Genetic analysis of RNA polymerase I unveiling new role of the Rpa12 subunit during transcription

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Most transcriptional activity of exponentially growing cells is carried out by RNA Polymerase I (Pol I), which produces a large rRNA precursor. The Pol I transcription cycle is achieved through complex structural rearrangements of the enzyme, revealed by recent structural studies. In the yeast *S. cerevisiae* the Pol I subunit Rpa49, particularly its C-terminal tandem winged helix domain (Rpa49Ct), is required supports both initiation and elongation of the transcription cycle. Here, we characterized novel extragenic suppressors of the growth defect caused by the absence of Rpa49. We identified suppressor mutations on the two largest subunits of Pol I, Rpa190 and Rpa135, as well as Rpa12. Suppressor mutants *RPA135-F301S* and *RPA12-S6L* restored normal rRNA synthesis and increased Pol I density on rDNA genes in the absence of Rpa49Ct. Most mutated residues cluster at an interface formed by the jaw in Rpa190, the lobe in Rpa135, and subunit Rpa12 when mapped on the structure of Pol I. Our genetic data in *S. cerevisiae* suggest a new role for Rpa12 at the jaw/lobe interface during transcription cycle.
**Introduction**

The nuclear genome of eukaryotic cells is transcribed by three RNA polymerases (Chambon, 1975). RNA polymerase II (Pol II) transcribes most of the genome and is responsible for all messenger RNA production. RNA polymerases III and I are specialized in the synthesis of a limited number of transcripts. RNA polymerase III (Pol III) produces small structured RNAs, including tRNAs and the 5S ribosomal RNA. RNA polymerase I (Pol I) produces a single transcript, the large polycistronic precursor (the 35S pre-rRNA in yeast), processed by multiple successive steps into the mature rRNAs (25S, 18S, and 5.8S in yeast). Despite producing a single transcript, Pol I is by far the most active eukaryotic RNA polymerase, responsible for up to 60% of the total transcriptional activity in exponentially growing cells (Warner, 1999). Pol I rRNA synthesis constitutes the first step of ribosome biogenesis and is a rate limiting process for cell growth. The strongly transcribed rRNA genes can be visualized using the DNA spread method developed by Miller et al, 1969, in which the 35S rRNA genes (rDNA) adopt a “Christmas tree” conformation, with up to 120 polymerases per transcribed gene (Miller & Beatty, 1969). Altered cell proliferation, often associated with a modified rate of rRNA synthesis via the deregulation of Pol I activity, has been associated with various types of cancer (Drygin et al, 2010).

The full subunit composition and structural data are now available for the three nuclear RNA polymerases of the budding yeast *Saccharomyces cerevisiae* (Fernández-Tornero et al, 2013; Engel et al, 2013); (Hoffmann et al, 2015; Cramer et al, 2001). Pol I contains a core of shared or homologous subunits that are largely conserved in eukaryotes and archaea, as for the other two nuclear RNA polymerases (Werner, 2008). The two largest subunits (Rpa190 and Rpa135) carry the catalytic site. Rpb5, Rpb6, Rpb8, Rpb10, and Rpb12 are shared with Pol II and Pol III, whereas Rpc40 and Rpc19 are only shared with Pol III. This nine-subunit core is associated with the stalk, a structure formed in Pol I by the heterodimeric complex
Rpa43/Rpa14, which is involved in docking the essential Rrn3 initiation transcription factor to the enzyme (Peyroche et al, 2000; Yamamoto et al, 1996; Blattner et al, 2011). The Pol I-Rrn3 complex interacts with promoter bound factors, the core factor (CF), forming the initially transcribing complex (ITC) (Engel et al, 2017; Sadian et al, 2017; Han et al, 2017; Keener et al, 1998). Additionally, Pol I and Pol III contain subunits that are functionally and structurally related to Pol II-specific basal transcription factors, called the "Built-in Transcription Factors" (Kuhn et al, 2007; Geiger et al, 2010). Their presence in Pol I and Pol III results in a higher number of subunits, from 12 subunits in Pol II, to 14 and 17 for Pol I and Pol III, respectively, and correlates with substantial transcript production from a few genes (Werner, 2008). The heterodimer formed by Rpa34 and the N-terminal domain of Rpa49 (Rpa49Nt) in Pol I (i.e. Rpc53 and Rpc37 in Pol III) is related to the basal transcription factor TFIIF, and stimulates endogenous transcript cleavage activity (Geiger et al, 2010; Landrieux et al, 2006; Wu et al, 2011). Rpc34 in Pol III and the Rpa49 C-terminal domain (Rpa49Ct) bear a tandem winged helix motif similar to TFIIE (Geiger et al, 2010; Landrieux et al, 2006). Rpa49Ct binds DNA and is involved in initiation and elongation (Pilsl et al, 2016; Geiger et al, 2010). Finally, Rpa12 (Pol I) andRpc11 (Pol III) C-terminal domains are both directly involved in stimulating endogenous transcript cleavage activity, similar to that of TFIIS for Pol II (Van Mullem et al, 2002; Kuhn et al, 2007).

Yeast genetic studies of Pol III and Pol I “Built-in Transcription Factors” have revealed striking differences, despite their clear similarities. Each Pol III subunit is essential for cell growth, but none of the Pol I “Built-in Transcription Factors” is needed for cell survival. Deletion of Rpa34 or invalidation of Rpa49Nt, by removing the TFIIF-like heterodimer, has a very mild growth effect in vivo (Gadal et al, 1997; Beckouet et al, 2008; Geiger et al, 2010). In contrast, full or C-terminal deletion of Rpa49 leads to a strong growth defect at all temperatures, which is more severe below 25°C (Liljelund et al, 1992; Beckouet et al, 2008;
Geiger et al, 2010). Full deletion of Rpa12 leads to a strong growth defect at 25 and 30°C, and is lethal at higher temperatures (Nogi et al, 1993). Deletion of the C-terminal extension of Rpa12 abolishes stimulation of intrinsic cleavage, without any detectable growth defect (Van Mullem et al, 2002; Kuhn et al, 2007). Finally, yeast strains carrying the triple deletion of RPA49, RPA34 and RPA12 are viable, but accumulate the growth defects associated with each of the single mutants (Gadal et al, 1997).

Pol I is functional in the absence of Rpa49, but shows well-documented initiation and elongation defects, both in vivo and in vitro (Liljelund et al, 1992; Beckouet et al, 2008; Albert et al, 2011; Pilsl et al, 2016; Gadal et al, 2002; Huet et al, 1975). Restoration of active rRNA synthesis, in the absence of Rpa49, has been used to identify factors involved in initiation and elongation, such as Hmo1 and Spt5, or other Pol I subunits, such as Rpa43 (Gadal et al, 2002; Beckouet et al, 2008; Viktorovskaya et al, 2011). We made use of the spontaneous occurrence of extragenic suppressors of the growth defect of the rpa49 null mutant (Liljelund et al, 1992) to better understand the role of this subunit and reveal new interactions between the various subunits of RNA pol I during transcription. We showed that the suppressing phenotype was caused by specific point mutations in the two largest Pol I subunits, Rpa190 and Rpa135. We identified a small area at the interface between the jaw and lobe modules of the two largest subunits, in which the Rpa12 linker domain is inserted, by mapping the mutations onto the structure of Pol I. Rpa12 turned out to be essential for suppression. Moreover, Pol I carrying a single point mutation in Rpa12 was able to correct the growth defect and initiate transcription in the absence of Rpa49. Overall, our genetic data, gathered in the absence of Rpa49 or Rpa49Ct domain, suggest that the jaw/lobe interface, including the N-terminus of Rpa12, is important for the stabilization of Pol I that allows efficient transcription.
**Results**

