Evidence that a complex of SIR proteins interacts with the silencer and telomere-binding protein RAP1

Paolo Moretti, Katie Freeman, Lavanya Coodly, and David Shore

Department of Microbiology, Columbia University College of Physicians & Surgeons, New York, New York 10032 USA

The maintenance of transcriptional silencing at HM mating-type loci and telomeres in yeast requires the SIR2, SIR3, and SIR4 proteins, none of which appear to be DNA-binding proteins. Here we show that SIR3 and SIR4 interact with a carboxy-terminal domain of the silencer, telomere, and UAS-binding protein RAP1. We identified SIR3 and SIR4 in a two-hybrid screen for RAP1-interacting factors and showed that SIR3 interacts both with itself and with SIR4. The interaction between RAP1 and SIR3 can be observed in vitro in the absence of other yeast proteins. Consistent with the notion that native SIR proteins interact with the RAP1 carboxyl terminus, we show that mutation of the endogenous SIR3 and SIR4 genes increases transcriptional activation by LexA/RAP1 hybrids. To test the importance of the RAP1–SIR3 interaction for silencing, we identified mutations in the RAP1 carboxyl terminus that either diminish or abolish this interaction. When introduced into the native RAP1 protein, these mutations cause corresponding defects in silencing at both HMR and telomeres. We propose that RAP1 acts in the initiation of transcriptional silencing by recruiting a complex of SIR proteins to the chromosome via protein–protein interactions. These data are consistent with a model in which SIR3 and SIR4 play a structural role in the maintenance of silent chromatin and indicate that their action is initiated at the silencer itself.

[Key Words: Transcriptional silencing, transcriptional activation; mating type; telomere position effect; SIR proteins; protein-protein interactions]

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In the yeast Saccharomyces cerevisiae, the RAP1 gene encodes an essential regulatory protein that functions as both an activator and repressor of transcription (Shore and Nasmyth 1987; Kurtz and Shore 1991; Sussel and Shore 1991; Kyrion et al. 1993). RAP1 DNA-binding sites have been identified within the promoter elements of a large number of genes, including most ribosomal protein genes and a number of glycolytic enzyme genes. In cases where deletion analyses have been performed, these RAP1-binding sites behave as upstream activation sites (UASs) (Rotenberg and Woolford 1986; Vignais et al. 1987; Buchman et al. 1988b; Chambers et al. 1989; Nishizawa et al. 1989; Bitter et al. 1991). Conversely, RAP1-binding sites at the HMR-E and HML-E silencers are required for complete repression of mating-type genes at the HMR and HML loci (Brand et al. 1987; Kimmerly et al. 1988; Mahoney et al. 1991; McNally and Rine 1991). At telomeres, multiple RAP1-binding sites are found within the terminal poly(C_1-3A) repeats (Longtine et al. 1989; Gilson et al. 1993), where the protein is involved in both the regulation of telomere structure and telomeric silencing (Conrad et al. 1990; Lustig et al. 1990; Sussel and Shore 1991; Kyrion et al. 1992, 1993).

Several lines of evidence indicate that RAP1 is a context-dependent regulator. First, the specific sequence of a RAP1-binding site does not determine its regulatory function: Silencer-associated sites can function to activate transcription when placed within promoters and vice versa (Shore and Nasmyth 1987). Second, in both silencer and promoter contexts, RAP1 usually requires other regulatory proteins bound nearby to execute its proper function. For example, it appears that the juxtaposition of a RAP1-binding site with an autonomously replicating sequence (ARS) consensus element (ACS) and an ABF1-binding site can constitute a silencer element (McNally and Rine 1991). A complex of six proteins called the origin recognition complex (ORC) binds to the ACS and cooperates with RAP1 at silencers (Bell et al. 1993; Foss et al. 1993; Micklem et al. 1993). At several glycolytic gene promoters the activator protein GCR1 has been shown to bind near RAP1 and contribute to the activation of these genes (Baker 1991; Huie et al. 1992).

Two types of experiments indicate that the activation and silencing functions of RAP1 are at least in part encoded by genetically separable domains of the protein. Missense mutations (rap1<sup>m</sup>) have been identified in a carboxy-terminal region of RAP1 that lead to defects in silencing but do not affect activation (Sussel and Shore...
Moretti et al.

1991]. In a different approach, GAL4 DNA-binding domain (GAD)/RAP1 hybrid proteins have been used to map both activation and silencing domains within RAP1 [Hardy et al. 1992a]. A carboxy-terminal region of RAP1 (amino acids 630–695), just beyond a centrally located DNA-binding domain, activates transcription in the context of GAD/RAP1 hybrids, whereas overexpression of the last 150 amino acids of the protein (678–827) can interfere with silencing, suggesting that this domain may play a role in silencing in the context of the native protein.

To explain how RAP1 might work as a context-dependent regulator, we proposed that when the carboxy-terminal domain is bound to a silencer it interacts specifically with factor(s) involved in repression. Alternatively, when bound to a promoter, we imagine that the adjacent activation region functions to stimulate transcription, either by interacting directly with the basal transcriptional machinery or through interactions with coactivators [Hardy et al. 1992a]. The inhibition of silencing brought about by the overexpression of the RAP1 carboxyl terminus could be accounted for by the titration (‘squelching’) of a RAP1-interacting factor involved in repression. To begin to test these ideas, we employed a strongly derepressing GAD/RAP1 hybrid in the two-hybrid system [Fields and Song 1989; Chien et al. 1991] and identified a new yeast gene, RIF1 [RAP1 interacting factor 1] [Hardy et al. 1992b]. A deletion of the RIF1 gene results in derepression of an HMR-E silencer, whose ARS element has been deleted, and in the elongation of telomeres. These two phenotypes are characteristic of silencing-defective rapi mutations and lead to the suggestion that one function of RAP1 is to direct the binding of RIF1 to silencers and telomeres [Hardy et al. 1992b]. However, RIF1 is only required for silencing at HMR when the A site (an ARS consensus element) is deleted from the HMR-E silencer, whereas the RAP1-binding site is required for full repression under all circumstances tested, suggesting that RIF1 might not be the only RAP1-interacting protein involved in silencing.

