues conserved from Nipbl to Scc2. Endogenous SCC2 is deleted and covered by wild-type SCC2 on a URA3 selectable plasmid, whereas a second mutant copy of scc2 is present on a HIS3 selectable plasmid. 10-fold serial dilutions of the indicated strains were spotted on SD-HIS-URA and SD-HIS+5FOA. Growth on 5FOA indicates the viability of mutation. The R787G and G1242V mutations result in reduced growth.

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FACT and the Chd1 chromatin remodeler.^{21,22} Cohesin loading at budding yeast pericentromeres is enhanced through Scc2-Scc4 recruitment by the inner kinetochore protein Ctf19.²³ Replication factors, such as components of the pre-replication complex in *Xenopus* egg extract and the Mcm2-7 replicative helicase in HeLa cells, may play a role in the recruitment of the cohesin loader.^{17,24} Localization of the cohesin loader on chromosomes is not fully understood but multiple protein complexes facilitate its recruitment.

The Mediator complex has been implicated in cohesin loading for promoter-enhancer interactions in vertebrates.²⁵ The Mediator complex is a transcriptional co-regulator required for activation and repression of RNA Pol II transcription.^{26,27} Mediator is a large, evolutionarily conserved, protein complex with four submodules: head, middle, tail, and a reversibly binding Cdk8 kinase module that regulates interaction with RNA Pol II.^{28,29} In yeast, Mediator consists of 21 core subunits and a four subunit Cdk8 kinase module with the Med14 subunit acting as an assembly scaffold.³⁰ Mediator co-localizes with NIPBL and cohesin at enhancers and promoters of actively expressed genes in mouse embryonic stem cells²⁵ and shows a synergistic effect on gene expression and limb development when partially reduced along with NIPBL in mice and zebrafish.³¹ In yeast, Mediator shows a similar binding pattern to RSC and the cohesin loading complex, binding at nucleosome depleted regions and promoters of RNA Pol II genes in addition to tRNA and snRNA genes,³² buts it is unclear if Mediator is part of the loading mechanism in genomes lacking promoter-enhancer interactions.

To better understand the loading of cohesin, we investigated the structural and functional domains of Scc2. We demonstrate that mutations in the evolutionarily conserved C-terminal HEAT repeat domains are lethal in budding yeast. A broad range of Scc2 interactors was identified by mass spectrometry. Comparison with previous mass spectrometry for human NIPBL reveals interactions with chromatin complexes, splicing factors and RNA-binding proteins are evolutionarily conserved. Based on mass spectrometry and a high copy genetic suppressor screen, we identified both a genetic and physical interaction with Mediator. An engineered interaction between Med14 and Scc2C is sufficient for cell growth and cohesin loading. The depletion of Med14 results in loss of cohesion and decreased binding of Scc2 at RNA Pol II-transcribed genes. Mediator, NIPBL, and cohesin have been suggested to regulate gene expression by linking enhancers to promoters in higher eukaryotes. Our studies suggest an evolutionarily conserved role for Mediator in direct recruitment of Scc2 to RNA Pol II-transcribed genes and sister chromatid cohesion.

RESULTS

Identification of essential domains in Scc2

The most common and most severe mutations in CdLS are in the *NIPBL* gene.¹³ To further investigate the structure and function of the yeast cohesin loader, Scc2, we sought to identify essential

domains. Using an amino acid alignment of Scc2 and NIPBL, as well as cataloged CdLS NIPBL mutations,¹³ we identified CdLS missense mutations that occur at amino acids conserved between yeast and humans (Figure S1A). We introduced these mutations into an SCC2 plasmid shuffle strain. We identified two mutations in the HEAT domain of SCC2 that result in significantly reduced growth, $scc2^{R787G}$ and $scc2^{G1242V}$ (Figures 1A and 1B). Additional essential residues were identified using an alanine block scanning approach, changing five consecutive amino acids to alanine (Figure 1C). Regions targeted were amino acids 701-800 and 1201-1300, the regions containing scc2^{R787G} and scc2^{G1242V}, respectively, as well as amino acids 501-600 containing the previously studied temperature-sensitive scc2^{E534K} (scc2-4) allele.^{11,33} A structural model of S. cerevisiae Scc21-1493 was created based on the A. gossypii Scc2¹⁻¹⁴⁷⁹ crystal structure³⁴ (Figure 1A). The E534K mutation is in the N-terminal globular domain (GD1) whereas R787G and G1242V mutations are located on opposite ends of the hook-shaped HEAT domain. Mutations were mapped to the crystal structure of Chaetomium thermophilum Scc2^{385–1840 16} (Figure 1D). The E534K, R787G, and G1242V mutations all occur at surface residues conserved from fungus to humans. Interestingly, the R787G and G1242V mutations occur at or proximal to amino acids that when mutated decreased the ability to interact with and load cohesin.^{16,34} Additionally, R787G and the 781–795 alanine scan mutations fall within a positive patch of amino acid residues that interact with DNA in the gripping state intermediate of topological cohesin loading onto DNA while G1242V is proximal to the interface between Scc2 and the SMC heads^{35–37} (Figure S1B). Our data suggest the gripping domain, along with other regions of the HEAT repeats, are critical for Scc2 function.

Scc2 mutants have misregulated gene expression profiles

To investigate the biological processes affected in Scc2 mutants, gene expression analysis was performed in the scc2^{R787G} and scc2^{G1242V} mutants. A plasmid containing either SCC2^{WT}, scc2^{R787G}, scc2^{G1242V}, or an empty vector (null) was introduced into an Scc2 auxin-inducible degron (Scc2-AID) background. Scc2-AID degradation was confirmed by growth assay on an auxin, 3-indoleacetic acid (IAA), plate and western blot (Figures S2A and S2B). The Scc2 mutant growth phenotypes in the Scc2-AID background were consistent with the plasmid shuffle (Figure S2A) and the degron tagged Scc2 was no longer detectable by western blot after culturing the cells with auxin (Figure S2B). Strains were grown in triplicate and RNA sequencing was performed after 3 h of auxin treatment. No genes were differentially expressed in the scc2^{R787G} strain, and only modest effects in the scc2G1242V and null strains were observed, suggesting direct effects on gene expression are minimal (Figure 2A). Cell growth was not affected until 6 h of auxin treatment, even though the degron tagged Scc2 was undetectable by western blot after 3 h (Figure S2C). RNA sequencing

See also Figure S1.

⁽C) Identification of additional essential residues in Scc2 using the plasmid shuffle assay for alanine block mutants in Scc2.

⁽D) Mutated amino acids mapped to the C-terminal crystal structure surface map of *Chaetomium thermophilum* Scc2^{385–1840}. Point mutants are indicated in red, lethal alanine block mutants in green, and non-lethal alanine block mutants in yellow.

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with a 9-h auxin treatment revealed hundreds of differentially expressed genes in the mutant and null strains (Figure 2A). Indirect effects from Scc2 loss may be more prevalent at this time point relative to the 3-h time point. As expected, the number of differentially expressed genes correlates with the severity of the growth phenotype of each strain, with the most differentially expressed genes in the null strain.

The gene expression profile of each strain can be visualized in the heatmap showing counts per million reads (CPM) at the genes differentially expressed in the null strain (Figure 2B). The scc2^{R787G} mutant expression profile more closely resembles the null strain whereas the scc2G1242V mutant clusters with the wild type (WT). Gene ontology (GO) term analyses for upregulated genes in both the Scc2 mutants and the null show enrichment for genes involved in translation and ribosome assembly and biogenesis. Downregulated genes are enriched for genes required for response to oxidative stress and the oxidationreduction process (Figure 2C). Many of the changes observed in gene expression are likely indirect consequences of the absence of functional Scc2. Cytometry shows the R787G and null mutations result in arrest at G2/M of the cell cycle, whereas the G1242V mutation shows less severe defects (Figure S2D). Nonetheless, these results are highly similar to the previously published gene expression profile in a strain bearing the scc2-4 mutation, grown at permissive temperature with minimal cell cycle defects, which also showed upregulation of ribosome

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Figure 2. Scc2 mutants have misregulated gene expression profiles

(A) Few genes were differentially expressed after 3 h of auxin treatment while hundreds of genes were differentially expressed with auxin treatment extended to 9 h. RNA sequencing was performed in Scc2^{degron} strains containing a plasmid copy of WT, null, *R787G*, or *G1242V SCC2*.