**Isolation of extragenic suppressor mutants of the growth defect in absence of Rpa49**

We characterized extragenic suppressors of the \textit{RPA49} deletion to better understand how cell growth is achieved in the absence of Rpa49. \textit{RPA49} full-deletion mutants show a strong growth defect at 30°C and are unable to grow at 25°C. However, spontaneous suppressors have been previously observed (Liljelund \textit{et al}, 1992). We reproduced this observation and quantified the frequency of occurrence of individual clones able to grow at 25°C. There was a low frequency of colony occurrence, comparable with the spontaneous mutation rate of a single control gene (\textit{CAN1}; $< 5 \cdot 10^{-6}$). We isolated more suppressors by irradiating the cells with UV light. UV irradiation, resulting in a survival rate of approximately 50%, increased the frequency of suppressor mutations by approximately 10 fold. We identified clones that grew at 25°C after three days and selected individual colonies, called SGR for suppressor of growth defect of \textit{RPA49} deletion, with various growth rates. We ranked SGR from 1 to 186 based on their growth rates at 25°C; \textit{SGR1} had a growth rate comparable to the wild-type (WT) condition (Figure 1A). We crossed the 186 SGR clones with a strain of the opposite mating type bearing only the deletion of \textit{RPA49} to obtain diploid cells homozygous for the \textit{RPA49} deletion and heterozygous for each suppressor. The restoration of growth of the diploids at 25°C showed that all suppressor phenotypes obtained were fully or partially dominant. We focused on the most efficient suppressor clones, SGR1 and SGR2, and performed tetrad analysis to follow segregation of the observed suppression phenotype. Each suppressor phenotype was linked to a single locus in the genome and neither SGR mutant had a strong growth defect (SGR1 in Figure 1A). We used global genomic mapping of SGR1 and SGR2, derived from "genetic interaction mapping" (GIM) methods (Decourty \textit{et al}, 2008) (Materials and Methods; Suppl. Figure 1), and found a genomic linkage close to genes encoding the two largest Pol I subunits: \textit{RPA135} for SGR1 and \textit{RPA190} for SGR2 (Suppl.
Figure 1). Sequencing of the genomic DNA revealed that SGR1 bears a double mutation, whereas SGR2 bears a single one (RPA135-I218T/R379K and RPA190-A1557V alleles, respectively). Furthermore, we identified an additional mutant, SGR3, in RPA135, RPA135-R305L. The heterogeneity of the growth induced by strong UV mutagenesis prevented suppressor cloning from the 183 remaining SGR clones.

We next used the dominant phenotype of these suppressors to isolate more alleles which suppress the deletion phenotype of Rpa49 in RPA190 and RPA135. We constructed a library of randomly generated mutants (see Materials and Methods) by propagating plasmids bearing WT RPA135 or RPA190 in a mutagenic E.coli strain. After phenotypic selection of rpa49Δ mutants bearing a mutagenized Rpa190 or Rpa135 subunit at 25°C, each plasmid bearing a suppressor allele was extracted, sequenced, and re-transformed into yeast to confirm the suppressor phenotype. We thus isolated nine novel alleles of Rpa190 and 13 of Rpa135 that were able to restore growth of rpa49 deletion mutant at 25°C (supplementary Table 1). We evaluated the suppression strength based on growth restoration at 25°C relative to WT growth, as for the original SGR strains.Suppressor alleles more effective than SGR1, 2, or 3 were identified using mutagenesis of Rpa190 or Rpa135 (suppl. Table 1). In conclusion, we identified novel alleles of the two largest Pol I subunits as extragenic suppressors of the rpa49Δ-associated growth defect.

**Rpa190 and Rpa135 mutant alleles can bypass the need of Rpa49 for optimal growth**

The growth of the strains bearing one of six alleles (RPA190-E1274K, RPA190-C1493R, RPA190-L1262P, RPA135-R379G, RPA135-Y252H, and RPA135-F301S) was evaluated by a 10-fold dilution test (Figure 1B), showing good suppression by all in the absence of Rpa49. Previous genetic studies have isolated other genetic backgrounds that alleviate the growth defect of rpa49Δ at 25°C, such as rpa43-35,326 (Beckouet et al, 2008), decreased rDNA copy number (Albert et al, 2011), Hmo1 over-expression (Gadal et al, 2002), or Spt5
truncations (Viktorovskaya et al., 2011). For all these mutants, rRNA synthesis was only partially restored in the absence of Rpa49 and significant transcription defects remained. Here, we focused on the RPA135-F301S allele, the most effective growth suppressor of the Rpa49 deletion: the rpa49Δ RPA135-F301S double mutant grew almost as well at 25°C as the WT strain (Figure 1B). In the absence of Rpa49, Rpa34 does not associate with transcribing Pol I (Beckouet et al., 2008). The rpa49ΔCt allele does not have the TFIIE-like module, including the tandem wing helix motif (corresponding to residues 187-415). In strains bearing the rpa49ΔCt allele, the TFIIF module (Rpa34 and Rpa49Nt) remains associated with the polymerase (Liljelund et al., 1992; Beckouet et al., 2008). However, yeast bearing rpa49ΔCt or rpa49 full-deletion have a similar growth defect (Liljelund et al., 1992; Beckouet et al., 2008). We sought further insight into the effect of the suppressors by integrating the RPA135-F301S point mutation into the endogenous gene in three genetic backgrounds: WT, rpa49Δ (full deletion), or rpa49ΔCt. The growth rate was determined in each of these yeast strains at 30°C, in the presence or absence of RPA135-F301S. The suppressor allele RPA135-F301S had no effect on growth in the WT strain (doubling time of 102 min). The doubling time was 180 min for the rpa49Δ strain and RPA135-F301S suppression restored growth to a doubling time of 135 min. We observed similar suppression on the rpa49ΔCt background (data not shown).

**Pol I bearing RPA135-F301S allele restores efficient rRNA synthesis in the absence of Rpa49Ct in vivo**

We used yeast mutant cells with a low and stable number of rDNA repeats to better associate the growth phenotype of the RPA135-F301S allele to nascent rRNA synthesis and Pol I density on transcribed genes in vivo. Almost all rRNA genes in such strains are in the active state and transcribe their rRNA genes with a very high Pol I loading rate (Cioci et al., 2003; Albert et al., 2011). Accordingly, this genetic background is better suited to study variations in the number of polymerase molecules per rRNA gene, as the cells of these strains have a fixed
number of rRNA copies which are all in the open chromatin state. We generated three strains on this background (bearing rpa49ΔCt, RPA135-F301S, or rpa49ΔCt RPA135-F301S alleles) and determined the doubling time (Figure 2A) and de novo synthesis of rRNA (Figure 2B). The presence of the RPA135-F301S allele in Pol I effectively compensated the growth defect caused by the absence of Rpa49. Labeling of the nascent rRNA was performed using a 2-min pulse with 3H adenine. We performed the labeling in three independent cultures because of heterogeneity in the cell cultures of the rpa49ΔCt mutant. RNA synthesis was reduced approximately five-fold for rpa49ΔCter, even under permissive conditions (30°C) (compare Fig. 2B, lane 1 to lanes 2 to 5). Pol I activity in the presence of RPA135-F301S was similar to that of the WT enzyme in the absence of Rpa49Ct (compare Figure 2B, lanes 5 and 6). Thus, RPA135-F301S appeared to restore rRNA production in the absence of Rpa49.