Several trans-acting factors, in addition to RIF1 and the known silencer-binding proteins ORC, RAP1, and AFB1 [Shore et al. 1987; Buchman et al. 1988a] are involved in the repression of HM loci and telomeres. Three SIR (silence information regulator) genes [SIR2, SIR3, and SIR4] are required to maintain the repressed state [Haber and George 1979; Klar et al. 1979; Rine et al. 1979; Rine and Herskowitz 1987; Aparicio et al. 1991], and SIR1 is important in the establishment of repression at HM loci [Pillus and Rine 1989; Chien et al. 1993]. Although all four of these SIR genes have been cloned and sequenced [Shore et al. 1984; Ivy et al. 1986; Marshall et al. 1987; Stone et al. 1991], the precise role of their gene products in silencing remains unknown. None of the SIR proteins appears to bind DNA directly.

To test the possibility that the SIR proteins might work in part by interacting directly with RAP1, we extended our previous screen for RAP1-interacting proteins, in this case using fusions between the bacterial DNA-binding protein LexA and the carboxyl terminus of RAP1. This new two-hybrid screen identified SIR3 and SIR4 as RAP1-interacting factors. In direct tests using the two-hybrid system we also show that SIR3 can interact with itself and with SIR4. We demonstrate that SIR3 can bind to RAP1 in vitro in the absence of other yeast proteins. In support of the idea that the endogenous SIR proteins interact with the RAP1 carboxyl terminus, we show that mutations in three SIR genes (SIR2, SIR3, and SIR4) and RIF1 increase the activation potential of LexA/RAP1 hybrid proteins when these hybrids contain an intact RAP1 carboxyl terminus. We also show that the RAP1–SIR3 interaction does not require the native SIR1–4 genes, and is improved by mutation of RIF1. Finally, we demonstrate that mutations in RAP1 that diminish SIR3 binding in the two-hybrid system cause defects in silencing at both HM loci and telomeres when incorporated into the native RAP1 gene. These results suggest that silencing is initiated by direct protein–protein interactions at the RAP1 carboxyl terminus that recruit SIR3 and SIR4 to DNA.

Results

Identification of SIR3 and SIR4 as RAP1-interacting factors using the two-hybrid system

Previous studies using the two-hybrid system indicated that the RIF1 protein interacts with the carboxyl terminus of RAP1 [Hardy et al. 1992b]. In an attempt to determine whether SIR proteins are also able to interact with this domain of RAP1, we extended our previous screen. RIF1 was isolated from a library generated by partial Sac3A digestion of total yeast genomic DNA inserted into the pGAD2 vector [Chien et al. 1991]. We therefore screened the two other reading frame libraries (in the pGAD1 and pGAD3 vectors), using a slightly longer RAP1 hybrid [LexA/RAP1(635–827)] and the reporter strain CTY10-5D [see Materials and methods]. From ~200,000 independent transformants, >30 clones were isolated that activated the reporter gene in a LexA/RAP1(635–827)-dependent manner. Southern blotting and partial DNA sequence analysis revealed that RIF1 had not been reisolated in either of these screens and indicated that two clones contained parts of the SIR3 and SIR4 genes.

The GAD/SIR3 clone that we isolated encodes the carboxy-terminal two-thirds of SIR3 (from amino acids 307 to 978) fused in-frame to the GAD-coding sequences. It is worth noting that this hybrid does not contain the amino acids affected in SIR3 suppressor mutations that restore silencing in histone H4 (HF2) mutant strains [Johnson et al. 1990]. The GAD/SIR4 clone contains only the carboxy-terminal 11% of the protein (from amino acids 1204 to 1358), also fused in-frame to GAD-coding sequences. This small carboxy-terminal fragment of SIR4 is sufficient to allow self-association as judged by the two-hybrid system [Chien et al. 1991].

Table 1 shows that both GAD/SIR3 and GAD/SIR4 require that RAP1 sequences be fused to LexA for activation to occur, as they fail to activate with either LexA
Interaction of SIR proteins with RAP1

**Table 1. Identification of SIR3 and SIR4 as RAP1-interacting proteins by the two-hybrid system**

| Hybrid | DNA-binding domain | activation domain | Transcriptional activation* |
|--------|--------------------|-------------------|-----------------------------|
| LexA/RAp1 [635–827] | GAD/SIR3 [307–978] | + + + |
| LexA | GAD/SIR3 [307–978] | – |
| LexA/RAp1 [635–827] | GAD | – |
| LexA/RAp1 [635–827] | SIR3–2μ | – |
| LexA/RAp1 [635–827] | GAD/SIR4 [1204–1358] | + |
| LexA | GAD/SIR4 [1204–1358] | – |
| LexA/RAp1 [635–827] | GAD | – |
| LexA/RAp1 [635–827] | SIR4–2μ | – |

*[+] Signal (strong blue color) detected in ~3–5 hr in β-gal assay on nitrocellulose filters; [+ +] signal detected after 8–12 hr of incubation; (–) no signal detected after 24 hr of incubation.

**The SIR3 protein interacts with itself and SIR4 in the two-hybrid system**

Previous genetic studies have suggested that the SIR3 and SIR4 proteins might interact with each other [Ivy et al. 1986; Marshall et al. 1987]. In addition, SIR4 has been shown to interact with itself in the two-hybrid system [Chien et al. 1991]. We thus decided to use this method to ask directly whether SIR3 and SIR4 can interact with each other and whether SIR3 can self-associate as does SIR4. Two LexA/SIR3 fusions were constructed, one coding for nearly the full-length SIR3 protein (amino acids 2–978) and the other for the carboxy-terminal two-thirds of SIR3 (amino acids 307–978). This shorter hybrid contains the same region of SIR3 that is fused to the activation domain of GAL4 in the clone isolated from the pGAD1 fusion library. Using the GAD/SIR3 and GAD/SIR4 clones isolated from the library screen, we were able to demonstrate a specific interaction with both LexA/SIR3 hybrids (Table 2). The shorter LexA/SIR3 hybrid appeared to interact more strongly than the larger hybrid with both GAD/SIR3 and GAD/SIR4. We have not investigated the cause of this difference, which might be attributable to differences in protein stability or folding, or to an inhibitory effect of the SIR3 amino terminus. In the same experiment, GAD/RIF1 was tested and no interaction with SIR3 was detected [data not shown]. The interaction between SIR3 and SIR4 was confirmed by constructing two LexA/SIR4 fusions, coding for 33% or 7% of the carboxyl terminus of SIR4, corresponding to the GAL4 DNA-binding domain fusions used previously to demonstrate SIR4 self-association [Chien et al. 1991]. Both of these LexA/SIR4 hybrids interact strongly with the GAD/SIR3 [307–978] clone (Table 2). These results demonstrate that the carboxyl termini of SIR3 and SIR4 can interact with each other, as well as with RAP1. The formation of both homodimers and heterodimers (or higher multimers) by SIR3 and SIR4 suggests that these two proteins may form a complex containing at least four subunits.