(B) The gene expression profile of $scc2^{R787G}$ is similar to null, whereas $scc2^{G1242V}$ clusters more closely to WT. Heatmap showing the log₂ transformed read counts per million (CPM) of the differentially expressed genes in the null strain after 9 h auxin treatment.

(C) Differentially expressed genes in *scc2*^{*R*787G}, *scc2*^{*G*1242V}, and null compared with WT are enriched for GO terms translation and ribosome biogenesis for upregulated genes and oxidative stress for downregulated genes.

See also Figure S2.

biogenesis and rRNA processing genes and downregulation of genes involved in oxidative phosphorylation.³³ Upregulation of rRNA processing genes was also present in the GO term analyses of the null and $scc2^{R787G}$ strains. Ribosomal protein genes (RPs) were upregulated in $scc2-4^{33}$ and indeed similar profiles are observed in each of the $scc2^{R787G}$, $scc2^{G1242V}$, and null strains compared with WT. Mutations at R787 and G1242 impair interaction with cohesin and cohesin loading is dimin-

ished in the *scc2-4* mutant.^{11,16} Therefore, the gene expression profile in the three different Scc2 mutant strains may reflect a consistent transcriptional response to defects in the loading of cohesin. The *scc2-4* mutant strain is associated with defects in the production of ribosomal RNA, ribosome assembly, and splicing,³³ and similar defects may be present in the R787G and G1242V mutants given the similarity in gene expression profiles. These findings reveal the E534K (*scc2-4*), R787G, and G1242V mutations all result in similar gene expression profiles and likely lead to defects in similar biological processes.

MED14 is a suppressor of scc2 mutants

To identify the genetic interactors of SCC2, we performed overexpression screens with strains bearing the $scc2^{R787G}$ and $scc2^{G1242V}$ mutations. The yeast genomic tiling vector collection was transformed into both the $scc2^{R787G}$ and $scc2^{G1242V}$ SCC2 plasmid shuffle strains. The genomic tiling vector collection consists of 1,588 vectors covering 97% of the S. *cerevisiae* genome with each 2μ *LEU2* selectable plasmid containing \sim 4–5 genes. Multiple tiling vectors rescued the $scc2^{R787G}$ and $scc2^{G1242V}$ growth phenotypes when overexpressed. Candidate suppressor genes were cloned into a *LEU2* selectable galactose inducible vector and tested for suppression in the $scc2^{R787G}$ and $scc2^{G1242V}$ SCC2 plasmid shuffle strains. In addition to SCC2, the two strongest suppressors of the $scc2^{R787G}$ and $scc2^{G1242V}$ mutants were the SCC4 component of the cohesin loading complex and the

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Figure 3. Overexpression of MED14 acts as a growth enhancer in strains bearing point mutations in SCC2

(A) Identification of a genetic interaction between Scc2 and Mediator. A genomic tiling vector screen identified *MED14* and *SCC4* as possible suppressors of *scc2*^{*R787G}</sup> and <i>scc2*^{*G1242V*} mutations. *MED14* and *SCC4* were cloned into a *LEU2* selectable galactose overexpression plasmid and transformed into the *scc2*^{*R787G*} and *scc2*^{*G1242V*} plasmid shuffle strains. 10-fold serial dilutions of the indicated strains were spotted on Gal-HIS-LEU-URA and Gal-HIS-LEU+5FOA. (B) Increase in cell growth is observed in the *MED14* overexpression strain compared with the empty vector (EV) control on 5FOA (B) *MED14* overexpression does not affect protein levels of cohesin or Scc2. Cohesin and Scc2 Myc tagged strains containing the *MED14* galactose overexpression plasmid (MED14 OE +) or empty vector (MED14 OE –) were grown in Gal-LEU and levels of cohesin subunits and Scc2 were analyzed by immunoblot. GAPDH and H2A levels are included as loading controls.</sup>

(C) Overexpressing *MED14* does not increase bulk Scc2 or cohesin on chromatin. Chromatin fractionation was performed on yeast cells and levels of Myc tagged Scc2 or Scc1 in the whole-cell extract (WCE), supernatant (SUP), and chromatin pellet (CP) were visualized by immunoblotting. GAPDH and H2A served as loading controls and positive controls for the SUP and CP fractions, respectively. See also Figure S3.

MED14 subunit of the Mediator complex (Figure 3A). Suppression is specific to the *MED14* subunit, as overexpression of other subunits of Mediator (*MED1, MED2, MED3, MED8, MED17, MED18*, and *MED19*) did not rescue growth (Figure S3A). Med14 is crucial for the intermodular interactions of the Mediator complex, contacting the three main Mediator modules and acting as a scaffolding protein for assembly of the head, middle, and tail modules.^{30,38} *MED14* overexpression also showed a limited rescue effect in the *scc2^{E534K}* (*scc2-4*) mutant (Figure S3B).

MED14 overexpression could potentially upregulate transcription of SCC2 and/or cohesin subunits, thereby rescuing the mutants. To test this possibility, we examined protein levels of Myc epitope-tagged Scc2 and cohesin subunits following MED14 overexpression (Figure 3B). Protein levels of Scc2 and the four core subunits of cohesin are not significantly affected. If the mechanism of rescue is through upregulating transcription, overexpression of MED14 might also be expected to rescue scc4 or cohesin mutations. However, the rescue is specific to mutations in scc2, as no rescue was observed in scc4 or cohesin temperature-sensitive mutants (Figures S3C and S3D). A study in human mammary stem cells showed that increased dosage of the MED14 scaffold stabilized the Mediator complex.³⁹ To test the possibility that increased levels of Mediator stabilize Scc2 on chromatin and therefore increase binding of cohesin, chromatin fractionation was performed (Figure 3C). Overexpressing MED14 did not alter the bulk amount of cohesin or Scc2 associated with chromatin. These findings suggest a genetic interaction between the Scc2 cohesin loader and Mediator subunit Med14. The mechanism of rescue does not appear to be increased transcription leading to higher levels of the loading complex or cohesin. Moreover, *MED14* overexpression does not create a global increase in chromatin association of the cohesin loader. These results suggest *MED14* overexpression may create more functional binding or support binding at a subset of locations.

Scc2 and Mediator physically interact

Nipbl and Mediator co-localize at enhancers and promoters and interact in higher eukaryotes.²⁵ After observing a genetic interaction between *SCC2* and *MED14* in yeast, we tested if Scc2 and Med14 physically interact by co-immunoprecipitation (coIP). Scc2-Myc co-precipitated with Med14-Flag in whole-cell extract (WCE). The coIP of Scc2 and Med14 persists after treating WCE with benzonase, suggesting the interaction is DNA independent (Figure 4A). Further supporting a physical interaction between Mediator and the cohesin loader, Scc2 co-precipitates with Med1 and Med16 subunits of Mediator (Figures S4A and S4B). Mutations in Scc2 did not impact the interaction with Med14 (Figure 4B). Additionally, the point mutations did not impact the coIP of Scc2 with Scc4 or two subunits of cohesin (Figures S4C–S4E). To determine if the interaction between the



Figure 4. Scc2 and Mediator physically interact

(A) Scc2 and Med14 interact by co-immunoprecipitation independent of DNA. Myc tagged Scc2 was immunoprecipitated from WCE. Co-precipitation of Flagtagged Med14 was analyzed by immunoblotting. Benzonase treating WCE to remove DNA did not affect the observed co-precipitation.