Rpa49 is involved in initiation and elongation. We evaluated Pol I density on transcribed genes by performing Miller spreads, the only technique that currently allows the counting of individual Pol I molecules on a single rRNA gene (Miller & Beatty, 1969; Albert et al., 2011). We previously showed that full deletion of rpa49 resulted in a three-fold decrease of Pol I density per gene in the same genetic background (Miller & Beatty, 1969; Albert et al., 2011). Cells of the strain expressing the rpa49ΔCt allele showed a four-fold decrease of Pol I density per gene (Figure 2C). Expression of the RPA135-F301S allele in the presence of Rpa49 or Rpa49Ct had no influence on Pol I density (Fig. 2C, RPA135-F301S). In contrast to strain rpa49ΔCt, the double mutant rpa49ΔCt RPA135-F301S showed significantly higher pol I occupancy (46 instead of 21 Pol I molecules per gene, on average). Indeed, the double mutant restored full rRNA production, although it exhibited only half of the WT Pol I density per gene.
Overall, these results show that the presence of the \textit{RPA135-F301S} allele in a strain lacking Rpa49Ct restores rRNA synthesis to WT levels and increases Pol I density on rRNA genes, indicative of productive transcription initiation and rRNA synthesis.

\textbf{Most suppressors are clustered in a hotspot of Pol I}

Structural data of Pol I are now available, showing Pol I in an inactive form (Fernández-Tornero \textit{et al}, 2013; Engel \textit{et al}, 2013), complexes of Pol I and Rrn3 (Pilsl \textit{et al}, 2016; Engel \textit{et al}, 2016; Torreira \textit{et al}, 2017), complexes of Pol I with other initiation factors (Engel \textit{et al}, 2017; Han \textit{et al}, 2017; Sadian \textit{et al}, 2017), and in elongating forms (Neyer \textit{et al}, 2016; Tafur \textit{et al}, 2016), respectively. We mapped Rpa135 and Rpa190 residues that suppressed the growth defect of the mutant strain \textit{rpa49ΔCt} onto the structure of WT Pol I in which the structure of Rpa49 was determined (Han \textit{et al}, 2017) (Figure 3A). Most of the suppressor mutations which provided growth recovery (see supplementary table 1) appeared to be clustered at a specific interface between the two largest subunits, Rpa190 and Rpa135 (figure 3B), between the lobe (Rpa135 - salmon) and the jaw (Rpa190 - blue). This small region is characterized by the presence of a beta-strand in the structure of Rpa12 (Rpa12 – yellow: residues 46-51, Figure 3B), following four beta-strands of Rpa190, thus forming a shared five-strand anti-parallel beta-sheet. The Rpa12 beta-strand also faces the Rpa135 lobe domain (residue 252 to 315 of Rpa135), in which six independent mutations were found, including \textit{RPA135-F301S}. Notably, Rpa135 contains a pattern of three amino acids "DSF" (D299, S300, F301), which are highly conserved among eukaryotic species. Notably, we found suppressor mutations for each of these three amino acids (see supplementary table 1). The side chains of the mutated residues are apparently involved in electrostatic interactions that stabilize this jaw/lobe interface (Figure 3C). Substituted residues resulted in destabilization of this interface (see supplementary table 1), suggesting a specific rearrangement of the interface lobe/jaw in each mutant.
Point mutations in Rpa12 suppress the growth defect when Rpa49 is missing

We then tested whether mutated alleles of RPA12 itself could behave as suppressors by generating a library of randomly mutagenized RPA12 (see Materials and Methods). Two dominant alleles (RPA12-S6L and RPA12-T49A) efficiently suppressed the growth defect of rpa49Δ and of rpa49ΔCt (Figure 4A, not shown here for rpa49ΔCt). However, these RPA12 suppressor alleles did not fully restore WT growth when expressed in the rpa49Δ mutant (compare the size of the colonies in Figure 4A.). We integrated the RPA12-S6L point mutation into the genome at its native locus in a low rDNA-copy-number genetic background, with or without rpa49Ct deletion, and analyzed the restoration of rRNA synthesis by pulse labeling (Figure 4B) and Pol I loading by Miller spreading (Figure 4C). rRNA synthesis and Pol I loading were largely restored when RPA12-S6L was expressed in strain rpa49ΔCt, similar to that for the RPA135-F301S mutant. The 3D structure of Pol I shows that RPA12-S6L and RPA12-T49A obtained by random mutagenesis are specifically located in the "hotspot" at the jaw/lobe interface. Threonine 49 of Rpa12 is located on the beta-strand (Rpa12 aa 46-51) (Figure 3B), facing residues D299, S300, and F301 of Rpa135, and Rpa190-E1274 (Figure 3B). This residue also appears to be highly conserved among eukaryotic species. The second mutation, RPA12-S6L, is located in the N-terminal domain of Rpa12. In conclusion, all point mutations in Rpa190, Rpa135, and Rpa12 detected in the hotspot domain of the jaw/lobe interface can substitute the requirement for Rpa49Ct in vivo.

Rpa14, Rpa34, and the expander/DNA mimicking loop of Rpa190, are not involved in the suppression phenotype

The structural determination of Pol I revealed the presence of an extended loop inside the DNA-binding cleft folded in a "expander/DNA mimicking loop" conformation when Pol I is in an inactive, dimeric form (Engel et al, 2013; Fernández-Tornero et al, 2013). This element is not found in Pol II or III. These residues are inserted between the four beta-strands of
Rpa190, corresponding to the mutation hotspot. A small deletion of this Rpa190 domain (1361-1390) results in a slight slow-growth phenotype (Fernández-Tornero et al, 2013). We determined whether the expander/DNA mimicking loop was responsible for the suppression phenotype. We generated a novel allele, rpa190Δloop (deletion of residues 1342-1411), which had no associated growth defect (Figure 5A). We were unable to generate a viable double mutant when combining this mutation with the rpa49 full deletion. Thus, the DNA-mimicking loop is required for Pol I activity in the absence of Rpa49. We next tested whether deletion of this loop influences suppression by the RPA135-F301S allele. Note that the rpa190Δloop combined with RPA135-F301S has no phenotype. There was no difference in the growth of the rpa49Δ RPA135-F301S double mutant and that of the triple mutant rpa49Δ RPA135-F301S rpa190Δloop (Figure 5A). Thus, the expander/DNA mimicking loop of Rpa190 is not required for suppression, but is required for the viability of the rpa49 deletion mutant.

Rpa34 forming a heterodimer with Rpa49Nt, and Rpa14 being essential in absence of Rpa49, we also introduced RPA135-F301S into yeast strains lacking theses Pol I subunits (Figure 5B and C). The growth of RPA135-F301S/rpa14Δ and RPA135-F301S/rpa34Δ double mutants was not different from that of the single mutants. However, RPA135-F301S suppressed the growth defect of the viable double mutant, rpa34Δ rpa49Δ, lacking both Rpa49 and Rpa34, the heterodimer partner of Rpa49 (Figure 5B). The double deletion mutant lacking both Rpa49 and Rpa14 was not viable, similar to the rpa190Δloop rpa49Δ double mutant (Gadal et al, 1997). Introduction of the suppressor RPA135-F301S, by genetic crossing, resulted in a triple mutant (rpa14Δ rpa49Δ RPA135-F301S) that could grow, but much more slowly (Figure 5C).
Overall, $RPA135-F301S$ had no effect when combined with the single deletions $rpa190\Delta$loop, $rpa34\Delta$, or $rpa14\Delta$. However, $RPA135-F301S$ suppression of $rpa49\Delta$-associated growth defect does not require Rpa190 DNA mimicking loop, Rpa34, or Rpa14.