**SIR3 binds to RAP1 in vitro**

To ask whether the SIR3 protein can interact directly with RAP1 we used an in vitro protein-binding assay. Sequences encoding the carboxyl terminus of RAP1 were fused to the glutathione S-transferase [GST] gene, and the resulting hybrid proteins were expressed in *Escherichia coli* and partially purified by binding to glutathione-agarose beads. As a source of SIR3 protein the entire SIR3-coding sequence was transcribed in vitro using T7 polymerase and translated in vitro using a rabbit reticulocyte lysate in the presence of 35S-labeled methionine [see Materials and methods for details]. Labeled SIR3 pro-

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**Table 2. SIR3–SIR3 and SIR3–SIR4 interactions detected by the two-hybrid system**

| Hybrid | DNA-binding domain | activation domain | Activation (β-gal Units) |
|--------|--------------------|-------------------|-------------------------|
| LexA/SIR3 [2–978] | GAD | 3.0 |
| LexA/SIR3 [2–978] | GAD/SIR3 [307–978] | 49 |
| LexA/SIR3 [2–978] | GAD/SIR4 [1204–1358] | 77 |
| LexA/SIR3 [307–978] | GAD | 3.0 |
| LexA/SIR3 [307–978] | GAD/SIR3 [307–978] | 1066 |
| LexA/SIR3 [307–978] | GAD/SIR4 [1204–1358] | 1938 |
| LexA/SIR4 [839–1358] | GAD | 5 |
| LexA/SIR4 [839–1358] | GAD/SIR3 [307–978] | 1171 |
| LexA/SIR4 [1252–1358] | GAD | 49 |
| LexA/SIR4 [1252–1358] | GAD/SIR3 [307–978] | 591 |
tein was incubated with a set of GST/RAP1 hybrid proteins bound to glutathione–agarose beads. After the beads were washed, bound proteins were eluted and analyzed by SDS-PAGE and autoradiography. As shown in Figure 1, SIR3 protein is able to interact with GST/RAP1 fusions with amino-terminal endpoints between amino acids 562 and 653. (The genetic properties of the corresponding LexA/RAP1 hybrids are described in detail below.) We believe that these interactions are specific because binding is dependent on RAP1 sequences, and labeled RAP1 protein, tested in the same conditions, fails to bind to the GST/RAP1 hybrids (Fig. 1). In our assay conditions binding by SIR3 is more efficient with the longer GST/RAP1 fusion proteins, but an interaction is detectable with the shorter hybrids. These results indicate that the SIR3 protein can interact physically with the carboxyl terminus of RAP1 in the absence of other yeast proteins. So far, we have been unable to detect an interaction in vitro between the RAP1 carboxyl terminus and SIR4. Consequently, we do not know whether the interaction detected in vivo is direct.

**RAP1 sequences required for interaction with SIR3, SIR4, and RIF1**

A series of LexA/RAP1 constructs with different RAP1 fusion points (between amino acids 635 and 702) was used to determine the amino-terminal boundary of RAP1 sequences required for the interaction with SIR3, SIR4, and RIF1. The primary conclusion to be drawn from these experiments (see Fig. 2) is that amino acids 679–827 of RAP1 are sufficient for the interaction with both SIR3 and RIF1. However, SIR4 is able to interact only with the LexA/RAP1[635–827] hybrid. All of the hybrids that interact with RIF1 appear to do so with equal strength (β-galactosidase level of ~5000–7000 units). In contrast, the interaction between SIR3 and this same set of LexA/RAP1 hybrids varies in a manner not strictly related to the length of RAP1 sequences present. For example, the RAP1–SIR3 interaction seems to be stronger for three smaller LexA/RAP1 hybrids [amino-terminal endpoints at 655, 667, and 679], than for two longer protein fusions [endpoints at 647 and 653]. However, the longest fusion tested [LexA/RAP1(635–827)] gives a value close to that of the three shorter hybrids. Because the RAP1–RIF1 interactions are constant, we imagine that the variability seen with GAD/SIR3 reflects a true difference in the interaction with this set of LexA/RAP1 hybrids rather than a variation in the amount or stability of the different hybrids (see below).

Next, we asked whether mutations in the RAP1 carboxyl terminus also affect protein–protein interactions detected by the two-hybrid system. To do this, we used LexA/RAP1 hybrids with amino-terminal fusion junctions at amino acids 647, 653, 655, 667, and 679 of RAP1. All of these LexA/RAP1 hybrids fail to activate transcription by themselves and are affected slightly or not at all by carboxy-terminal mutations. Data in Table 3 show that incorporation of a small linker insertion mutation at amino acid 825 of RAP1 [825*], which results in the addition of 5 amino acids very near the carboxyl termi-
The first value given in each entry is the number of β-gal units measured for the interaction. The number in parenthesis is the fold decrease relative to the value obtained for the corresponding wild-type LexA/RAP1 hybrid (i.e., not containing the 825* mutation).

Mutation of SIR2, SIR3, SIR4, or RIF1 increases the transcriptional activation potential of LexA/RAP1 hybrids

Previous studies of a series of GAL4 DNA-binding domain/RAP1 hybrids had identified a transcriptional activation domain in RAP1 (between amino acids 630 and 695) that partially overlaps its carboxy-terminal silencing domain [Hardy et al. 1992a]. We reasoned that if the native SIR proteins do interact directly with the carboxyl terminus of RAP1 they might modulate its ability to function as a transcriptional activator. To test this idea, we examined the activation properties of a related series of LexA/RAP1 hybrid proteins in strain CTY10-5D, which contains a LexA operator- lacZ reporter gene, and each of five different derivatives of this strain containing gene disruptions of either SIR1, SIR2, SIR3, SIR4, or RIF1.