(B) R787G and G1242V mutations do not affect the Scc2-Med14 interaction. Immunoprecipitations from the lysate of Myc tagged wild type, no tag, R787G, and G1242V Scc2 were analyzed for coIP of Med14-Flag by immunoblotting. The ratio of Scc2 to Med14 in the Scc2 mutants is comparable with WT in the Myc pull-downs.

(C) Diagram indicating inter- (green) and intra- (purple) protein cross-links of the Scc2-Scc4 complex. Positions of Scc2 domains are marked.

(D) Mediator is detected by mass spectrometry of purified Scc2. Cross-linking mass spectrometry was performed on Myc tagged Scc2 purified from WCE with anti-Myc magnetic beads with on bead disuccinimidyl sulfoxide (DSSO) cross-linking and on bead digestion. The heatmap of 1e6 log_2 transformed dNSAF values shows enrichment of Mediator subunits in seven MS runs with purified Scc2-Myc compared with six no tag control runs. The loader and cohesin subunits are included as a positive control. Asterisks indicate statistical significance at p < 0.05.

See also Table S2 and Figures S4 and S5.

cohesin loader and Mediator is dependent on the presence of cohesin or the chromatin deposition of Scc2 mediated by Scc4, we performed coIP of Scc2 and Med14 after depletion of auxin-inducible degron tagged Scc1 or Scc4. Depletion of Scc1-AID and Scc4-AID was confirmed by a growth assay (Figure S4F). Neither depletion impacted the interaction between Scc2 and Med14 (Figures S4G and S4H), suggesting the interaction is independent of cohesin and Scc4.

To further define the interaction between Scc2 and Mediator we performed cross-linking mass spectrometry analysis of *in vivo*

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purified Scc2-Myc endogenously expressed from its native promoter. Although cross-linking data has been reported from recombinant cohesin loading complex,³⁴ this is the first cross-linking proteomics performed with *in vivo* purified Scc2. Although many proteins were enriched in our proteomic analysis of Scc2, Scc4 was the only protein identified with inter-protein cross-links to Scc2 (Figure 4C). The inter- and intra-protein cross-links observed between Scc2 and Scc4 are consistent with previous reports,^{15,34} suggesting the method was successful. In addition to previously observed intra-molecular cross-links, cross-links within Scc2 were identified that may contribute to the flexible conformations exhibited by the Scc2 hook and the Scc2N-Scc4 head of the cohesin loading complex observed by electron microscopy.¹⁴

Subunits of the Mediator complex, as well as known interactors of Scc2, were detected by MS in the Scc2-Myc purifications. Scc4 was the most abundant protein after Scc2, and all subunits of cohesin were present (Figure 4D). We detected peptides from 19 Mediator subunits in at least one of the Scc2 mass spectrometry runs. The results of Mediator co-purification with Scc2 are summarized in the heatmap (Figure 4D). Five Mediator subunits were significantly enriched, one from the head module (Med17), two from the tail (Med5 and Med15), and two from the kinase module (Med12 and Med13). We conclude that the interaction of the cohesin loader and Mediator complex, previously identified in higher eukaryotes,²⁵ is conserved in yeast.

Evolutionarily conserved interactions between Scc2 and a broad range of chromatin-associated proteins

In addition to Mediator, mass spectrometry identified many chromatin remodelers, proteins implicated in cohesin loading, and chromatin-modifying complexes not previously reported to interact with Scc2. Multiple subunits from the RSC complex, a chromatin remodeler that acts as an adapter for the cohesin loader,²⁰ were enriched in the Scc2 purifications (Figure S5A). We also observed significant enrichment of other complexes implicated in cohesin recruitment such as the FACT and Chd1 chromatin remodelers^{21,22} and the MCM2-7 and ORC components of the pre-replication complex (pre-RC)^{17,24} (Figure S5A). The Irc5 and Chl1 helicases, which have been implicated in cohesin deposition,⁴⁰⁻⁴⁵ were also significantly enriched (Figure S5A). The histone acetyltransferases SAGA and NuA4 and the methyltransferase COMPASS, chromatin modifier complexes involved in the regulation of transcription, were examples of newly discovered and highly enriched interactors, as was the Ccr4-Not complex involved in mRNA regulation and transcription initiation (Figure S5B). In addition to cohesin, subunits from condensin and Smc5/6 complexes were enriched in the Scc2 purifications (Figure S5C), consistent with a role for Scc2 in the loading of these complexes.⁴⁶

Proteomics of human cohesin and Nipbl revealed interactions with RNA-binding proteins, splicing factors, MCM proteins, and Mediator.⁴⁷ Many splicing and RNA-binding proteins were likewise found as Scc2 interactors in yeast (Table S2). In addition to identifying protein complexes that could potentially be involved in cohesin localization, our data demonstrate that many interactions are evolutionarily conserved. Mediator is hypothesized to form a complex with Nipbl and cohesin to loop enhancers to promoters in metazoan genomes. The interaction of

Mediator and Scc2 in yeast, an organism that lacks long-range enhancer-promoter interactions as well as canonical topologically associating domains (TADs), raises the possibility that this interaction may have been co-opted for genome organization in metazoans but may have a more elemental function.²⁵

Mediator as a chromatin receptor for Scc2C

The functional modularity of the cohesin loader has been displayed through structural and biochemical studies.^{14,15,17} Scc4 binds the N-terminal domain of Scc2 and is required for chromatin targeting in vivo,¹⁴ whereas Scc2C is capable of loading cohesin onto naked DNA in vitro.18,48 Tethering of Scc2C to the RSC complex, as well as other chromatin remodelers, bypasses the requirement for Scc4 and is sufficient for cohesin loading.²⁰ We did not detect significant enrichment of remodelers that fail to rescue cell growth in the tethering assay, such as the Ino80 and Swr1 complexes. RSC, Isw1, and Chd1, remodelers that restore viability, were all enriched in our proteomics experiment (Figure S5). To test Mediator as a chromatin receptor for Scc2C, Med14 was tagged with GFP at the C terminus and GFP binding protein (GBP) was fused to the N terminus of Scc2C in an Scc2 auxin-inducible degron background. Strikingly, Med14 is a viable receptor for GBP-Scc2C and sustained robust cell growth upon Scc2 depletion at levels similar to the Sth1 subunit of the RSC complex (Figure 5A). Cohesin levels were measured by ChIP/qPCR at several locations. Binding at chromosome arms and centromere-adjacent regions was diminished in cells expressing only GBP-Scc2C, whereas the tethering of Med14-GFP to GBP-Scc2C restored cohesin at both arms and centromere-adjacent regions to approximately half of the wild type (Figure 5B). This suggests that the Mediator complex can act as an Scc2C receptor and facilitate cohesin loading, similar to RSC.

If Mediator plays a significant role in the recruitment of Scc2 and cohesin, then sister chromatid cohesion may be compromised without Mediator. We evaluated sister chromatid cohesion by monitoring lacO-LacI-GFP-marked pericentromere, arm, or telomere sites in Med14 auxin-inducible degron tagged strains. Degron function and efficient depletion of Med14 was confirmed by growth assay and western blot (Figures 5C and 5D). The strains containing the degron tagged Med14 exhibited markedly reduced growth with the addition of auxin (Figure 5C). Med14 was depleted by adding auxin to cells arrested in G1 using alpha-factor. The cells were then released into a G2/M arrest by nocodazole treatment, depleting Med14 within one cell cycle (Figures S6A and S6B). Degron tagged Med14 was not detectable after auxin treatment (Figure 5D). Med14 depletion resulted in defective sister chromatid cohesion at all loci examined (Figure 5E). Levels of cohesin subunits in WCE were not affected by Med14 depletion (Figure S6C), suggesting the cohesion defects cannot be attributed to decreased levels of cohesin. These results indicate a crucial role of Mediator in facilitating sister chromatid cohesion in yeast.

