



Epitope tag-induced synthetic lethality between cohesin subunits and Ctf7/Eco1 acetyltransferase

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ABSTRACT

Ctf7/Eco1-dependent acetylation of Smc3 is essential for sister chromatid cohesion. Here, we use epitope tag-induced lethality in cells diminished for Ctf7/Eco1 activity to map cohesin architecture in vivo. Tagging either Smc1 or Mcd1/Scs1, but not Scs3/Irr1, appears to abolish access to Smc3 in *ctf7/eco1* mutant cells, suggesting that Smc1 and Smc3 head domains are in direct contact with each other and also with Mcd1/Scs1. Thus, cohesin complexes may be much more compact than commonly portrayed. We further demonstrate that mutation in *ELG1* or *RFC5* anti-establishment genes suppress tag-induced lethality, consistent with the notion that the replication fork regulates Ctf7/Eco1.

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1. Introduction

Cohesins tether together sister chromatids from early S-phase until anaphase onset and are essential for both high fidelity chromosome transmission and transcription regulation. Cohesin complexes consist of Smc1, Smc3, Mcd1/Scs1 (herein Mcd1) and Scs3/Irr1 and accessory factors Pds5 and Rad61/WAPL [1]. While many of the components that make up the cohesin complex are known, the structure of this complex in vivo remains under intense debate. Based on early electron microscopy (EM) studies and subsequent biochemical assays, a popular model of cohesin is that Smc1 and Smc3 dimerize by hinge–hinge interactions and that their separated head domains are bridged by Mcd1. From this, a huge triangular ring is posited to result with an inner diameter of 35 nm [2,3]. More recent Atomic Force Microscopy, Fluorescence Resonance Energy Transfer (FRET) and EM studies suggest that Smc1 and Smc3 head domains may remain in contact with one another and that their coiled-coil domains fold back to form shortened structures in which the ATPase heads are positioned closely to the hinges [4–6]. These subunit interactions provide for numerous configurations including one ring, bracelets, snaps, two-ring hand-cuffs and C-clamps [7–12].

A major obstacle in discerning the physiologically relevant cohesin architecture in vivo is that there is evidence supporting both one ring and two ring models. Both models rely heavily on analyses of epitope-tagged subunits. Intriguingly, these studies reveal that alternate tags produce cohesin complexes of different biochemical properties [3,5,13]. A critical issue thus becomes the extent to which these tagged cohesin proteins give rise to complexes that behave like wild type complexes in vivo. In either case, chromatid-bound cohesins must be converted to a pairing competent state by the establishment factor Ctf7/Eco1 (herein Ctf7). Ctf7 is an acetyltransferase that modifies Smc3 to produce sister chromatid pairing – linking establishment and cohesin architecture to sister chromatid pairing [14].

Here, we pursue a novel assay to demonstrate in vivo that epitope-tagged cohesin subunits previously used to support one-ring embrace models are unfit when challenged genetically in vivo. We uncovered cohesin tag-induced lethality in cells diminished for Ctf7 acetyltransferase activity. In turn, we exploited this in vivo cohesin architecture assay to ascertain subunit domains proximal to the site of Smc3 acetylation and thus potential regulators of Ctf7-dependent acetylation reactions. In addition, we demonstrate the molecular basis of anti-establishment factors (Rfc5 and Elg1) in regulating sister chromatid pairing reactions [15,16, Maradeo, Garg and Skibbens, submitted for publication].

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2. Materials and methods

2.1. Strain construction

All strains in this study are W303 background (Table 2). To generate Smc1-13MYC integration product PCR was performed using primers TTGACCTATATAGATATTATTAGTTATTTGACGGGTTA-TAGCAGAGGTTGGTTTCATAGAGAATTCGAGCTCGTTTAAAC and AGACAGCAACAAGAAAACCTCGTGAAGATCATAACTTTGGACTTGAGCAATTACGCAGAACGGATCCCCGGGTTAATTA with pFA6a-13MYC-KAN Mx6 Longtine vectors.