LexA/RAP1 hybrids that have amino-terminal endpoints at amino acids 562 and 630 have an increased ability to activate transcription of the reporter gene in sir3, sir4, and rif1 mutants [Fig. 3, rows 1 and 2]. The level of β-galactosidase activity in these strains is increased 2- to 3.5-fold compared with that of wild-type cells. The next hybrid in the series, LexA/RAP1(635–827), shows a particularly striking effect. This hybrid is unable to activate transcription in wild-type cells but converts into an activator in four of the mutant strains tested [Fig. 3, row 3]. The β-galactosidase levels increase 7-fold in the sir2 mutant, 16-fold in the rif1 mutant, and almost 30-fold in the sir3 and sir4 mutants relative to the SIR+ RIF+ reporter strain. It is interesting to note that a sir1 mutation had no effect on this or any other hybrid tested. Hybrids with RAP1 amino-terminal endpoints at amino acids 647 and 653 did not activate transcription in any genetic background tested [Fig. 3, rows 4 and 5], nor did shorter hybrids with endpoints at amino acids 655, 667, 679, 691, or 702 [data not shown]. The effect of the sir2, sir3, sir4, and rif1 mutations on LexA/RAP1 hybrids appears to be specific as no effect is observed with two different and unrelated transcriptional activators, LexA/GAL4 [Fig. 3, bottom] and LexA/SNF6 [data not shown].

Mutation of the RAP1 carboxyl terminus increases activation by LexA/RAP1 hybrids and abolishes SIR and RIF1 repression

Taken together, the results described above suggest that the repressing effect of SIR and RIF1 genes on RAP1

![DNA-Binding Domain Hybrid](image)

| DNA-Binding Domain Hybrid | Activation in Wild-Type Cells (β-gal Units) | Fold increase in mutant cells |
|---------------------------|------------------------------------------|-----------------------------|
|                           |                                          | sir1 | sir2 | sir3 | sir4 | rif1 |
| LexA RAP1(562-827)        | 283                                      | 0.8  | 1.3  | 2.0  | 1.9  | 2.6  |
| LexA RAP1(630-827)        | 363                                      | 0.9  | 1.1  | 2.0  | 2.9  | 3.5  |
| LexA RAP1(635-827)        | 6.0                                      | 1.1  | 7.0  | 28   | 27   | 16   |
| LexA RAP1(647-827)        | 2.0                                      | 1.0  | 0.9  | 1.1  | 1.3  | 1.1  |
| LexA RAP1(653-827)        | 2.0                                      | 1.0  | 1.0  | 0.8  | 1.1  | 1.0  |
| LexA GAL4(768-881)        | 10,000                                   | 1.0  | 1.0  | 1.0  | 1.0  | 1.0  |

Interaction of SIR proteins with RAP1

**Figure 3.** Transcriptional activation by Lex/RAP1 hybrids is increased by mutations in SIR2, SIR3, SIR4, and RIF1. Transcriptional activation measured in β-gal units in wild-type cells (CTY10-5D) is normalized to a value of 10,000 units for LexA/GAL4(768-881) [LexA/GAD], which was included as a control in all experiments. Values for the hybrids in mutant cells are expressed as a fold increase over that in wild-type cells and are normalized to a value of 10,000 units for LexA/GAD in that strain. LexA/GAD values in all mutant strains were essentially indistinguishable from the wild-type parent.
hybrids is attributable to a direct interaction between the SIR and RIF1 proteins and the RAP1 carboxyl terminus. A prediction of this model is that carboxy-terminal mutations in RAP1 might increase the activation potential of some hybrids and relieve SIR and RIF repression.

To test this idea, we introduced carboxy-terminal mutations into the series of hybrids shown in Figure 2. As shown in Figure 4, the 825* linker-insertion mutation causes an 8- and 10-fold increase in the activation potential of hybrids with amino-terminal endpoints at positions 562 and 630, respectively. The same insertion mutation has an even more dramatic effect on the LexA/RAPI(635–827) hybrid, increasing its ability to activate transcription by ~200-fold. In contrast, the 825* mutation has no effect on hybrids beginning at amino acids 647 or 653, both of which still fail to activate the reporter gene. It is worth noting that the wild-type versions of these two hybrids [LexA/RAPI(647–825*) and LexA/RAPI(653–827)] were also unaffected by sir and rif1 mutations (Fig. 3). A similar pattern is seen with LexA/RAPI hybrids truncated at amino acid 799 of RAP1 [Fig. 4, bottom]. In this case, however, hybrids beginning at amino acids 647 and 653 are partially activated by the 28-amino-acid truncation mutation. To ask whether any of the carboxy-terminal mutant LexA/RAPI hybrids [Fig. 4] are subject to repression by SIR or RIF1 proteins, we examined their activation properties in the series of mutant strains described above. Unlike the wild-type hybrids, none of the carboxy-terminal mutant hybrids were affected by mutations in sir1, sir2, sir3, sir4, or rif1 (data not shown).

**RAP1–SIR3, SIR3–SIR3, and SIR4–SIR3 interactions do not require endogenous SIR proteins**

The simplest explanation for the genetic results described above is that RAP1, SIR3, and SIR4 interact directly with one another. This notion is strongly supported in the case of the RAP1–SIR3 interaction, as biochemical experiments show that these two proteins can interact in vitro in the absence of other yeast proteins. However, it is possible that some or all of the two-hybrid interactions that we observe in vivo require the mediation of other SIR proteins or are modulated in some way by SIR proteins. To test this possibility we used derivatives of the CTY10-5D reporter strain described above, which contain mutations of SIR1, SIR2, SIR3, or SIR4. The SIR dependency of the interaction between RAP1 and SIR3 was analyzed by using LexA/RAPI hybrids with amino-terminal endpoints at amino acids 647 and 653. The interactions between SIR3 and itself and between SIR3 and SIR4 were tested by using the two LexA/SIR3 hybrids (see Table 2) and the GAD/SIR hybrids isolated from the library screens (see Table 1). It has not been possible to extend this analysis to the interaction between RAP1 and SIR4, as the only hybrid that interacts with SIR4 [LexA/RAPI(635–827)] becomes active when SIR2, SIR3, or SIR4 are mutated (Fig. 3).