Mediator facilitates the recruitment of Scc2 to RNA Pol II gene groups

To further define the role of Mediator in the localization of the cohesin loader, we performed calibrated ChIP-seq of Scc2 in a Med14 conditional depletion background. Scc2-Myc was





Figure 5. Mediator as a chromatin receptor for Scc2C

(A) Tethering of GBP-Scc2C to Med14-GFP bypasses the need for Scc2. Cells expressing the indicated components were streaked on to YPD medium containing 3-indoleacetic acid (IAA) to degrade the Scc2^{degron}. The Sth1 subunit of RSC is included as a positive control.

(B) Tethering GBP-Scc2C to Med14-GFP reconstitutes cohesin loading. Cells were synchronized in G1 with alpha-factor, Scc2^{degron} was depleted with IAA, and cells were released into nocodazole-imposed mitotic arrest. Cohesin levels were assessed by ChIP against HA-tagged Scc1 followed by quantitative real-time PCR at three chromosome arms and two centromere-adjacent cohesin binding sites and a negative control site. Means and SEM of three independent experiments are shown.

(C) Validation of Med14^{degron} in the presence of auxin (IAA). 10-fold serial dilutions of yeast strains used to examine sister chromatid cohesion, containing a lacO-LacI-GFP marked pericentromere, arm, or telomere site and either WT or degron tagged Med14, were spotted onto YPD+IAA to deplete Med14^{degron}. Depletion of Med14 results in diminished cell growth.

(D) Med14 protein is degraded in sister chromatid cohesion strains after the addition of auxin. Cells were synchronized in G1 with alpha-factor, Med14^{degron} was depleted with IAA, and cells were released into nocodazole-imposed G2/M arrest. Cells were pelleted and WCE prepared and analyzed for the depletion of Med14 by immunoblot.

(E) Mediator promotes sister chromatid cohesion. Cells were synchronized in G1 with alpha-factor, Med14^{degron} was depleted with IAA, and cells were released into nocodazole-imposed mitotic arrest. Cells were fixed and DAPI stained, and the percent of large-budded cells with a single DAPI mass containing 2 GFP spots, indicating loss of sister chromatid cohesion, was scored. A representative image of a cell with cohesion (1 spot) and loss of cohesion (2 spots) is shown. Means and SEM of three independent experiments are shown. A minimum of 100 cells were scored for each replicate; WT CEN, ARM, TEL n = 523, 443, 570 and Med14 degron CEN, ARM, TEL n = 603, 576, 637; **** p <.0001; two-way ANOVA test. See also Figure S6.

introduced in a Med14-AID-FLAG background. The functionality of the Med14 degron was assessed by the inhibition of cell growth with auxin (Figure 6A). Cultures were arrested in G1, auxin added (if depleting), and released into G2/M arrest (Figures S6D and S6E). Depletion of Med14 was verified by western blot and levels of Scc2 were not affected (Figure 6B). Enrichment of DNA in the ChIP compared with a control with primary antibody omitted was confirmed by polymerase chain reaction (PCR) (Figure S6F).

Peaks within 1 kb of a transcription start site (TSS) were mapped to genes by biotype and the ratio of peaks without and with Med14 (IAA/noIAA) was examined (Figure 6C). Although the

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average Scc2 binding at all peaks was not affected, we found that depletion of Med14 results in a modest average decrease in Scc2 peaks at protein coding genes, with the largest decrease at the snoRNA, and ribosomal protein genes. Metagene analysis confirms decreased Scc2 binding at the RNA Pol II-transcribed snoRNA and ribosomal protein genes (Figure 6D). This suggests Mediator facilitates Scc2 recruitment to RNA Pol II gene groups in budding yeast.

DISCUSSION

The co-localization of the cohesin loader and Mediator at gene promoters and enhancers has been observed in multiple studies with their interaction proposed to induce enhancer-promoter interactions.^{25,51,52} Here, we show the interaction between the cohesin loader and Mediator is conserved in yeast and provide evidence that Mediator acts as a chromatin adapter for the cohesin loader, facilitating the recruitment of Scc2 to RNA Pol II gene groups. Yeast lack enhancers and long-range interactions for gene activation are rare. Our findings suggest a more evolutionarily conserved function for Mediator in cohesin loader recruitment, independent from enhancer-promoter interactions.

Our analysis identified essential domains in the C-terminal HEAT repeat domain of S. cerevisiae Scc2 and a genetic interaction between SCC2 and MED14. Previous studies have highlighted the essential residues of the HEAT domain of the cohesin loader. Mutations in the Scc2 ortholog in Chaetomium thermophilum, corresponding to budding yeast Scc2 R787 and G1242 residues, result in diminished binding with an N-terminal

Figure 6. Mediator facilitates Scc2 recruit-

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(A) Validation of Med14^{degron} in Scc2-Myc tagged strain used for calibrated ChIP-seq. 10-fold serial dilutions were spotted on YPD and YPD+IAA.

(B) Med14^{degron} is depleted in the presence of IAA. Cells grown in triplicate were synchronized in G1 with alpha-factor, Med14^{degron} was depleted with IAA, and cells were released into nocodazoleimposed mitotic arrest. Cells were then cross-linked with formaldehvde. A portion of the culture was removed before cross-linking for gel electrophoresis and levels of Med14 and Scc2 were monitored by immunoblot. H2A served as a loading control.

(C) Scc2 peaks are most affected at snoRNA and RP genes after Med14 depletion. A spike-in of Mis4-Myc tagged S. pombe was added to each culture for calibration, and the Scc2-Myc immunopurified chromatin was extracted for ChIP-seq. The experiment was performed in triplicate with Med14 depleted (IAA) and Med14 present (noIAA). Peaks within 1 kb of transcription start sites were mapped to genes by biotype. The boxplot shows the log₂ ratio of Scc2 peaks with Med14 depleted over no depletion (IAA/noIAA).

(D) Metagene analysis of the ChIP-seq data shows that Scc2 peaks decrease at snoRNA and ribosomal protein genes when Med14 is depleted.

See also Figure S6.

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fragment of the Scc1 subunit of cohesin. A budding yeast double mutant with one mutation adjacent to the R787G mutation (K788A R792A) is defective in loading

cohesin.^{16,34} Recent studies in S. pombe show that the C-terminal HEAT domain of Scc2 contains two DNA binding domains, one of which corresponds to a region including S. cerevisiae R787, that are crucial for initial binding of cohesin with DNA in an intermediate of topological loading.35,36 We speculate that the growth phenotypes observed with the R787G and G1242V mutations are due to diminished cohesin loading. The overexpression of Scc4, known to be essential for localization of the loading complex to chromatin,¹⁴ rescued the growth of these mutants. Increased levels of Scc4 may increase the localization of the loading complex, allowing for an increase in cohesin loading and account for the increase in cell viability. The observation that the Med14 subunit of Mediator rescues these mutants similar to Scc4, suggests a role for Mediator in cohesin loader recruitment. Increased dosage of MED14 in human cells led to higher protein levels of multiple Mediator subunits due to stabilization of the complex rather than changes in transcription.³⁹ We speculate that the overexpression of the Med14 subunit in yeast similarly drives an increase in the Mediator complex because it is the scaffolding subunit.