**Rpa12 is required for suppression of the rpa49\Delta-associated growth defect**

We next evaluated whether specific regions of Rpa12 may be involved in the suppression of the $rpa49\Delta$-associated growth defect (Figure 6A). The C-terminal region of Rpa12 (TFIIS-like) is inserted towards the active center of Pol I to stimulate intrinsic cleavage activity but is displaced during productive initiation and elongation steps. C-terminal deletion of Rpa12 resulted in normal growth (Van Mullem et al, 2002) (Figure 6A- lane 2), although the $Rpa12\Delta Ct$ allele is unable to stimulate cleavage activity in vitro (Kuhn et al, 2007). Full deletion of $RPA12$ resulted in a slight growth defect at 24°C, which was stronger at 30°C (Nogi et al, 1993) (Figure 6A-lane 3). In contrast, $rpa49$ deletion resulted in a growth defect at 30°C, which was stronger at 24°C (Liljelund et al, 1992) (Figure 6A-lane 4). Combining $rpa12\Delta Ct$ with $rpa49\Delta$ resulted in a mildly synergistic phenotype, with a stronger growth defect at both 24°C and 30°C (Figure 6A – lane 5). The double mutant lacking both full Rpa12 and Rpa49 subunits was viable, but had a major growth defect (Gadal et al, 1997) (Figure 6A – lane 6).

Random mutagenesis of $RPA12$ did not lead to the isolation of $RPA12$ alleles containing suppressor mutations in the C-terminal domain. We explored whether the C-terminal extension of Rpa12 is necessary for suppression of the $rpa49\Delta$ phenotype by introducing the Rpa12 C-terminal truncation into the strain bearing both $rpa49\Delta$ and suppressor alleles $RPA12-S6L$ or $RPA12-T49A$. Thus, $RPA12-S6L$ and $RPA12-S6L-\Delta Ct$ resulted in similar suppression of the $rpa49\Delta$ growth defect (Figure 6B). $RPA12-T49A$ behaved similarly to $RPA12-S6L$ (data not shown).
We then introduced the RPA135-F301S allele and assessed the suppression phenotype of the strain lacking Rpa49 at 25°C, with or without the entire Rpa12 subunit (Figure 6C). We constructed a strain with RPA12 under the control of the regulatable pGAL promoter. The growth defect of rpa49Δ was completely suppressed by the RPA135-F301S allele when RPA12 was expressed (Figure 6C, left panel), whereas suppression mediated by RPA135-F301S was completely abolished in its absence (Figure 6C, middle panel). Residual growth of the rpa49 rpa12 double mutant was detectable after 10 days (Figure 6B, right panel) and was the same with or without the RPA135-F301S allele.

In conclusion, we show that Rpa12, particularly its N-terminal portion, is required for the suppression of the growth defect mediated by RPA135-F301S when Rpa49 is depleted.

**Rpa12 is required for efficient promoter-dependent transcriptional activity in vitro**

The suppressor mutants suggest a functional interaction between Rpa49 and Rpa12. The absence of Rpa49 or its C-terminal extension strongly reduced Pol I loading on rDNA genes (Albert et al., 2011; Beckouet et al., 2008; Figure 2C), indicating an initiation defect in vivo. The in vitro results of promoter-dependent transcription assays in the absence of Rpa49, particularly of Rpa49Ct, are consistent with the in vivo data, in which their absence results in an almost initiation-inactive form of the polymerase, whereas nonspecific RNA synthesis is not as strongly affected (Pilsl et al., 2016). Miller spread experiments have been previously performed for mutant strains lacking Rpa12 (Prescott et al., 2004), but Pol I occupancy of the rDNA genes was not analyzed, and is difficult to interpret due to the known instability of Rpa190 in the absence of Rpa12 (Van Mullem et al., 2002). However, it is possible to test whether subunit Rpa12 supports transcription initiation in vitro. We thus performed promoter-dependent transcription and non-specific transcription assays comparing Pol I, lacking Rpa12, to WT Pol I (Figure 7).
In vitro, transcription on tailed templates was significantly reduced in the absence of Rpa12 when the amount of Pol I was held constant (approximately 14% using 4 nM Pol relative to that of WT Pol I). In promoter-dependent transcription the reduction in transcription efficiency was much more pronounced (approximately 1.2% of that of WT Pol I) indicating that Rpa12 is involved in the initiation step.
Discussion

Here, we investigated how the growth of cells can improve in the absence of Rpa49. We showed that altering a very specific area of Pol I resulted in a functional Pol I molecule lacking this subunit. We propose that the mutated area, which includes a part of subunit Rpa12, undergoes a conformational change that supports the initiation of transcription.

Suppressor mutants are not at the Rrn3-Pol I stalk interface

Our previous studies suggested the specific involvement of Rpa49 in the association and dissociation of Rrn3 from the Pol I stalk (Albert et al., 2011; Beckouet et al., 2008). Here, we show that genetically modified polymerases lacking Rpa49 or Rpa49Ct, with a single modified residue in Rpa190, Rpa135, or Rpa12, at a position diametrically opposed to the position that binds to Rrn3, can initiate transcription and that strains harboring them grow normally. We propose that there is a second important interface which is involved in Pol I recruitment to the rDNA gene, in addition to the interface between Rrn3 and the stalk.

The role of Rpa12 in suppression

The Rpa12 subunit is involved in stimulating the intrinsic cleavage activity of Pol I through a TFIIS-like domain at its C-terminus. Purified Pol I with Rpa12 lacking the C-terminal domain has no cleavage activity (Kuhn et al., 2007). Furthermore, the C-terminal domain of Rpa12 can contact the active site of the polymerase in the inactive conformation and is retrieved in both initiation competent and elongating forms of the polymerase. Direct evidence that cleavage is not involved in suppression of the growth defect came from the experiments showing a fully functional suppressor phenotype for *RPA12ΔCt-S6L*, which lacks the domain required for stimulating cleavage.

The N-terminal domain of Rpa12, at the surface of Pol I, is involved in the recruitment of the largest subunit, Rpa190, at 37°C (Van Mullem et al., 2002) and is required for docking of this
subunit to the enzyme. A linker region of Rpa12 connects the N-terminal module (Rpb9-like) at the surface of Pol I to the mobile C-terminal region (TFIIS-like) and is therefore indirectly required for cleavage. *In vitro*, purified Pol I that lacks Rpa12 has less activity than WT Pol I in promoter-dependent transcription assays. Mutations in other Pol I domains, such as deletions in Rpa34, Rpa14, or the Rpa190-DNA mimicking loop, did not influence suppression of the rpa49 deletion growth defect by the RPA135-F301S allele. In contrast, the Rpa12 linker was absolutely required for efficient suppression. Accordingly, the RPA135-F301S allele was unable to restore efficient growth when Rpa12 was absent. Thus, Rpa12 and RPA135-F301S cooperate to support initiation if Rpa49 is absent.