None of the two-hybrid interactions tested require SIR gene function [data not shown]. However, the interaction between RAP1 and GAD/SIR3 appears to be slightly weakened by the absence of SIR2, particularly in the case of the longer LexA/RAPI hybrids where β-galactosidase levels in the sir2 mutant are 40–50% that of wild type [data not shown]. We conclude from these results that the RAP1–SIR3, SIR3–SIR3, and SIR4–SIR3 interactions detected by the two-hybrid assay are either direct or require the activity of other genes that we have not tested. It is worth emphasizing that a rather small carboxy-terminal domain of SIR4, consisting of only 97 amino acids, appears to be sufficient to confer both self-association and binding to SIR3, as both interactions are independent of endogenous SIR gene function. At present, we cannot rule out the possibility that RAP1 mediates these SIR interactions.

**RIF1 and SIR3 compete for binding to the carboxy terminus of RAP1**

It has been shown recently that SIR proteins and RIF1 may have opposing functions with respect to telomere position effect and length regulation [Kyrion et al. 1993; Palladino et al. 1993]. Here, we have shown that both SIR3 and RIF1 interact with a common set of LexA/RAPI hybrid proteins in the two-hybrid system [Fig. 2]. We therefore decided to ask whether RIF1 competes with SIR3 for binding to the RAP1 carboxy terminus. To test

| DNA-Binding Domain Hybrid | Activation in Wild-Type Cells (β-gal Units) | Fold Increase over Wild-type |
|---------------------------|-------------------------------------------|-----------------------------|
| LexA RAP1(562–825)        | 2235                                      | 7.9                         |
| LexA RAP1(630–825)        | 3678                                      | 10.1                        |
| LexA RAP1(635–825)        | 1395                                      | 232                         |
| LexA RAP1(647–825)        | 2                                         | 1.0                         |
| LexA RAP1(653–825)        | 2                                         | 1.0                         |
| LexA RAP1(653–799)        | 1642                                      | 5.8                         |
| LexA RAP1(652–799)        | 5207                                      | 14.3                        |
| LexA RAP1(630–799)        | 3641                                      | 607                         |
| LexA RAP1(635–799)        | 60                                        | 30                          |
| LexA RAP1(647–799)        | 110                                       | 55                          |
| LexA RAP1(653–799)        |                                           |                             |

Figure 4. Transcriptional activation by LexA/RAPI hybrids is increased by mutation of the RAP1 carboxy terminus. A series of LexA/RAPI hybrids with different amino-terminal RAP1 fusion endpoints (see Fig. 2) and either a linker insertion mutation two amino acids before the carboxy terminus of RAP1 [825*] or a truncation of the carboxy terminus at amino acid 799 were assayed as before (Fig. 2). The fold increase in activation compared to the corresponding hybrids with no mutation at the carboxy terminus (see Fig. 2) is given in the right-hand column.
this possibility we examined the interaction between GAD/SIR3 and a series of LexA/RAP1 hybrids in a strain containing a disruption of the RIF1 gene. As shown in Table 4, the interaction between GAD/SIR3 and LexA/RAP1 hybrids with amino-terminal endpoints between amino acids 647 and 655 appears to be stronger in rif1 mutant cells than in the isogenic RIF1 + parent strain. Furthermore, the variability in the strength of the RAP1-SIR3 interaction is abolished, with the level of β-galactosidase reaching a nearly constant value of ~5000–6000 units. Only the interaction with the shortest hybrid tested [LexA/RAP1(679–827)] is unaffected by the rif1 mutation. These results suggest that the native SIR3 and RIF1 proteins might compete for binding to the RAP1 carboxyl terminus.

Mutations in the RAP1 carboxyl terminus that diminish SIR3 binding affect silencing at HMR and at telomeres

We have shown above that mutations in the carboxyl terminus of RAP1 affect the ability of LexA/RAP1 hybrids to interact with SIR3 in the two-hybrid system (Table 3). If this putative RAP1–SIR3 interaction is important for silencing in vivo, we would expect that the same carboxy-terminal mutations in the context of native RAP1 would lead to silencing defects. To test this idea, we constructed strains in which the only copy of RAP1 contained one of four different carboxy-terminal mutations. These strains also contain either a TRP1 reporter gene under control of the HMR-E silencer, or a URA3 gene adjacent to a telomere created at the ADH4 locus (Gottschling et al. 1990). Because the HMR-E silencer is a redundant regulatory element (Brand et al. 1987; Kimmerly et al. 1988), we also tested a TRP1 reporter strain carrying a mutation in the ACS (A element) or the ABF1 site (B element) at the silencer. The hmr3A silencer is particularly sensitive to mutations in RAP1 (Sussel and Shore 1991; Sussel et al. 1993). Silencing of the telomeric URA3 gene was monitored by assaying for growth in the presence of 5-fluoro-orotic acid (FOA), which kills cells expressing URA3.

In strains containing the rap1-7 mutation [a linker insertion at amino acid 825, referred to earlier as 825*] a very slight defect is observed in silencing at HMR, as indicated by some growth in the absence of tryptophan, but only when the A element at HMR-E is mutated (Fig. 5A, row 3). However, the rap1-7 mutation causes a significant loss of telomeric silencing, as evidenced by a 1000-fold drop in FOA resistance in the strain containing a telomeric URA3 gene (Fig. 5B, row 3). More severe

| DNA-binding domain hybrid | Activation in wild-type cells | Activation in rif1 mutants | Fold increase |
|---------------------------|-------------------------------|---------------------------|---------------|
| LexA/RAP1 (647–827)       | 325                           | 4612                      | 13.7          |
| LexA/RAP1 (653–827)       | 487                           | 5335                      | 9.7           |
| LexA/RAP1 (655–827)       | 878                           | 5662                      | 6.4           |
| LexA/RAP1 (667–827)       | 3562                          | 6124                      | 1.7           |
| LexA/RAP1 (679–827)       | 4680                          | 5769                      | 1.2           |

All strains contain GAD/SIR3 (307–978).
mutations of the RAP1 carboxyl terminus, truncations at amino acids 716, 703, and 695 (rap1-8, rap1-9, and rap1-10, respectively) cause partial derepression of the hmrAB silencer and complete derepression of the hmrAA silencer [Fig. 5A, bottom two panels, rows 4–6]. However, the wild-type HMR-E silencer appears to be unaffected by these rap1 mutations [Fig. 5A, top panel, rows 4–6]. All three of these rap1 deletions also result in a complete loss of telomeric silencing [Fig. 5B, rows 4–6]. An unrelated deletion of the amino terminus of RAP1 (rap1-6) shows no defect in either HMR or telomeric silencing, indicating the specificity of the carboxy-terminal deletions [Fig. 5A,B, row 2]. Taken together, these data support the notion that the RAP1–SIR3 interaction is important for silencing, as there is a correlation between the strength of this interaction as measured by the two-hybrid system and the efficiency of repression at telomeres and RAP1-dependent silencers.