2.2. Genetic analysis

Diploid strains were sporulated in 0.3% potassium acetate and tetrads dissected on YPD agar media. The genotypes of the resultant spores were analyzed for wild type, single, double, triple and quadruple mutants. Isolated spores were grown at log phase in YPD followed by 10-fold serial dilutions and then maintained

Table 1A
mcd1:myc rfc5-1 ctg7^{eco1-1} X mcd1:myc elg1.

Genotype	Observed	Expected
<i>mcd1:myc</i>	11	15
<i>mcd1:myc elg1</i>	10	15
<i>mcd1:myc rfc5-1</i>	11	15
<i>mcd1:myc ctg7^{eco1-1}</i>	0	15
<i>mcd1:myc elg1 rfc5-1</i>	17	15
<i>mcd1:myc elg1 ctg7^{eco1-1}</i>	13	15
<i>mcd1:myc rfc5-1 ctg7^{eco1-1}</i>	9	15
<i>mcd1:myc elg1 rfc5-1 ctg7^{eco1-1}</i>	13	15
Dead	36	0
Total	120	

on YPD agar plates at a range of temperatures. For Table 1A, first *MCD1-18MYC elg1* mutant cells were crossed to *eco1-1 rfc5-1* mutant cells, sporulated, dissected and genotypes analyzed. No *MCD1-18MYC eco1-1* double mutant strains were recovered but both *MCD1-18MYC eco1-1 rfc5-1* and *MCD1-18MYC eco1-1 elg1* triple mutants were obtained. To bias the recovery of a *MCD1-18MYC eco1-1* double mutant, *MCD1-18MYC elg1* was crossed to *MCD1-18MYC eco1-1 rfc5-1*, sporulated, dissected and genotypes analyzed.

3. Results and discussion

Sister chromatid cohesion, via conversion of cohesins to a pairing competent state, is established during S phase by Ctf7-dependent Smc3 acetylation [17,18]. Numerous studies employ cohesin-tagged subunits to characterize cohesin structure and acetylation. However, studies that challenge the physiological relevance of these constructs *in vivo* are severely limited. In the process of generating new strains to further characterize cohesion and acetylation regulation, we discovered that C terminally MYC-tagged Mcd1 is lethal when combined with *ctf7* alleles. Strains containing *MCD1-MYC* were crossed to cells containing *ctf7^{eco1-1}*, sporulated, dissected and the resulting tetrads analyzed. Of an expected 24 *MCD1-MYC ctf7^{eco1-1}* spores, no viable isolates were recovered (Table 1A, data not shown). In contrast, *MCD1-MYC* expression is not lethal in combination with other cohesion mutant strains (*scc2*, *scc4* or *pds5*), obviating arguments that the lethality reported here is simply based on compounded cohesion defects [19,20]. Nor is *MCD1-HA* expression lethal in *ctf7* mutant strains, augmenting the specificity of the synthetic lethality reported here [21]. These results reveal that *mcd1:myc* is a severe but cryptic mutant allele of *MCD1* and that the MYC-based C-terminal extension specifically diminishes the ability of Ctf7/Eco1 to acetylate Smc3 – the only known essential substrate for Ctf7 acetylation.