Cohesion defects are observed with deficiencies in other purported chromatin adapters of the cohesin loading complex.^{20,23,53,54} Likewise, we report a cohesion defect upon Mediator depletion. We find that depletion of Mediator decreased cohesion at a pericentromeric, arm, and telomeric site. Therefore, Mediator is important to maintain normal levels of cohesion. A recent study in S. cerevisiae also implicates Mediator in sister chromatid cohesion.⁵⁵ A strain forced to use the meiosis-specific kleisin subunit of cohesin during mitotic growth selectively

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accumulated compensatory mutations in the Mediator complex that improved sister chromatid cohesion. Interestingly, mutations that introduced early stop codons in subunits of the kinase module showed the highest increase in cell fitness. This suggests the Mediator complex devoid of the kinase module, which is the version that regulates RNA Pol II transcription, may also promote cohesion. A genetic screen with an S. pombe mutant defective in acetylation of Smc3 identified genetic interactions of cohesin with Mediator and the CCR4-NOT complex, which is also present in our Scc2 purifications.⁵⁶ Other recent studies in budding yeast found that segmental amplifications containing SCC2 and MED14 were acquired as adaptations to DNA replication stress⁵⁷ and that an oxidative stress-induced association of Mediator and cohesin plays a role in base excision repair of DNA lesions.⁵⁸ Evidence is accumulating for a functionally significant and evolutionarily conserved interaction between Mediator and Scc2.

Physical interaction between the cohesin loader, cohesin ring, and Mediator has previously only been observed in higher eukaryotes.^{25,47,58} Mass spectrometry of Mediator purified from neural stem cells contained NIPBL and both SMC subunits of cohesin as well as other complexes that were present in our Scc2 purification such as subunits of the COMPASS and SWI/ SNF complexes.⁵⁹ Mediator is postulated to interact with transcription factor activation domains through dynamic shapeagnostic "fuzzy" interfaces. Multiple subunits of cohesin are predicted to contain such domains, providing potential Mediator interaction interfaces.⁶⁰ We show that Scc2-Mediator interaction is present in yeast and that Scc2C fusion to Mediator is sufficient for cohesin loading, similar to the RSC complex.²⁰ We propose Mediator acts as a chromatin receptor for Scc2 at a subset of RNA Pol II gene groups, namely, snoRNA and ribosomal protein genes. Interestingly, both Scc2 and Mediator localize at snoRNA genes and promote proper snoRNA transcription in yeast, 33,61 and human Mediator regulates the transcription of snRNA genes.⁶² We speculate that the evolutionarily conserved functional interaction between Mediator and the cohesin loader occurs at a subset of RNA Pol II- highly transcribed genes.

Nucleosome-depleted DNA is required for the loading of cohesin. Chromatin remodelers that promote the recruitment of Scc2-Scc4 may generate a nucleosome-free region required to load cohesin.²⁰ The human ISWI remodeler has been implicated in cohesin loading,⁶³ fitting with the emerging idea that the Scc2-Scc4 receptor landscape is multifactorial. Mediator may participate in creating or marking the nucleosome-depleted DNA environment where cohesin loading occurs. Mediator interacts with multiple chromatin remodelers such as SWI/SNF, RSC, and Chd1^{32,64–66} and Mediator itself may contribute to nucleosome displacement in both yeast and humans.^{67,68} Whether Mediator works together with chromatin remodelers to load cohesin or is sufficient to recruit Scc2-Scc4 and generate nucleosome depleted DNA in the absence of remodelers is an area for future investigation.

Mutations in NIPBL and the human SWI/SNF complex result in congenital developmental disorders with closely related clinical features, known as CdLS and Coffin-Siris syndrome, respectively.^{12,69} Mutations in Mediator are also linked to developmental syndromes and cognitive defects in humans. Interestingly, a screen for chromosomal rearrangements in clinically diagnosed CdLS patients, lacking mutations in cohesin or the cohesin loader, identified patients with chromosomal imbalances containing Mediator, helicase, and nucleosome remodeler genes.⁷⁰ Mutations in the Mediator complex result in FG, Lujan, and X-linked Ohdo syndromes as well as schizophrenia.^{71–74} Investigation into how Mediator, the cohesin loader, and its chromatin receptors interact to load cohesin and regulate genome architecture and gene expression will deepen our understanding of the etiologies of these human diseases.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
 - Lead contact
 - Materials availability
 - Data and code availability
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
- METHOD DETAILS
 - Yeast strain construction
 - SCC2 mutant plasmid shuffle
 - SCC2 point mutants suppressor screen
 - RNA sequencing
 - Whole cell extract co-immunoprecipitation
 - Scc2 purification and mass spectrometry
 - Calibrated ChIP-seq
 - Quantitative ChIP
 - Sister chromatid cohesion assay
- QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. cub.2022.05.019.

ACKNOWLEDGMENTS

We thank the Molecular Biology, Bioinformatics, Microscopy, and Media Prep facilities at the Stowers Institute, Kansas City, MO, USA for their assistance. We thank Dan Bradford for his assistance in performing the genomic tiling vector screen. We thank the members of the Gerton Lab at the Stowers Institute for their advice and discussions. For technical support, we thank Cindy Maddera and Rhonda Egidy. The graphical abstract for this manuscript was created with BioRender.com. This study was supported by the Stowers Institute for Medical Research. F.U. received funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation program (grant agreement no. 670412) and the Francis Crick Institute, which receives its core funding from Cancer Research UK (FC001198), the UK Medical Research Council (FC001198), and the Wellcome Trust (FC001198). S.M. was supported by an EMBO long-term fellowship.

AUTHOR CONTRIBUTIONS

M.M. and J.L.G. conceived the study. M.M. carried out the characterization of Scc2 mutants, suppressor screen, Scc2 mutant RNA-seq, analysis of Scc2 proteomics, cohesion assay, and Scc2 ChIP-seq experiments. C.S. performed the computational analysis for the RNA-seq and ChIP-seq experiments. S.M. performed the tethering experiment and cohesin ChIP experiment. Y.H., Y.Z.,

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Z.W., and L.F. performed the Scc2 proteomic experiments. S.M. and F.U. provided discussion and suggestions for writing the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: January 4, 2021 Revised: April 7, 2022 Accepted: May 9, 2022 Published: June 1, 2022

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Anti-Mouse IgG HRP conjugated	GE Healthcare	RRID# AB_772210
Anti-Rabbit IgG HRP conjugated	GE Healthcare	Cat# NA934V
Anti-c-Myc 9E10	BioLegend	RRID# AB_2148451
Anti-c-Myc-Peroxidase	Roche	RRID# AB_390910
Monoclonal ANTI-FLAG M2	Sigma-Aldrich	RRID# AB_259529
Monoclonal ANTI-FLAG M2-Peroxidase	Sigma-Aldrich	RRID# AB_439702
Anti-HA-Peroxidase	Roche	RRID# AB_390917
Histone H2A antibody	Active Motif	RRID# AB_2687477
GAPDH antibody	GeneTex	RRID#AB_11174761
Anti-HA F7	Santa Cruz	Cat# sc-7392
Anti-HA (12CA5)	Roche	RRID# AB_514505
Chemicals, peptides, and recombinant proteins		
α-factor	Zymo Research	Cat# Y1001
Nocodazole	Sigma-Aldrich	Cat# M1404
Indole-3-acetic acid (IAA)	Sigma-Aldrich	Cat# 45533
Formaldehyde Solution	Sigma-Aldrich	Cat# 252549
Proteinase K	ThermoFisher	Cat# EO0491
Benzonase nuclease	Sigma-Aldrich	Cat# E1014
RNase A	ThermoFisher	Cat# EN0531
Propidium iodide solution	Sigma-Aldrich	Cat# P4864
GelRed Nucleic Acid Gel Stain	Biotium	Cat# 41103
5-Fluoroorotic Acid	Zymo Research	Cat# F9001
DAPI	ThermoFisher	Cat# 62248
cOmplete Protease Inhibitors Cocktail	Sigma-Aldrich	Cat# 11836145001
Triton X-100	Sigma-Aldrich	Cat# T9284
NP-40 Alternative	Sigma-Aldrich	Cat# 492016
Sodium Deoxycholate	Sigma-Aldrich	Cat# S1827
2-Chloroacetamide	Sigma-Aldrich	Cat# C0267
Dithiothreitol	ThermoFisher	Cat# R0861
Sequencing grade modified trypsin	Promega	Cat# V5111
Copper (II) sulfate	Sigma-Aldrich	Cat# C1297
Critical commercial assays		
Pierce ECL 2 western blotting substrate	ThermoFisher	Cat# 80196
Nitrocellulose Blotting Membrane	Sigma-Aldrich	Cat# GE10600002
Dynabeads Protein G	ThermoFisher	Cat# 10004D
Pierce Anti-c-Myc Magnetic Beads	ThermoFisher	Cat# 88842
NuPAGE LDS Sample Buffer	ThermoFisher	Cat# NP0007
NuPAGE bis-tris 4-12% gradient protein gel	ThermoFisher	Cat# NP0323BOX
Gateway LR Clonase II Enzyme mix	ThermoFisher	Cat# 11791020
Gateway BP Clonase II Enzyme mix	ThermoFisher	Cat# 11789020
TruSeq Stranded Yeast Ribo-Zero Gold Kit	Illumina	Cat# MRZY1324
TruSeq Stranded RNA LT kit with Ribo-Zero Gold	Illumina	Cat# RS-122-2301
KAPA HTP Library Prep Kit for Illumina	Roche	Cat# KK8234
PowerUp SYBR Green Master Mix	ThermoFisher	Cat# A25742