Discussion

The role of RAP1 in transcriptional silencing

Previous genetic studies have demonstrated the importance of the RAP1 carboxyl terminus in transcriptional silencing at HM loci and telomeres [Kurtz and Shore 1991; Sussel and Shore 1991; Hardy et al. 1992a; Kyrion et al. 1993]. The results reported here provide evidence for a molecular mechanism to explain the role of this RAP1 domain in silencing. We propose that RAP1 works at HM silencers and at telomeres by recruiting a complex of SIR proteins via direct protein–protein interactions with its carboxyl terminus. This conclusion is based on four independent lines of evidence. First, SIR3 and SIR4 hybrid proteins interact with the RAP1 carboxyl terminus in the two-hybrid system, and SIR3 can interact with both itself and SIR4. Second, SIR3 protein interacts with the RAP1 carboxyl terminus in vitro in the absence of other yeast proteins. Third, mutation of the endogenous SIR genes increases the activation potential of RAP1 in the context of LexA/RAP1 hybrid proteins, providing independent evidence that the native SIR proteins can interact directly with the RAP1 carboxyl terminus. Finally, mutations in RAP1 that reduce or abolish the RAP1–SIR3 interaction in the two-hybrid system have a corresponding effect on silencing when incorporated into the native RAP1 protein. The ability of the RAP1 carboxyl terminus to bind SIR proteins may be sufficient to establish silencing, as hybrid proteins containing only this region of RAP1 fused to the GAL4 DNA-binding domain can establish repression when targeted to mutated silencers containing GAL4 binding sites [S. Buck and D. Shore, unpubl.]. We do not have enough information at present to know whether SIR4 interacts directly with either SIR3 or RAP1.

On the basis of the multiple interactions detected among RAP1, SIR3, and SIR4, we propose a model in which SIR3 and SIR4 form a heteromeric complex that interacts with RAP1 at HM silencers and telomeres. The fact that both SIR3 and SIR4 [Chien et al. 1991] appear to interact with themselves and with each other suggests that these two proteins might be capable of forming a large complex containing at least two copies of each protein. One can also imagine that this putative SIR3–SIR4 complex is capable of initiating the assembly of a structure that extends along the chromatin fiber from silencers and telomeres. Such a polymerization model would provide an explanation for the ability of HM silencers and telomeres to exert their repressive effects at a considerable distance. Results from previous studies are consistent with a structural role for SIR3 and SIR4, as both exhibit striking gene dosage effects. For example, excess SIR4 gene dosage or overexpression of a carboxy-terminal fragment of the protein can interfere with silencing at HM loci and telomeres [Ivy et al. 1986; Marshall et al. 1987; Sussel and Shore 1991; Renaud et al. 1993]. However, a single extra copy of SIR4 can suppress different silencing defects at HMR (Sussel et al. 1993). On the other hand, increased gene dosage of SIR3, but not SIR4, can increase the frequency of silencing and the extent of propagation of silent chromatin from telomeres [Renaud et al. 1993]. Taken together, these data suggest that the precise roles of SIR3 and SIR4 may differ at both HM loci and telomeres.

At present, the role of SIR2 in the putative RAP1–SIR complex is unclear. Mutations in SIR2 have only a modest effect on activation by LexA/RAP1 hybrids compared with those of SIR3 and SIR4, and SIR2 was not identified in a two-hybrid screen for RAP1-interacting proteins. Perhaps SIR2 interacts with either SIR3, SIR4, or both proteins but does not contact RAP1 directly. It should be noted that the role of SIR2 in the cell is apparently different from that of either SIR3 or SIR4, as sir2 mutations affect rDNA recombination whereas mutations in either SIR3 or SIR4 do not [Gottlieb and Esposito 1989]. The recent observation that overexpression of SIR2 results in histone deacetylation in vivo [Braunstein et al. 1993] suggests that SIR2 may have a more direct role in modifying chromatin. One might imagine, therefore, that SIR2 is either loosely or transiently associated with a RAP1–SIR3/SIR4 complex or that SIR2 recognizes an altered chromatin conformation created by the action of this complex.

Data presented here raise the possibility that the native RIF1 and SIR3 proteins bind competitively to the RAP1 carboxyl terminus. We interpret this result in terms of a model in which RIF1 binding to RAP1 exerts a negative effect on silencing that primarily affects telomeres. Consistent with this model, rif1 mutants have been shown to display improved telomeric repression [Kyrion et al. 1993]. Recent studies of rap1 mutants provide further support for this model and clarify the different roles of RAP1 at HM loci and telomeres [S. Buck and D. Shore, unpubl.].

Protein–protein interactions and the context dependence of RAP1 function

Previous studies clearly indicate that the regulatory function of RAP1 (either repression or activation) is de-
interactions at these genes. However, GCR1 appears to make a direct protein–protein interaction with RAP1 and can function at some RAP1-containing promoters in the absence of its own DNA-binding domain [Tornow et al. 1993]. Furthermore, it seems that not all GCR1-dependent promoters contain binding sites for this protein [Santangelo and Tornow 1990]. The GAL11 protein also helps RAP1 to activate at many different promoters yet does not appear itself to be a DNA-binding protein [Nishizawa et al. 1990]. Therefore, protein–protein interactions between coactivators [e.g., GCR1 and GAL11] and RAP1 may play an important role at many promoters. How these interactions are targeted to promoters without apparently interfering with the silencing function of RAP1 at HM loci and telomeres is not well understood.