**Modification of the jaw/lobe interface facilitates DNA cleft closure**

RNA polymerase I undergoes major conformational changes during the transcription cycle, mainly affecting the width of the DNA-binding cleft(Fernández-Tornero, 2018). During the initiation of transcription, the cleft aperture narrows from a semi-open configuration, as seen in cryo-EM structures of the enzyme bound to Rrn3 (Engel et al, 2016; Pilsl et al, 2016; Torreira et al, 2017), to a fully closed conformation observed in transcribing complexes (Engel et al, 2017; Han et al, 2017). This allows gripping of the transcription bubble inside the cleft (Figure 8A). Following Rrn3 release, DNA binding is further secured by the Rpa49-linker, which crosses the cleft from the lobe to the clamp, passing over the downstream DNA, and the Rpa49Ct domain, which binds the upstream DNA in the vicinity of the clamp (Han et al, 2017; Tafur et al, 2016).

Cleft closure is achieved by the relative movement of two structural units, located on opposite sides of the cleft, which pivot using five hinges(Fernández-Tornero et al, 2013). The unit consisting of the shelf and clamp modules is apparently rigid, whereas that consisting of the core and lobe modules, which is in the vicinity of the mutated residues of our study, undergoes internal rearrangements (Movie 1). The most prominent reorganization within this
latter unit affects the Rpa190 jaw domain, the outer rim of which shifts away from the DNA by approximately 3.7 Å, using the lobe/jaw interface as a hinge. This movement also involves the linker region of subunit Rpa12, which contains a beta-strand (residues 46-50) that completes a four-stranded beta-sheet in the Rpa190 jaw domain. As a result, a short alpha-helix within the Rpa12 linker region shifts its position by approximately 3.0 Å.

**Possible roles of Rpa12 in regulating DNA cleft closure**

Rearrangements in the jaw of Rpa190 and Rpa12 linker regions are likely essential to allow pivoting of the shelf-clamp unit against the core-lobe unit. Without such motion, cleft closure would be impossible (Movie 1). We propose that RPA135-F301S or RPA12-S6L favor DNA capture by increasing the flexibility of the lobe/jaw/Rpa12 interface of Pol I relative to that of the WT polymerase. Structural analysis suggests that the Rpa49Ct and its linker domains are involved in securing the closed cleft conformation (Tafur et al, 2016; Han et al, 2017). Cleft closure is likely destabilized in the rpa49ΔCt mutant (Figure 8B). We propose that the Rpa12 linker domain contributes to the hinge in the lobe/jaw interface. Rpa12 may be involved in cleft-closure, which is necessary to guide the DNA towards the active center, or it may stabilize DNA bound Pol I. Our mutated Pol I probably captures DNA more efficiently than WT polymerase. This structural rearrangement is strongly suggested by our genetic data and the requirement of Rpa12 for promoter-dependent in vitro initiation. There is indirect evidence that similar domains are involved in the initiation of transcription of Pol II and Pol III. Rpb9 in Pol II is similar to the Rpa12-N-terminal module, but lacks the TFIIS domain. Rpb9 is required for proper start site selection and transcription fidelity (Hull et al, 1995; Walmacq et al, 2009). Furthermore, mutations in the lobe domain of Rpb2, adjacent to Rpb9, alter both Pol II–TFIIF binding and the transcription start site (Chen et al, 2007). For Pol III, TFIIS has been shown to stimulate transcription initiation in vitro and in vivo (Ghavi-Helm et
al, 2008). However, motion at the lobe/jaw interface during cleft closure is only detectable in Pol I.
Figure legends

**Figure 1.** Alleles of *RPA190* and *RPA135* suppress the growth defect of the *rpa49Δ* mutant at various levels. (A) The SGR1 mutant restores growth of the *rpa49Δ* mutant. Ten-fold serial dilutions of wild-type (WT), *rpa49Δ* single mutant, SGR1 single mutant, and SGR1/*rpa49Δ* double mutant strains were spotted on rich media to assess growth at 30 and 25°C for three days. (B) Ten-fold dilutions of WT and *rpa49Δ* compared to *rpa49Δ* carrying various plasmids: pGL190_3 (*RPA190*-E1274K), pGL190_11 (*RPA190*-C1493R), pGL190_23 (*RPA190*-L1262P), pGL135_6prim (*RPA135*-R379G), pGL135_54 (*RPA135*-Y252H), or pGL135_33 (*RPA135*-F301S). Growth was evaluated after three days at 25°C. The strains and plasmids used are listed in supplementary Tables 2 and 3, respectively.

**Figure 2.** The *RPA135*-F301S allele restores growth and rRNA synthesis, and modulates Pol I occupancy of rDNA genes in the absence of Rpa49Ct. (A) Doubling times of WT, *rpa49ΔCt*, *RPA135*-F301S, and the *rpa49ΔCt/RPA135*-F301S double mutant in a low copy number background (see supplementary table 2). (B) *In vivo* labeling of newly synthesized RNAs. WT (lane 1), *rpa49ΔCt* (lanes 2-4), *RPA135*-F301S (Lane 5), and the *rpa49ΔCt/RPA135*-F301S double mutant (Lane 6) were grown to an OD$_{600}$ of 0.8. Cells were then pulse-labeled with [8-$^3$H] adenine for 2 min. Samples were collected, and total RNA extracted and separated by gel electrophoresis. (C) Representative Miller spreads of WT, *rpa49ΔCt*, *RPA135*-F301S, and the *rpa49ΔCt/RPA135*-F301S double mutant. Panels on the right of each micrograph show interpretive tracing of the genes. Polymerases that appear on the gene are shown on the tracing by black dots. The number of polymerases counted on the genes is indicated below. N represents the number of individual spread genes used for quantification (see Materials and Methods).
Figure 3. Mapping of the modified residues in Rpa190, Rpa135, and Rpa12 alleles on the structure of Pol I. (A) Two different views of the original initial transcribing complex model and its 14 different subunits (PDB 5W66(Han et al, 2017)). (B) Most mutated suppressor residues are clustered at the interface between the jaw (Rpa190, blue) and lobe (Rpa135, salmon) modules of Pol I. Note that Rpa12 (residue 46 to 51; yellow) is part of this interface. (C) Enlargement of the areas containing the modified residues (Rpa190-N863, -S1259, -L1262, -E1274, -C1493, Rpa135-Y252, D299, S300, F301, R305, and Rpa12-S6, T49) shown in panel B. The figure was prepared with Pymol using the crystal structure of Pol I PDB 4C3I (Fernández-Tornero et al, 2013)).

Figure 4. The RPA12-S6L allele restores growth and rRNA synthesis and modulates Pol I occupancy of rDNA genes in the absence of Rpa49Ct. (A) Ten-fold dilutions of the rpa49Δ mutant carrying various plasmids: an empty pRS316 plasmid (rpa49Δ), YCp50-26 bearing RPA49 (WT), pRS316-A12-S6L (RPA12-S6L), or pRS316-A12-T49A (RPA12-T49A). Growth was evaluated after three days at 25°C or two days at 30°C. (B) In vivo labeling of newly synthesized RNA. For comparison, WT (lane 1) and rpa49ΔCt (lanes 2) depicted in figure 2B are shown with the RPA12-S6L single mutant and the rpa49ΔCt/RPA12-S6L double mutant. (C) Representative Miller spreads of WT, rpa49ΔCt, RPA12-S6L, and the rpa49ΔCt/RPA12-S6L double mutant.