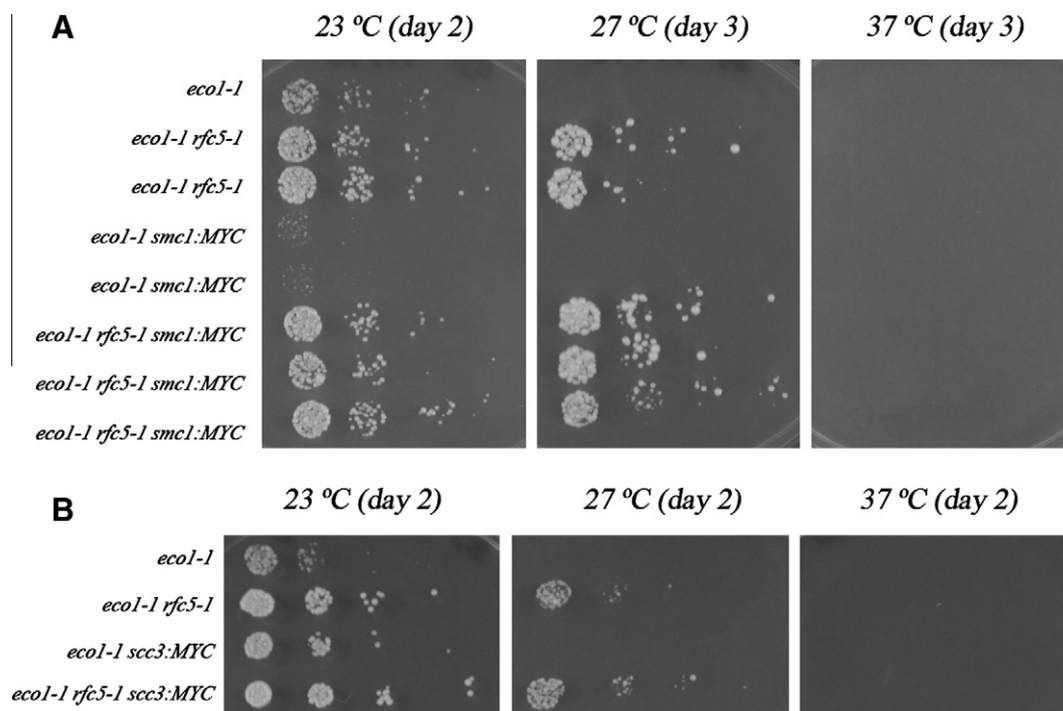


Fig. 1. (A) *SMC1-13MYC* exhibits conditional growth defects in combination with *ctf7^{eco1-1}* mutant cells that can be rescued with mutation in *rfc5*. 10-fold serial dilutions of single, double and triple mutant cells. Colony growth shown for cells on rich medium plates grown at 23, 27, and 37 °C for number of days indicated. (B) *SCC3-18MYC ctf7^{eco1-1}* mutants exhibit no conditional growth phenotypes. 10-fold serial dilutions of single, double and triple mutant cells. Colony growth shown for cells on rich medium plates grown at 23, 27, and 37 °C for number of days indicated.

The head domain of Smc1, composed of both N- and C-termini, interacts closely with the C terminus of Mcd1 [3]. If the bulky MYC adduct on Mcd1 significantly diminishes Ctf7 access to Smc3, then MYC-tagging Smc1 C terminus within *ctf7* mutant cells should phenocopy this lethality. To test this prediction, we crossed strains containing *SMC1-13MYC* allele into *ctf7^{eco1-1}* mutant cells, sporulated, dissected and analyzed the resulting tetrads. Of the expected 9 *SMC1-13MYC ctf7^{eco1-1}* spores, 7 spores were inviable. After prolonged incubation, we were able to recover 2 viable spores, but these are dramatically growth-compromised such that both are near-inviable at temperatures that support robust growth of *ctf7^{eco1-1}* single mutant cells (Fig. 1A and Table 1B). These findings reveal that *smc1-myc* is also a severe but cryptic allele of *SMC1* and suggests that the bulky 13MYC moiety extending from the C-terminus of Smc1 similarly blocks efficient Ctf7-dependent Smc3 acetylation. In combination, these results support a compacted cohesin structure such that Smc1 and Smc3 head domains are in direct contact with each other and also with Mcd1. These results appear to undermine a popular model in which Smc1 and Smc3 are separate but bridged by Mcd1 to form a huge open ring-like structure capable of entrapping two sister chromatids (Fig. 2).

To date, little evidence supports a model in which Scc3/Irr1 (herein Scc3) is in direct contact with Smc1 or Smc3 head domains other than its binding to Mcd1 [3,5]. To augment the specificity of the mapping results described above, we tested whether expression of Scc3-18MYC is lethal in *ctf7* mutant cells. We crossed strains containing *SCC3-18MYC* to *ctf7^{eco1-1}* mutant cells, sporulated, dissected and analyzed the resulting tetrads. In contrast to either *mcd1-myc* or *smc1-myc*, expression of *SCC3-MYC* in *ctf7* mu-

tant cells provided for robust growth at permissive temperature identical to that of single mutant *ctf7* cells (Fig. 1B). Thus, MYC addition to Scc3 is transparent to Ctf7 access to Smc3, positioning Scc3 away from the Smc3 acetylation site (see Table 1C).