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited data		
ChIP sequencing data	This study	GEO: GSE201194
RNA sequencing data	This study	GEO: GSE201194
Mass spectrometry data	This study	MassIVE: MSV000086339, ProteomeXchange: PXD022100
Unprocessed images presented in this manuscript	This study	https://www.stowers.org/research/ publications/libpb-1710
Experimental models: Organisms/strains		
All Saccharomyces cerevisiae strains used in this study are listed in Table S1	Lab stock and this study	N/A
Oligonucleotides		
POA1-F AAACGGCCACATCAAATACC	Muñoz et al. ²⁰	N/A
POA1-R TCCAAGGGACTCCGAATATG	Muñoz et al. ²⁰	N/A
MRP10-F ACCCCCTCTTCCCAGACTAA	Muñoz et al. ²⁰	N/A
MBP10-B CCAGCACATTTAGGGCTCAT	Muñoz et al. ²⁰	N/A
MET10-F ACTTGTGTGGCCCTACTTGG	Muñoz et al. ²⁰	N/A
MET10-B CGACTITGATGCCTCTTTCC	Muñoz et al ²⁰	N/A
	Muñoz et al ²⁰	N/A
	Muñoz et al 20	N/A
	Muñoz et al 20	N/A
	Muñoz et al 20	N/A
	Muñoz et al 20	N/A
GLT1-E GGGTGTGGAGTTTGTGGTCT	Muñoz et al 20	N/A
		N/A
	This study	N/A
Yeast genomic tiling collection	OpenBiosystems	Cat# YSC4613
pRS413-SCC2-7MYC Point and alanine block mutants	This study	N/A
pHyg-AID*-6FLAG	Addgene	Cat# 99519
pHyg-AID*-6HA	Addgene	Cat# 99520
Software and algorithms		
Fiji (Image J) V2.3.1/1.53f	https://imageJ.net/	RRID# SCR_002285
FlowJo v9	FlowJo	RRID# SCR_008520
Prism V9.3.1	GraphPad	RRID# SCR_002798
PyMOL	http://www.pymol.org/	RRID# SCR_000305
ProLuCID	Xu et al. ⁷⁵	N/A
DTASelect v1.9	Tabb et al. ⁷⁶	N/A
NSAF v0.0.1	Zhang et al. ⁷⁷	N/A
STARTApp	Nelson et al. ⁷⁸	N/A
ReactomePA	Yu and He ⁷⁹	RRID# SCR_019316
HiSeq Control Software v2.2.58	Illumina	N/A
Illumina Primary Analysis version RTA v2.4.11	Illumina	N/A
Illumina Secondary Analysis bclfastq2 v2.17	Illumina	N/A
R 3.6-4.0	https://www.r-project.org/	RRID# SCR_001905
Bioconductor 3.10-3.13	http://www.bioconductor.org	RRID# SCR_006442
edgeR v3.28.0	http://www.bioconductor.org/ packages/edgeR/	RRID# SCR_012802

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
TopHat 2.1.1	http://ccb.jhu.edu/software/ tophat/index.shtml	RRID# SCR_013035
bowtie2 bowtie2-2.4.2	http://bowtie-bio.sourceforge.net/ bowtie2/index.shtml	RRID# SCR_016368
STAR STAR_2.6.1c	http://code.google.com/p/rna-star/	RRID# SCR_004463
Macs2 2.1.2	https://github.com/macs3-project/ MACS	RRID# SCR_013291
Ensembl 96	http://www.ensembl.org/	RRID# SCR_002344

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by lead contact Jennifer L. Gerton (jeg@stowers.org).

Materials availability

Yeast strains and plasmids generated in this study are available for distribution.

Data and code availability

- RNA-seq data and ChIP-seq data have been deposited at GEO and are publicly available as of the date of publication. All mass
 spectrometry files have been deposited to the ProteomeXchange via the MassIVE repository and are publicly available as of the
 date of publication. Accession numbers are listed in the key resources table. Original images and data underlying this manuscript can be accessed from the Stowers Original Data Repository (ODR). The ODR link is listed in the key resources table.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

All Saccharomyces cerevisiae yeast strains used in this study were of the W303 or BY4741 background and are listed in Table S1. Cells were cultured at 30°C. α -factor was used at a concentration of 7.5 μ g/ml, nocodazole at 15 μ g/ml and indole-3-acetic acid (IAA) at 1 mM.

METHOD DETAILS

Yeast strain construction

To construct the *SCC2* mutant plasmid shuffle strains, *SCC2-7MYC* including the promoter region was amplified from yeast genomic DNA and cloned into pRS413. Plasmid pRS413-*SCC2-7MYC* point mutations and alanine block mutations were made by site-directed mutagenesis and transformed into the *SCC2* plasmid shuffle strain. Endogenous genes were epitope tagged by gene targeting using polymerase chain reaction (PCR) products. The auxin degron strains were constructed using pHYG-AID*6FLAG (Addgene #99519) or pHYG-AID*6HA (Addgene #99520) and yeast strain BY26972 from the Japan National BioResource Project (NBRP::Yeast).

SCC2 mutant plasmid shuffle

SCC2 plasmid shuffle yeast strains were grown overnight in synthetic media lacking histidine (SD-HIS). The following day cultures were diluted to OD600 1.0 and 10 fold serial dilutions were spotted on SD-HIS-URA and SD-HIS-URA+5FOA agarose plates and grown at 30°C.

SCC2 point mutants suppressor screen

SCC2 point mutant plasmid shuffle strains R787G and G1242V were transformed with the yeast genomic tiling vector collection (Open Biosystems) as per standard high-throughput 96-well yeast protocols. Transformants were selected on SD-HIS-LEU-URA. Transformants were cultured overnight in SD-HIS-LEU at 30°C in 96 well plates and pinned to SD-HIS-LEU-URA and SD-HIS-LEU+5FOA 96 well format agarose plates. *Scc2-R787G* strains were pinned at 1:5 and 1:10 dilutions while *scc2-G1242V* strains were pinned at 1:20 and 1:40 dilutions. Candidate suppressor genes were amplified by PCR and cloned into the *LEU2* selectable gal overexpression vector pAG425 using the Gateway (ThermoFisher) cloning system.