Figure 5. Rpa14, Rpa34, and the DNA mimicking loop of Rpa190 are not required for suppression. Deletion of the DNA mimicking loop of Rpa190 (A) or RPA34 (B) does not modulate the suppression activity of RPA135-F301S. (C) RPA135-F301S suppresses the synthetic lethality between rpa14Δ and rpa49Δ. Ten-fold serial dilutions were performed seeded onto rich media and growth evaluated after three days at 24°C.
Figure 6. **RPA12 alleles can modulate the rpa49Δ-associated growth defect.** (A) Growth of the double mutants: rpa49Δ rpa12ΔCt, or rpa49Δ combined with full depletion of rpa12. Depletion of Rpa12 was achieved using a pGAL-RPA12 construct on glucose containing medium (strain OGT30-1c). Ten-fold serial dilutions of OGT30-1c bearing pRS316-A12 (WT), pRS316-A12-DCter expressing Rpa12 bearing a C-terminal deletion of residues 65-125 (rpa12ΔCt), or an empty plasmid pRS316 (-rpa12) were seeded onto media. The growth of rpa49Δ combined with RPA12 depletion was tested using strain OGT30-3c bearing pRS316-A12 (rpa49Δ), pRS316-A12-DCter (rpa12ΔCt rpa49Δ), or an empty plasmid pRS316 (-rpa12 rpa49Δ). Growth was assessed after four days at 24 or 30°C. (B) The C terminus of Rpa12 is not required for suppression. Ten-fold serial dilutions of OGT30-1c (RPA49-WT), bearing pRS316-A12 (WT) or pRS316-A12-DCter (rpa12ΔCt), and OGT30-3c (rpa49Δ), bearing pRS316-A12 (WT), pRS316-A12-DCter (rpa12ΔCt), pRS316-A12-S6L (RPA12-S6L), or pTD10 (RPA12-S6L-ΔC) were seeded onto media. Growth was assessed after four days at 24°C. (C) Suppression activity of RPA135-F301S is abolished in the absence of Rpa12. RPA12, under a regulatable promoter (pGAL) was either expressed (+RPA12; left panel) on galactose containing medium or repressed (-RPA12; right panel) on glucose containing medium. Ten-fold serial dilutions of RPA49-WT, rpa49Δ, RPA49-WT, RPA135-F301S, or rpa49Δ RPA135-F301S were grown at 24°C. Depletion of RPA12 abolishes the suppression activity of RPA135-F301S (compare the left to the middle and right panels). Extended incubation (right panel, 10 days) was used to detect growth of the double mutant -RPA12 rpa49Δ on plates.

Figure 7. **Subunit Rpa12 supports in vitro Pol I transcription initiation and non-specific transcription.** Tailed template assays (lane 1-3) and promoter-specific transcription (lane 4-6) were performed using 4 nM WT Pol I or 4 and 12 nM Pol I lacking Rpa12 (see Materials and Methods). Quantification relative to WT activity is shown for each lane.
Figure 8. Schematic representation of Pol I (A) Free monomeric Pol I with mobile Rpa49Ct and linker. (B) Initially transcribing complex (ITC) upon insertion of melted DNA in the presence of Rpa49 (purple). Rpa49Ct interacts with upstream DNA and the Rpa49-linker is folded, closing the cleft. Movements of Rpa12 and the jaw with respect to the lobe are indicated with arrows. (C) Pol I lacking Rpa49 is likely defective in stabilizing the closed conformation in the DNA-binding cleft, likely resulting in a looser grip on the DNA (red asterisks). (D) Suppressor mutations (green) facilitate movement of the jaw/lobe interface and gripping of the DNA by the Pol I enzyme, in the absence of Rpa49.
**MATERIALS & METHODS**

*Plasmids and yeast strain constructions*

The oligonucleotides used in this study are listed in supplementary Table 4. Plasmids and details of the cloning steps are described in supplementary Table 3. Randomly mutagenized *RPA190* and *RPA135* libraries were obtained by transformation and amplification of pVV190 and pNOY80, respectively, into XL1-red strains according to the manufacturer’s guidelines (XL1-Red Competent Cells, from Agilent Technologies). Yeast strains are listed in supplementary Table 2, and were constructed by meiotic crossing and DNA transformation (Schiestl & Gietz, 1989)(Sambrook *et al*, 1989). The yeast media and genetic techniques were described previously (Sherman *et al*, 1986).

yCNOD226-1a was obtained from yCNOD223-2a by switching the *KAN-MX* to the *NAT-MX* marker under the control of the MF(ALPHA)2/YGL089C promoter (alphaNAT-MX4), which allowed selection of MATα haploid cells (Decourty *et al*, 2008). OGT9-6a is an offspring of yCNOD226-1a crossed with BY4741. OGT8-11a is an offspring of BY4742 crossed with Y1196. Strains OGT9-6a and OGT8-11a were plated on rich media and UV irradiated (5W/m² during 5 second), resulting in 50% survival. LH514D and LH11D are suppressor clones of the growth defect selected from UV-irradiated OGT9-6a grown at 25°C. AH29R is a suppressor clone of the growth defect selected from UV-irradiated OGT8-11a grown at 25°C. Genetic interaction mapping (GIM) analysis of the *RPA49* deletion mutant was performed as described previously (Decourty *et al*, 2008). Microarray data were normalized using MATLAB (MathWorks, Inc., Natick, MA) as previously described (Albert *et al*, 2011). OGT15-7b is an offspring of LH514D with BY4741, followed by homologous recombination using PCR-amplified fragments generated with oligos 1716 and 1717 and pCR4-HIS3 as template. yTD16-1a was first transformed with plasmid pCJPF4-GAL49-1. Then, *RPA135* was tagged by homologous recombination using PCR-amplified fragments generated with
oligos 835 and 836 and genomic DNA of strain RPA135-TAP or yTD6-6c, generating yTD27-1 and yTD28-1a, respectively. Strain yTD25-1a bears a C-terminal deletion of RPA49 generated by homologous recombination using PCR-amplified fragments generated with oligos 208 and 1515 and pFA6-KAN-MX6 as template. yTD11-1a was derived from strain yTD25-1a after switching to HPH-MX by homologous recombination using pUC19-HPH cut by BamHI. C-terminal deletion of RPA49 in yTD29-1a and yTD30-1a were generated by homologous recombination using PCR-amplified fragments generated with oligos 649 and 650 and yTD11-1 genomic DNA as template, transformed into yTD27-1 and yTD28-1a, respectively. Genomic allelic insertion of RPA12-S6L in yTD31-1a and yTD23-1a was performed by homologous recombination using PCR-amplified fragments generated with oligos 1556 and 1557 and pRS316-A12-S6L-KAN as template, transformed into yTD27-1 and yTD29-1a, respectively.