Anti-establishment factors are in part defined by the discovery that their deletion rescue *ctf7* mutant cell phenotypes [15,16,18, 22,23]. Likely anti-establishment mechanisms include altering cohesin complex structure to prohibit pairing or sequestering Ctf7 protein to block cohesin acetylation. The tag-induced lethality reported above provides a robust and unique assay to further characterize the recently identified anti-establishment activities of Elg1-RFC and Rfc5 [15,16,Maradeo, Garg and Skibbens, submitted for publication]. If either Elg1 or Rfc5 block Ctf7 access to Smc3, then deletion of *ELG1* or mutation in *RFC5* (*rfc5-1*) should bypass the lethality resulting from co-expression of *ctf7^{eco1-1}* and *mcd1-myc*. To test this hypothesis, *mcd1-myc elg1* double mutant cells were crossed to *ctf7^{eco1-1} rfc5-1* double mutant cells, sporulated, dissected and the resulting tetrads analyzed. While no *ctf7^{eco1-1} mcd1-myc* double mutant isolates were recovered, these crosses produced expected frequencies of both *ctf7^{eco1-1} mcd1-myc elg1* and *ctf7^{eco1-1} mcd1-myc rfc5-1* triple mutant strains (Section 2). A similar level of bypass was obtained when we tested for *rfc5-1* bypass of *ctf7/eco1-1 smc1-myc* phenotypes, although we recovered slightly less than the expected number of spores (Table 1B). In contrast to the severe growth defects exhibited by *ctf7^{eco1-1} smc1-myc* even at 23°, *ctf7^{eco1-1} smc1-myc rfc5-1* triple mutant cells grew robustly at both 23 and 27 °C. Moreover, *ctf7^{eco1-1} smc1-myc rfc5-1* triple mutant cells significantly out-performed *ctf7^{eco1-1}* single mutants at all temperatures tested (Fig. 1A). Thus, both *ELG1*

Table 1B
smc1:myc X ctf7^{eco1-1} rfc5-1.

Genotype	Observed	Expected
Wild type	7	9
<i>smc1:myc</i>	10	9
<i>rfc5-1</i>	8	9
<i>ctf7^{eco1-1}</i>	11	9
<i>ctf7^{eco1-1} rfc5-1</i>	8	9
<i>smc1:myc ctf7^{eco1-1}</i>	2	9
<i>smc1:myc rfc5-1</i>	10	9
<i>smc1:myc ctf7^{eco1-1} rfc5-1</i>	3	9
Dead	13	0
Total	72	

Table 1C
SCC3:MYC X ctf7^{eco1-1} rfc5-1.

Genotype	Observed	Expected
Wild type	8	9
<i>SCC3:MYC</i>	8	9
<i>rfc5-1</i>	10	9
<i>ctf7^{eco1-1}</i>	7	9
<i>ctf7^{eco1-1} rfc5-1</i>	10	9
<i>SCC3:MYC ctf7^{eco1-1}</i>	12	9
<i>SCC3:MYC rfc5-1</i>	10	9
<i>SCC3:MYC ctf7^{eco1-1} rfc5-1</i>	6	9
Dead	1	0
Total	72	