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RNA sequencing

Scc2 degron strains carrying pRS413(HIS) plasmid with SCC2-WT, R787G, G1242V or empty vector (null) were cultured in triplicate (except for null strain in duplicate) overnight in SD-HIS. The following day, strains were diluted to \sim OD₆₀₀ = 0.1 in SD-HIS, after 90 minutes, 3-indoleacetic acid (IAA) was added to a concentration of 1mM and cells were cultured for an additional 9 hours to OD₆₀₀ range of 0.4 – 0.8. Cells were pelleted and RNA was extracted. RNA integrity was checked on a 1.2% agarose gel and run on an Agilent Technologies 2100 Bioanalyzer. Samples were ribo-depleted with TruSeq Stranded Yeast Ribo-Zero Gold Kit (Illumina, MRZY1324) per manufacturer's instructions. Following depletion, Illumina libraries were generated using the TruSeq Stranded RNA LT kit with Ribo-Zero Gold, 48; Set A (Illumina, RS-122-2301). Libraries were normalized and pooled together with an average base pair size of 290 and sequenced as 50 bp single reads on the HiSeq 2500 (Illumina) using HiSeq Control Software v2.2.58. Following sequencing, Illumina Primary Analysis version RTA 1.18.61 and Secondary Analysis bcl2fastq2 v2.17 were run to demultiplex reads from all libraries and generate FASTQ files. Reads were aligned to version sacCer3 of the Saccharomyces cerevisiae genome using tophat2. The resulting BAM files were analyzed in R. Bioconductor package edgeR was used to perform differential expression analysis. Data has been deposited at GEO: GSE201194.

Whole cell extract co-immunoprecipitation

Cultures were grown to mid-log phase. IAA was added to 1mM for 3 hours for depletion in degron strains. Whole cell extracts were prepared by resuspending cell pellets in lysis buffer (50 mM Tris (pH 7.5), 150 mM NaCl, 0.1% NP-40, 1 mM dithiothreitol (DTT), 10% glycerol and protease inhibitors) and vortexing with glass beads at 4°C. Cell extracts were diluted with dilution/wash buffer (50 mM Tris (pH 7.5), 150 mM NaCl, 0.1% NP-40). Diluted cell extracts were incubated with the Myc 9E10 antibody (BioLegend) overnight followed by 2 h with protein G dynabeads (Invitrogen-10004D) at 4°C. The beads were washed three times with dilution/wash buffer and proteins were eluted in NuPAGE LDS sample buffer at 95°C for 10 min. Immunoprecipitates were subjected to polyacrylamide gel electrophoresis (PAGE) on NuPAGE 4-12% Bis-Tris Protein Gels (Invitrogen) and Western blotting (Myc-HRP, FLAG-HRP, HA-HRP, GAPDH, H2A).

Scc2 purification and mass spectrometry

Yeast culture

For each yeast strain (either Scc2-13Myc in BY4741 or wild-type BY4741), 6L cultures were grown in YPD medium and the cell pellet frozen and stored at -80°C until use.

Myc-Affinity Capture and on-Beads Crosslinking

In brief, after binding to the Myc-beads and two rounds of washes, bound proteins were crosslinked on-beads with 1mM DSSO for 40min, then digested on-beads with trypsin for 30 min. Multiple rounds of Myc-AP-XL were performed from three independent batches of Scc2-Myc cells and one BY4741 (no tag) cell culture. Several technical replicate LC/MS analyses were acquired from the eluted proteins. In all, seven mass spectrometry runs were performed from the Scc2 purification and six runs of the no tag control to identify proteins enriched in the purified Scc2.

Cell pellets were resuspended in a total volume of up to 30 ml in BH0.15 extraction buffer (25 mM HEPES, pH 7.5, 2 mM MgCl2, 0.1 mM EDTA, 0.5 mM EGTA-KOH, 15% Glycerol, 0.1% NP-40, 150 mM KCl) with freshly added 100x protease inhibitor and 5mM DTT. Cells were frozen using liquid nitrogen and dry ice and lysed with a Waring Commercial Blender, 250 ml container with 5 small pieces of dry ice blended for 10-15 sec. The frozen cells were transferred to a 600 ml beaker and thawed in a warm water bath. 100 ml of 30 mg/ml heparin and 20 ml of 25U/ul SAN (Salt Active Nuclease; ArcticZyme) were added to the thawed cells, which were rotated at RT for 20 minutes. The extract was clarified by spinning the lysate in a tabletop centrifuge for 15 min at 4000 rpm, 4 °C. Supernatant was saved and the pellets resuspended to 30 ml total volume by adding BH0.15 extraction buffer to repeat the freeze/thaw/incubate steps once more. Extractions from two rounds were combined and clarified by centrifugation for a final time at 19,000 rpm for 30 min using a JA25.50 rotor in a floor centrifuge.

Pierce[™] Anti-c-Myc Magnetic Beads (150 µl) were washed with 1 ml of BH0.15 extraction buffer. The supernatant was discarded, and the washed beads were added to the protein lysate and incubated on a rotating wheel overnight at 4°C. Proteins bound to anti-Myc beads were washed twice with 1.5 ml BH0.15. Beads were transferred to a new 1.5 ml tube, washed with 1 ml of pre-elution rinse buffer (50 mM HEPES, pH 7.5, 75 mM KCl, 1 mM EGTA), and the supernatant was discarded. Proteins bound to anti-Myc beads were crosslinked by adding 150 µl pre-elution rinse buffer complemented with protease inhibitor cocktail (Sigma) and DTT (dithiothreitol; Thermo Fisher Scientific) with 0.6 µl of 250 mM disuccinimidyl sulfoxide (DSSO) to a final concentration of 1mM and left to crosslink at room temperature for 40 min. The crosslinking reaction was quenched by adding 7.5 µl of 1 M NH₄CO₃ (final 50 mM) and rotating at RT for 15 min. The supernatant was transferred to a new tube and saved as flow-through.

On-beads protein digestion

Crosslinked proteins bound to anti-Myc beads were washed twice with 500 µl 10 mM Tris-HCl pH7.5, 150 mM NaCl, then denatured, reduced and digested with 100 ml 50 mM Tris-HCl pH7.5, 2 M urea, 1 mM TCEP (Pierce), 5 µl Trypsin at 0.1 mg/ ml (Sequencing Grade Modified Trypsin; Promega), at 30 °C for 30 min. The supernatant was collected. Another 60 µl of 50 mM Tris-HCl pH7.5, 2 M urea, 5 mM 2-Chloroacetamide (CAM, Sigma) were added to the beads to alkylate free cysteines, and the resulting supernatant combined with the first one. This 2-step process was repeated once more. CAM was added to 2.5 mM and the reaction incubated in the dark at RT for 30 min. CaCl₂ was added to 2 mM along with 5 µl of Trypsin, and the digestion was incubated at 37 °C overnight. The digestion was quenched by adding formic acid to 5%. The combined supernatants from these multiple steps constitute the first "elution" (E1). These digestion steps were repeated once, and the collected supernatants combined as the second elution (E2). After



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digestion, the beads were resuspended in 150 μ l of 2% Sodium Deoxycholate (SDS; Sigma) with 50mM HEPES and incubated at 95°C for 30 min. This fraction was labeled as "SDS-E1". From each of the E1, E2, and SDS-E1 fractions, 14 μ l was set aside to analyze protein contents on silver-stained SDS-PAGE.