TGT135-3b was obtained after sporulation of y27138 transformed by pNOY80. TGT135-3b and OGT15-9d were mated to generate TGT12. yTD2-3b and yTD2-3d are offspring of TGT12 transformed with pGL135_33. yTD6-6c and yTD6-6b were generated by homologous recombination using pTD2_6c_135TAP cut with XhoI-NsiI, transformed into yTD2-3b and yTD2-3d, respectively. yTD48-1a was generated by deletion of the Rpa190 DNA-mimicking loop using homologous recombination with PCR-amplified fragments generated using oligos 1189 and 1194 and genomic DNA of strain SCOC2260 as template, transformed into BY4741. yTD51-2c, yTD51-8a, and yTD51-5a are offspring of yTD48-1a mated with yTD37-3a. yTD36-2b is an offspring of yCN224-1a mated with yTD40-1a. yTD37-3d and yTD37-7d are offspring of yCN224-1a mated with yTD41-1a. yTD38-3d is an offspring of yCN225-1a mated with yTD40-1a and yTD39-8a is an offspring of yCN225-1a mated with yTD41-1a.
Strain yTD53-1a was constructed by homologous recombination using a PCR-amplified fragment generated with oligos 1634 and 1635 and pFA6a-KanMX6-GAL::3HA as template. OGT30-1a and OGT30-3a are offspring of yTD53-1a mated with yTD6-6b. yTD40-1a and yTD41-1a were generated by homologous recombination using PCR-amplified fragments generated with oligos 700 and 1679 and pFA6a-HA-KIURA3 as template, transformed into OGT30-3a and OGT30-1a respectively, switching RPA135-TAP-tag to untagged RPA135.

**In vivo labeling and RNA extraction and analysis**

Metabolic labeling of pre-rRNA was performed as previously described (Hermann-Le Denmat et al., 1994) with the following modifications. Strains were pre-grown in synthetic glucose-containing medium lacking adenine at 30°C to an OD600 of 0.8 at. One milliliter cultures were labeled with 50 µCi [8-3H] adenine (NET06300 PerkinElmer) for 2 min. Cells were collected by centrifugation and the pellets frozen in liquid nitrogen. RNA was then extracted as previously described (Beltrame & Tollervey, 1992) and precipitated with ethanol. For high molecular weight RNA analysis, 20% of the RNA was glyoxal denatured and resolved on a 1.2% agarose gel. Low molecular weight RNAs were resolved on 8% polyacrylamide/8.3 M urea gels.

**Miller spread experiments and analysis**

Chromatin spreading was mainly performed as described previously with minor modifications (Osheim et al., 2009). Carbon-coated grids were rendered hydrophilic by glow discharge instead of ethanol treatment. Negatively stained chromatin was obtained by short incubation with heavy metal followed by quick drying of the sample. Images were obtained using a JEOL JEM-1400 HC electron microscope (40 to 120 kV) with an Orius camera (11Mpixels). The position of the RNA polymerase I molecules and the rDNA fiber were determined by
visual inspection of micrographs using Image J (http://rsb.info.nih.gov/ij/). Digital images were processed by software programs Image J and Adobe Photoshop® (v. CS6).

**In vitro promoter-dependent transcription assays**

*In vitro* promoter-dependent transcription reactions were performed as previously described (Pilsl *et al*., 2016; Tschochner, 1996) with some modifications. Briefly, 1.5 ml reaction tubes (Sarstedt safety seal) were placed on ice. Template (0.5 –1 µl; 50–100 ng DNA) was added, corresponding to a final concentration of 5–10 nM per transcription reaction (25-µl reaction volume). Core factor (1–2 µl; 0.5 to 1 pmol/µl; final concentration 20–40 nM) and 1–3 µl Pol I (final concentration 4–12 nM) were added to each tube. Then, 20 mM HEPES/KOH pH 7.8 was added to a final volume of 12.5 µl. Transcription was started by adding 12.5 µl 2X transcription buffer. The samples were incubated at 24°C for 30 min at 400 rpm in a thermomixer. Transcription was stopped by adding 200 µl Proteinase K buffer (0.5 mg/ml Proteinase K in 0.3 M NaCl, 10 mM Tris/HCl pH 7.5, 5 mM EDTA, and 0.6% SDS) to the supernatant. The samples were incubated at 30°C for 15 min at 400 rpm in a thermomixer. Ethanol (700 µl) p.a. was added and the tubes mixed. Nucleic acids were precipitated at -20°C overnight or for 30 min at -80°C. The samples were centrifuged for 10 min at 12,000g and the supernatant removed. The precipitate was washed with 0.15 ml 70% ethanol. After centrifugation, the supernatant was removed and the pellets dried at 95°C for 2 min. RNA in the pellet was dissolved in 12 µl 80% formamide, 0.1 M TRIS-Borate-EDTA (TBE), 0.02% bromophenol blue, and 0.02% xylene cyanol. Samples were heated for 2 min with vigorous shaking at 95°C and briefly centrifuged. After loading on a 6% polyacrylamide gel containing 7M urea and 1X TBE, RNAs were separated by applying 25 watts for 30–40 min. The gel was rinsed in water for 10 min and dried for 30 min at 80°C using a vacuum dryer. Radiolabelled transcripts were visualised using a PhosphoImager.
Suppl movie legends 1

Supplemental Movie 1. Conformational changes in Pol I upon initiation. The movie starts with the closed cleft conformation (PDB 5W66 (Han et al, 2017)), in which melted DNA occupies the cleft and then changes to the intermediate cleft conformation observed in monomeric Pol I (PDB 5M3M(Neyer et al, 2016)). Relevant structural regions have been variously colored and labeled, and residues mutated in this report are shown in red.
**Supplementary tables**

| Sub. | Allele | Strength | Hotspot | Location | Mutant effect |
|------|--------|----------|---------|----------|---------------|
| Rpa190 | N863T | Weak | Funnel | Funnel/ Rpa12-linker | Destabilization of Funnel/Rpa12 |
| Rpa190 | S1259L | Medium | Jaw | Jaw/ Shelf hinge | Hinge conformation |
| Rpa190 | L1262P | Weak | Jaw | Jaw/ Shelf hinge | Hinge conformation |
| Rpa190 | E1274K | Medium | Jaw | Jaw/ Rpa12-linker interface | Destabilization of Jaw/Rpa12 |
| Rpa190 | C1493R | Medium | Jaw | Jaw/ Rpa12-linker interface | Destabilization of Jaw/Rpa12 |
| Rpa135 | Y252H | Strong | Lobe | Jaw/ Lobe interface | Destabilization of Jaw/Lobe |
| Rpa135 | D299G | Medium | Lobe | Jaw/ Lobe interface | Destabilization of Jaw/Lobe |
| Rpa135 | S300F | Medium | Lobe | Jaw/ Lobe interface | Destabilization of Jaw/Lobe |
| Rpa135 | F301S | Strong | Lobe | Jaw/ Lobe interface | Destabilization of Jaw/Lobe |
| Rpa135 | F301L | Strong | Lobe | Jaw/ Lobe interface | Destabilization of Jaw/Lobe |
| Rpa135 | SGR3 R305L | Medium | Lobe | Jaw/ Lobe interface | Destabilization of Jaw/Lobe |
| Rpa12 | S6L | Medium | N-terminal | Jaw/ Rpa12-linker interface | Destabilization of Jaw/Rpa12 |
| Rpa12 | T49A | Medium | Linker | Jaw/ Rpa12-linker interface | Destabilization of Jaw/Rpa12 |
| Rpa190 | L608S | Medium | | | |
| Rpa190 | E611K | Medium | | | |
| Rpa190 | S936A | Weak | | | |
| Rpa190 | SGR2 A1557V | Weak | | | |
| Rpa135 | D157G | Medium | | | |
| Rpa135 | D157N | Medium | | | |
| Rpa135 | SGR1 I218/ R379K | Medium | | | |
| Rpa135 | R379G | Weak | | | |
| Rpa135 | G580D | Medium | | | |
| Rpa135 | C584Y | Weak | | | |
| Rpa135 | I913V | Medium | | | |