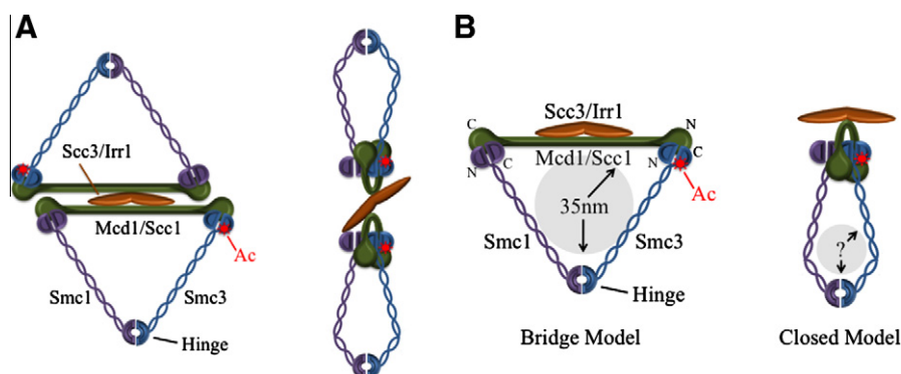


Fig. 2. (A) Left: dimerization cohesins in 'Bridge' conformation. Right: dimerization of cohesins in 'Closed' conformation. Both models reflect evidence of Mcd1–Mcd1 dimerization but that a single copy of Scc3 is present in a cohesin complex [4]. Note that several other pairing-competent cohesin structures (one ring, two ring, C-clamps, bracelet-type spirals) are equally feasible and posited in the literature. (B) Left: separated head domains of Smc1 (purple) and Smc3 (blue) are bridged by Mcd1 (green) to create a huge triangular ring structure of up to 35 nm. Right: Smc1 and Smc3 head domains are in intimate contact with each other and also with Mcd1 to produce a greatly diminished ring lumen. Numerous studies suggest that folding of Smc1 and Smc3 arms positions the hinge region near to the head domains and thus eliminates an embrace-type ring lumen (1). In either model, Scc3/ (orange) binds only through Mcd1/Scc1 and is distal from Smc1/3 head domains. Red sunburst – acetylation site on Smc3; N = N-terminus; C = C-terminus.

Table 2

Strains used in this study are W303 background.

YMM629	<i>MATα eco1-1:ADE rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	[24]
YMM630	<i>MATα eco1-1:ADE rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	[24]
YMM902	<i>Matα SMC1-13MYC:KAN ade2 his3 leu2 trp1 ura3</i>	This study
YMM708	<i>MATα SCC3-18MYC:TRP ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1</i>	[19] ^a
YMM833	<i>MATα MCD1-18MYC:TRP elg1::KAN ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	[25] ^a
YMM864	<i>MATα MCD1-18MYC:TRP eco1-1:ADE rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM926	<i>MATα eco1-1:ADE ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM927	<i>MATα eco1-1:ADE rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM928	<i>MATα eco1-1:ADE SMC1-13MYC:KAN ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM929	<i>MATα eco1-1:ADE SMC1-13MYC:KAN ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM930	<i>MATα eco1-1:ADE SMC1-13MYC:KAN rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM931	<i>MATα eco1-1:ADE SMC1-13MYC:KAN rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM932	<i>MATα eco1-1:ADE SMC1-13MYC:KAN rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM935	<i>MATα eco1-1:ADE ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM936	<i>MATα eco1-1:ADE rfc5-1:ADE ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM937	<i>MATα eco1-1:ADE SCC3-18MYC:TRP ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study
YMM940	<i>MATα eco1-1:ADE SCC3-18MYC:TRP rfc5-1:LEU ade2-1 his3-11,15 leu2-3112 trp1-1 ura3-1 can1-100</i>	This study

^a Backcrossed to W303.

deletion and *rfc5-1* mutation are individually sufficient to rescue cohesin tag-induced defects in Ctf7-dependent Smc3 acetylation and sister chromatid pairing reactions. In combination, these findings suggest a direct mechanism through which Elg1 and Rfc5 oppose establishment at the DNA replication fork by hindering Ctf7/Eco1 accessibility to Smc3.

The inability to biochemically recover cohesin–cohesin interactions was previously used to argue against two-cohesin ring models [3]. In this study, we used a genetic assay to reveal that these epitope-tagged cohesins do not function as wild type proteins in vivo but instead are in fact severe but cryptic alleles. Tag-induced lethality appears specific to Ctf7 function and thus Smc3 acetylation – given that similar tagged subunits support viability in other cohesin mutants. These findings undermine more speculative but plausible models that tag-induced lethality is based on adversely affecting some other cohesin mechanism (for instance, tag-based alteration of SMC ATPase cycles). Moreover, the tag-induced lethality reported here supports a compact cohesin architecture in vivo in which Smc1 and Smc3 head domains are in intimate proximity to one another and also to Mcd1 – in contrast to a huge ring model (predicated on separate but bridged Smc1,3 heads) conjectured to entrap two chromatin fibers (Fig. 2).

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