Given the fact that RAP1 appears to be much more abundant than any of the SIR proteins [S.M. Gasser, pers. comm.], it may seem surprising that deletion of SIR genes can have such a profound effect on the activity of LexA/RAPI hybrids. We suggest two reasons to account for these results. First, several of the LexA/RAPI hybrids that we have studied may have more inherently favorable interactions with silencing factors because of the absence of residues important for transcriptional activation. The particularly dramatic effect of sir and rif1 mutations on the LexA/RAPI[635–827] hybrid can thus be explained by the fact that this hybrid is actually missing only 5 amino acids from the amino terminus of the RAP1 activation domain. Second, it seems likely that the LexA operator–lacZ reporter gene used in these studies provides a useful system to examine RAP1 protein–protein interactions precisely because it lacks auxiliary regulatory elements normally associated with RAP1-binding sites at either promoters or HM silencers. The presence of eight consecutive LexA operators upstream of the lacZ reporter gene may create a situation resembling that at telomeres, where consecutive RAP1-binding sites appear to favor interactions with SIR proteins. This feature of the artificial reporter may allow one to detect RAP1–SIR interactions that would not normally occur at natural promoters because of competition by other regulatory proteins bound to nearby sites or the action of coactivators that may themselves participate in other protein–protein interactions at promoters. By modifying the reporter gene system to include other regulatory elements found either at promoters or HM silencers we may begin to obtain insights into the features of these elements that favor particular sets of RAP1 protein–protein interaction at different chromosomal loci.

**RAP1 and nuclear localization of telomeres**

Recent studies have shown that yeast telomeres appear to be clustered in the nucleus and localized to the nuclear periphery, perhaps directly attached to the nuclear envelope [Klein et al. 1992]. Strikingly, mutation of either SIR3 or SIR4 abolishes the perinuclear localization of telomeres and may also reduce their tendency to aggregate [Palladino et al. 1993]. Our work provides a plausible molecular explanation for these observations. We

Interaction of SIR proteins with RAP1
suggest that the association of SIR3 and SIR4 with RAP1 protein bound to the terminal [C_{1-3}A] sequences at telomeres leads, either directly or indirectly, to their attachment to the nuclear membrane. The SIR complex may bind directly to the nuclear envelope via a carboxy-terminal domain of SIR4 that is homologous to nuclear lamins (Difffley and Stillman 1989b). The ability of SIR3 and SIR4 to interact with themselves and each other may also explain the apparent aggregation of telomeres in vivo. Finally, attachment of RAP-bound chromosomal lamins (Diffley and Stillman 1989b). The ability of SIR3 attachment to the nuclear membrane. The SIR complex lomeres leads, either directly or indirectly, to their at-

Moretti et al.

Materials and methods

Media and strains

Growth and manipulation of yeast strains was done according to standard procedures (Rose et al. 1990). The yeast strain CTY10-5D (MATa ade2-1 trpl-901 leu2-3,112 his3-200 gal4 gal80 URA3::lexA op–lacZ) was used in all studies involving LexA hybrid proteins. This strain (a gift of C.-T. Chien and R. Sternglanz, State University of New York, Stony Brook) contains a lacZ reporter gene with 4 ColE1 operators [or eight binding sites for LexA dimers] inserted upstream of the transcription start site of a GALI–lacZ gene integrated at the URA3 locus. HIS3 gene disruptions of SIR1, SIR2, SIR4 (Kimmerly and Rine 1987), SIR3, and RFI1 in strain CTY10-5D were obtained by gene replacement (Rothstein 1991) and confirmed by Southern blotting. The SIR3 disruption was constructed by deleting a BglII–XhoI fragment [encoding amino acids 108–945] and replacing it with a fragment containing the HIS3 gene. The RFI1 disruption was constructed by replacement of an MluI–XhoI fragment of the gene with HIS3. This construct removes all of the amino-terminal RFI1-coding sequence, up to amino acid 1744. Libraries of partial Sau3Al-digested yeast genomic DNA in the vectors pGAD1 and pGAD3 (Chien et al. 1991) were generously provided by P. Bartel and S. Fields (State University of New York, Stony Brook). Plasmid DNAs were rescued from CTY10-5D by transformation into the E. coli strain BA1 (thr leu6 thi thyA trpC1117 hisB strB)3, selecting simultaneously for ampicillin resistance and leucine prototrophy. Strains containing the rap1-6, rap1-7, rap1-8, rap1-9, and rap1-10 alleles are MATa haploids in which the chromosomal copy of RAP1 is a rap1::LEU2 deletion/disruption that removes all of the RAP1 amino-terminal sequences up to amino acid 760. The mutant rap1 allele is present in these cells on a HIS3 CEN plasmid. These strains are derived from a series of hmr::TRP1 strains or a URA3–Tel VIII strain described previously (Susset and Shore 1991; Chien et al. 1993). They are all otherwise isogenic to strain W303-1B [MMATa HMRa ade2-1 can1-100 his3-11,15 leu2-3,112 trpl-1 ura3-1] (Thomas and Rothstein 1989).

Plasmids

The LexA protein and LexA/RAP1, LexA/SIR3, LexA/SIR4, and LexA/GAL4 [768–881] hybrid proteins were expressed from plasmid pBTM116 (2 µ origin, TRP1, pADH1–lexA; a gift of P. Bartel and S. Fields). Most RAP1 carboxy-terminal fragments were obtained from a set of C_{992–RAP1} fusions described previously (Hardy et al. 1992a) as EcoRI–PstI, BamHI–PstI, or Smal–PstI fragments. The RAP1(647–827) fragment was obtained from a XhoI linker insertion mutation. The RAP1(679–827) and RAP1(691–827) fragments were designed by digestion at a BsrBI or HindIII site in RAP1, respectively, followed by Klenow fragment repair of the ends. The carboxy-terminal deletion of RAP1 at amino acid position 799 was generated by digestion at a BslI site followed by Klenow fragment repair of the end. LexA/GAL4[768–881] [LexA/G43] was constructed by cloning an EcoRI–BamHI fragment from pGAD3 (Chien et al. 1991), which contains the GAL4 activation domain [amino acids 768–881], into EcoRI–BamHI-cut pBTM116. LexA/SNF6 was a gift of B. Laurent (Laurent and Carlson 1992). The carboxy-terminal mutation of RAP1 at amino acid position 825 was obtained by digestion at a EcoO109I site, followed by Klenow fragment repair of the ends and insertion of a 12-mer XhoI linker. The LexA/SIR3(2–978) fusion was created by a three-way ligation in which an EcoRI–Clal fragment of SIR3 generated by PCR [a generous gift of C.-T. Chien] was ligated to a Clal–BamHI fragment of the gene. The LexA/SIR3(307–978) fusion was created by joining SIR3 sequences [BglII–BamHI] coding for amino acids 307–978 to Lexa sequences in pBTM116. The LexA/SIR4(389–1358) and (1252–1358) fusions were created by ligating EcoRI–SalI fragments from plasmids pCTC17 and pCTC23 (Chien et al. 1991) into EcoRI–SalI-cut pBTM116. In some cases, the reading frame at the fusion junction of the pBTM116 plasmid was altered by filling in the EcoRI and BamHI sites within the polylinker of this vector. More detailed information about these constructs is available upon request.