LC/MS acquisition

Digested peptides were analyzed on an Orbitrap Fusion Lumos mass spectrometer equipped with a FAIMS Pro interface (Thermo Scientific, San Jose, CA) coupled to a Dionex Ultimate 3000 RSCLnano System. Peptides were loaded on an Acclaim[™] PepMap[™] 100 C18 0.3 mm i.D. x 5 mm length trap cartridge (Thermo Scientific, San Jose, CA) with loading pump at 2 ml/min via autosampler. A 75 µm i.d. analytical microcapillary column was packed in-house with 250 mm of 1.9 µm ReproSil-Pur C18-AQ resin (Dr. Masch GmbH, Germany). AgileSLEEVE (Analytical Sales & Products, Pompton Plains, NJ) was used to maintain column temperature at 40°C. The organic solvent solutions were water/acetonitrile/formic acid at 95:5:0.1 (v/v/v) for buffer A (pH 2.6) and at 20:80:0.1 (v/v/v) for buffer B. The chromatography gradient was a 25 min column equilibration step in 2% B; a 3 min ramp to reach 10% B; 90 min from 10 to 40 % B; 6 min to reach 95% B; a 9 min wash at 95% B; 0.1 min to 2% B; followed by a 12 min column re-equilibration step in 2% B. The nano pump flow rate was set to 180 nL/min. Orbitrap Fusion Lumos was set up with peptide identification method as: full MS1 resolution 120,000; ITMS2 isolation window 1.4 m/z, ITMS2 max ion injection time 50 ms, ITMS2 CID 35% with normal scan. FAIMS compensation voltages (CVs) were set up as -40V, -60V, and -80V.

MS dataset processing

Collected MS/MS spectra were searched with the ProLuCID algorithm⁷⁵ against a database of 12276 protein sequences combining 6010 non-redundant *Saccharomyces cerevisiae* proteins (NCBI, 2017-05-16 release), 193 common contaminants, and their corresponding 6138 randomized amino acid sequences. All cysteines were considered as fully carboxamidomethylated (+57 Da statically added), while methionine oxidation was searched as a differential modification. DTASelect v1.9⁷⁶ and *swallow*, an in-house developed software, were used to filter ProLuCID search results at given FDRs at the spectrum, peptide, and protein levels. Here, all controlled FDRs were less than 1%. All 13 data sets were contrasted against their merged data set, using Contrast v1.9 and in house developed *sandmartin* v0.0.1. Our in-house developed software, NSAF7 v0.0.1, was used to generate spectral count-based label free quantitation results dNSAF.⁷⁷ Mass spectrometry files have been deposited to the ProteomeXchange via the MassIVE repository with accession numbers: PXD022100 and MSV00086339.

Calibrated ChIP-seq

250 ml cultures per replicate were grown in YPD to OD600 \sim 0.2 and arrested 1.5 hrs with alpha-factor. IAA was added to 1mM and cultures were held in alpha-factor arrest for an additional 1.5 hrs before release into YPD containing IAA and nocodazole for 2 hrs until mitotic arrest was reached. 50ml of culture was removed to verify Med14 depletion by Western blot. Mis4-Myc tagged S. pombe cells were added as a spike-in control at a 1:4 OD ratio. Cells were fixed in 1% formaldehyde for 30 min at room temperature, then quenched with glycine (final concentration 0.125 M) for 5 min. Cells were washed twice with cold PBS, frozen in liquid nitrogen, and stored in -80°C until used. Cells were lysed by bead beating at 4°C in FA-Lysis SDS buffer (50 mM Hepes (pH7.5), 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.2% SDS, protease inhibitors). The lysate was sonicated using a Covaris SS20 sonicator for 25 min and centrifuged to remove cell debris. 10% of the total chromatin was processed for input and chromatin shearing was verified on a bioanalyzer. Chromatin extract was used to set up three IPs and one no antibody control per replicate and incubated with Myc antibody (BioLegend 626802) at 4°C overnight followed by 4hrs at 4°C with Protein G Dynabeads (Invitrogen 100-04D) prewashed in FA-Lysis buffer. Beads were washed at room temperature with the following sequence of buffers; FA-Lysis buffer, two washes in FA-Lysis buffer with 500 mM NaCI, TEL buffer (0.25M LiCl, 10 mM Tris-HCl pH 8.0, 1 mM EDTA, 1% NP-40, 1% sodium deoxycholate), twice with 1XTE. Dynabeads from the three IPs per replicate and three no antibody controls (one from each replicate) were combined during the final wash. Chromatin fragments were eluted using 200 μl of Elution Buffer (TE + 1% SDS) at 65°C 20 min with agitation. NaCl was added to a final concentration of 250mM, and crosslinks were reversed overnight at 65°C. Elutions were treated with RNAse 1hr at 37°C and proteinase K for 2hr at 55°C. DNA was purified by running the elutions through a Qiagen PCR purification column.

Libraries were prepared with a Sciclone liquid-handling robot using the KAPA HTP Library Prep Kit for Illumina (Roche KK8234) and Bio Scientific NEXTflex DNA barcodes (PerkinElmer). NEXTflex DNA barcode adapters were used at a dilution of 1:125 with 15 cycles of PCR. The resulting libraries were then quantified using a Bioanalyzer (Agilent Technologies) and a Qubit fluorometer (Life Technologies). Libraries were pooled and sequenced on an Illumina NextSeq 500 instrument as High Output 75 bp single read runs. Following sequencing, Illumina Primary Analysis version RTA v2.4.11 was run to demultiplex reads for all libraries and generate FASTQ files. Reads were aligned to version sacCer3 of the yeast genome from UC Santa Cruz using bowtie2 v2.4.2. Concordant alignments were analyzed in R using Bioconductor to generate coverage and normalize the data in rpm. Locations of enrichment for each protein were identified using MACS2 version 2.1.2. ChIP calibration was performed as previously described.^{49,50} The boxplot was constructed using the top 50th percentile of peaks by P-value quantified by the mean RPM value in a 100 base window at the peak center and peaks within 1 kilobase of transcription start sites were mapped to genes by biotype. Metagene plots were constructed by converting coverage of the Scc2 ChIP into rpm per base and then extracting out coverage for a given set of genes, defined as a range of base pairs upstream and downstream of the transcription start site (TSS), and plotting the average of the gene set at each base pair. Data has been deposited at GEO: GSE201194.

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Quantitative ChIP

Chromatin immunoprecipitation was performed as previously described.²⁰ Cells were grown in medium lacking methionine, arrested 1.5 hour with alpha-factor, and shifted to YPD medium containing IAA in addition to alpha-factor for 2 hours, before release to a nocodazole-imposed mitotic arrest. Cohesin levels were assessed by ChIP against HA tagged Scc1 followed by quantitative real-time PCR at three chromosome arm and two centromere cohesin binding sites and a negative control site.

Sister chromatid cohesion assay

Yeast strains containing either wild type *MED14* or *MED14-AID*^{*}, copper inducible lacI-GFP and lacO repeat arrays inserted at either PERICEN, TEL, or ARM sites. Cells were grown in YPD + 200 μ M Cu₂SO₄, arrested in G1 with alpha-factor, IAA added, and released into nocodazole-imposed mitotic arrest. Cells were fixed briefly in 70% ethanol, DAPI stained, and imaged using a Nikon widefield microscope. Z stacks with 15 images at 0.1 μ m intervals were acquired and merged by maximum intensity projection. Percentage of cells showing 2 separate GFP foci was quantified using ImageJ.

QUANTIFICATION AND STATISTICAL ANALYSIS

For the Scc2 mass spectrometry the dNSAF log_2 FoldChange and the p-value between the Scc2-Myc purifications and negative controls acquired in parallel were calculated with the linear model of STARTApp package.⁷⁸ After applying log_2 FoldChange >= 1 and p-value <= 0.05 as filtering criteria for proteins significantly enriched in the Scc2-Myc purifications, the gene subset was used for GO enrichment through enrichGO function in the R package ReactomePA.⁷⁹

DNA from the quantitative ChIP was quantified using the PowerUP SYBR Green Master Mix (ThermoFisher) and a Quant Studio 12K Real-Time PCR System (Thermo Fisher).

For the sister chromatid cohesion assay means and SEM of three independent experiments were used. Percentage of cells showing 2 separate GFP foci was quantified using ImageJ. A minimum of 100 cells were scored for each replicate. Exact number of cells counted (n) is indicated in the figure legend. For statistical analysis, a two-way ANOVA test was performed within GraphPad Prism. **** indicates p value <.0001.