Constructs for plasmid shuffling of rap1 mutant alleles were created using the pRS313 vector [Sikorski and Hieter 1989]. Plasmids expressing the rap1 alleles 6–10 were constructed from a series of XhoI linker insertion mutations cloned into pRS313. Plasmids expressing the rap1 alleles 8, 9, and 10 alleles contain the ADH1 terminator in place of the RAP1 terminator.

GST/RAP1 fusions were created in a version of the plasmid pGEX2TK in which the polylinker of the vector has been substituted with the polylinker of plasmid pLC20R [Marsh et al. 1984]. The RAP1 carboxy-terminal fragments were obtained from a set of C_{992–RAP1} fusions described previously [Hardy et al. 1992a]. The pT7–SIR3 construct has been made in plasmid pT7-BSalI, using a version of the SIR3 gene in which a Neol site has been created at codon 1. The pT7–RAP1 was made by site-directed mutagenesis of the RAP1 ATG to introduce a Neol site followed by cloning of a Ncol–XhoI fragment into pT7-BSalI [Brigati et al. 1993].

Isolation and identification of SIR3 and SIR4 using the two-hybrid system

The yeast LexA operator–lacZ reporter strain CTY10-5D was cotransformed with a plasmid expressing the LexA/RAP1[635–827] hybrid and a library of genomic DNA fragments in the pGAD3 or pGAD1 expression vectors, using the high-efficiency transformation method of Schiestl and Geitz (1989). Transformants were selected on SC-Trp-Leu medium at 30°C and screened for β-galactosidase activity using a nitrocellulose filter assay [Breeden and Nasmyth 1985]. Positive [blue] colonies were identified, purified, and retested. DNA prepared from positive clones was transformed into the LexA E. coli strain BA1, and Amp? Leu+ transformants were selected. Plasmid DNA was tested by transformation into the yeast reporter strain CTY10-5D containing no plasmid, the LexA/RAP1[635–827] plasmid, or one of three different control plasmids expressing LexA, LexA–lamin, or LexA–ADH1 hybrids. Clones that displayed LexA/RAP1[635–827]-dependent activation were sub-
jected to dideoxynucleotide sequencing using an oligonucleotide primer that hybridizes to the GAL4 activation domain sequences near the cloning junction (5'-TACCACTACAATGGATG-3').

Transcriptional activation assays

Transcriptional activation by LexA hybrids or LexA and GAD hybrid combinations was measured in strain CTY10-5D and its sir or rif1 mutant derivatives. Transformants were grown in selective liquid medium containing 0.05% glucose for 40 hr. Cells (5 ml) were pelleted, resuspended in 250 μl of Z buffer (60 mM Na2HPO4, 40 mM NaH2PO4, 10 mM KCl, and 1 mM MgSO4) containing 0.27% β-mercaptoethanol, and permeabilized by 3×1 min of rapid vortexing with 0.3 gram of glass beads. The suspension was centrifuged and 20–150 μl of the supernatant was used in a 1-ml α-galactosidase assay (Miller 1972; Breeden and Nasmyth 1987). Activities were normalized to protein concentration using the Bradford assay (Bradford 1976). Units of β-galactosidase activity were calculated by taking the average from at least three independent transformants of each construct. Values for independent transformants varied from each other by <20%. All numbers reported for LexA/RAP1 hybrids were normalized by using a value of 10,000 units for LexA/GAD in that strain. Activation by LexA/GAD in sir and rif1 mutants was essentially indistinguishable from that seen in the wild-type CTY10-5D parent strain.

In vitro protein-binding studies

GST/RAP1 fusion proteins were expressed in E. coli strain DH5α. Transformants were grown overnight in 10 ml of 2XTY medium containing 50 μg/ml of ampicillin, pelleted, grown for 3 hr in 50 ml of fresh medium, and induced for 1.5 hr with isopropyl-β-D-thiogalactopyranoside (IPTG) at 0.1 mM. The cells were then pelleted and resuspended in 1 ml of TEN buffer (100 mM Tris-HCl at pH 7.4, 1 mM EDTA, 50 mM NaCl) containing 1 mM phenylmethylsulfonylfluoride (PMSF) and 20 μg/ml of pepstatin A, and sonicated on ice for 15 sec. Insoluble material was pelleted at 10,000 rpm for 10 min in a Sorvall SS34. Supernatants were stored at −70°C or used immediately for adsorption on glutathione–agarose.

Typical binding reactions used 400 μl of crude bacterial extracts and 200 μl of glutathione–agarose slurry incubated at 4°C for 1 hr on a rocking platform. The agarose beads were washed five times with 1 ml of TEN buffer and resuspended in 100 μl of TEN buffer to obtain a 50% slurry. An aliquot (10 μl) of each purified fusion protein was diluted in an equal volume of 2× SDS sample buffer, heated for 3 min in boiling water, and analyzed by SDS-PAGE. Gels were stained with Coomassie blue.

SIR3 protein was synthesized in vitro using the TNT T7-coupled reticulocyte lysate system (Promega) according to the manufacturer’s instructions. An aliquot (5 μl) of the reaction was diluted with 20 μl of SDS sample buffer, heated for 3 min in boiling water, and analyzed by SDS-PAGE. The gels were treated with fixing solution for 30 min and Amplify (Amersham International) for 30 min, and then dried and exposed to X-ray film.

Typical protein-binding reactions were performed in 200 μl of binding buffer (20 mM HEPES at pH 7.5, 1 mM EDTA, 25 mM NaCl, 7 mM MgCl2, 0.05% NP-40, 17% glycerol), containing 20 μl of purified GST–RAP1 protein adsorbed to the 50% glutathione–agarose slurry (in binding buffer). The beads were preincubated with 100 μg/ml of BSA for 15 min, and incubated with 3 μl of in vitro-synthesized proteins for 1 hr. The beads were collected with a 5-sec pulse in a microcentrifuge and washed three times with 1 ml of binding buffer. Washed beads were resuspended in 30 μl of 2× SDS sample buffer, heated for 3 min in boiling water, and analyzed by SDS-PAGE. The gels were treated and exposed to X-ray film as before.

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P Moretti, K Freeman, L Coodly, et al.

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