**Supplementary table 1.** List of 24 individual suppressor mutations of the growth defect of *rpa49Δ* strain in the Rpa190, Rpa135, and Rpa12 subunits (Sub.). The suppressors were classified according to growth rate when combined with *rpa49Δ* mutant: weak, medium, or strong. SGR1, 2, and 3 depict alleles originally isolated after UV mutagenesis (see text). Thirteen of the 24 mutants, affecting 12 different positions, were found in a specific hot-spot shown in Figure 4.
## Supplementary table 2: Yeast strains used in this study

| Referred to as | strain | Genotype | source |
|----------------|--------|----------|--------|
| WT in figures 1 and 5 | BY4741 | MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 | Euroscarf |
| yT1196 | MATα his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0 rpa49Δ::KANMX4 | Euroscarf |
| yT2738 | MATα alpha his3Δ1/ his3Δ1 leu2Δ0/ leu2Δ0 ura3Δ0/ ura3Δ0 lys2Δ0/ LYS2 MET15/ met15Δ0 rpa135Δ::KANMX4/RPA135 rpa49Δ::HPHMX4/RPA49 | Euroscarf |
| TGT135-3b | MATα his3Δ11 leu2Δ0 ura3Δ0 lys2Δ0 rpa135Δ::KANMX4 +pNOY80 | Euroscarf |
| TGT12 | MATα alpha his3Δ1/ his3Δ1 leu2Δ0/ leu2Δ0 ura3Δ0/ ura3Δ0 lys2Δ0/ lys2Δ0 rpa135Δ::KANMX4/RPA135 rpa49Δ::HPHMX4/RPA49 | Euroscarf |
| rpa49Δ in figure 1, with the plasmids indicated in the legend | OGT9-6a | MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 rpa49Δ::KANMX6 | This study |
| rpa49Δ in figure 4A, with the plasmids indicated in the legend | OGT8-11a | MATα his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 rpa49Δ::KANMX6 | This study |
| SGR1 in figure 1 | OGT15-7b | MATα his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 HIS3::RPA135(I218T, R379K) | This study |
| SGR1 rpa49Δ in figure 1 | LHS14D | MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 rpa49Δ::alphaNAT | This study |
| SGR2 | AH29R | MATα his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 rpa49Δ::kanmx4 RPA190-1557V | This study |
| SGR3 | LH11D | MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 rpa49Δ::alphaNAT RPA135-3R305L | This study |
| RPA135-TAP | MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 RPA135-TAP::HIS3MX6 | 202233825 (Ghaemmaghami et al., 2003) |
| yTD16-1a | MATα ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 fob1[Delta]:::NAT-MX, rDNA copy no. ~25 | This study |
| WT in figures 2, 4B, and 4C | yTD27-1a | MATα ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 fob1[Delta]:::NAT-MX RPA135-TAP-HIS3, rDNA copy number ~25 + pCF4-LEU-GAL49 | This study |
| RPA135-F301S in figure 2 | yTD28-1a | MATα ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 fob1[Delta]:::NAT-MX RPA135-F301S-TAP-HIS3, rDNA copy number ~25, + pCF4-LEU-GAL49 | This study |
| yTD25-1a | MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 rpa49ΔC(186-416)::KANMX4 | This study |
| yTD11-1a | MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 rpa49ΔC(186-416)::HPHMX4 | This study |
| rpa49ΔC in figures 2, 4B, and 4C | yTD29-1a | MATα ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 fob1[Delta]:::NAT-MX RPA135-TAP-HIS3 rpa49ΔC(186-416)::HPH, rDNA copy number ~25, + pCF4-LEU-GAL49 | This study |
| rpa49ΔC | RPA135-F301S in figure 2 | yTD30-1a | MATα ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 fob1[Delta]:::NAT-MX RPA135-F301S-TAP-HIS3 rpa49ΔC(186-416)::HPH, rDNA copy number ~25, + pCF4-LEU-GAL49 | This study |
| RPA12-S6L in figures 4B and 4C | yTD31-1a | MATα ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 fob1[Delta]:::NAT-MX RPA135-TAP-HIS3 RPA12-S6L-KAN-MX, rDNA copy number ~25, + pCF4-LEU-GAL49 | This study |
| rpa49ΔC RPA12-S6L in figure 4B | yTD23_1a | MATα ade2-1 ura3-1 his3-11,15 trp1-1 leu2-3,112 can1-100 fob1[Delta]:::NAT-MX RPA135-TAP-HIS3 RPA12-S6L-KAN-MX rpa49ΔC(186-416)::HPHMX, rDNA copy number ~25 | This study |
| rpa49Δ in figures 6 and 7 | OGT15-9d | MATα his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 rpa49Δ::HPHMX | This study |
| SCOC2260 | Mata ade2-1 arg4 leu2-3,112 trp1-289 ura3-52 RPB6::TAP-KLURA3 rpa190-ΔLOOP-KANMX | This study |
| plasmids indicated in the legend | yTD48-1a | MATa his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 rpa190Δloop::KANMX | This study |
|--------------------------------|---------|-------------------------------------------------|------------|
| rpa190Δ loop                   | yTD51-2a | MATalpha his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 lys2Δ0 rpa190Δloop::KANMX RPA135-F301S::URA3Kl | This study |
| RPA135-F301S                   | yTD51-8a | MATa his3Δ1 leu2Δ0 lys2Δ0 met15Δ0 ura3Δ0 RPA135-F301S::URA3Kl rpa49Δ::HPHMX | This study |
| rpa49Δ in figures 5B and 5C    | yTD51-5a | MATa his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0 rpa49Δ::KANMX | This study |
| RPA135-F301S                   | yTD2-3b | MATa his3Δ1 leu2Δ0 ura3Δ0 lys2Δ0 rpa34Δ::NATMX | This study |
| rpa34Δ in figures 5B and 5C    | yCN223-2a | MATA his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 lys2Δ0 rpa49Δ::KANMX | This study |
| RPA135-F301S                   | yTD2-3a | MATa his3Δ1 leu2Δ0 ura3Δ0 lys2Δ0 rpa49Δ::NATMX | This study |
| rpa49Δ RPA135-F301S in figure 5C | yCN224-1a | MATalpha his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 lys2Δ0 rpa43Δ::KANMX | This study |
| rpa34Δ RPA135-F301S in figure 5B | yCN224-1a | MATalpha his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 lys2Δ0 rpa43Δ::KANMX | This study |
| rpa34Δ RPA135-F301S in figure 5B | yTD36-2b | MATalpha his3Δ1 leu2Δ0 ura3Δ0 lys2Δ0 rpa43Δ::NATMX RPA135-F301S::URA3 | This study |
| rpa34Δ rpa49Δ RPA135-F301S in figure 5B | yTD37-7d | MATalpha his3Δ1 leu2Δ0 ura3Δ0 lys2Δ0 rpa49Δ::NATMX rpa49Δ::HPHMX | This study |
| rpa49Δ RPA135-F301S in figure 5B | yTD37-3d | MATalpha his3Δ1 leu2Δ0 ura3Δ0 lys2Δ0 rpa49Δ::NATMX rpa49Δ::HPHMX RPA135-F301S::URA3 | This study |
| rpa49Δ in Figure 6 with the plasmids indicated in the legend | yCN225-1a | MATalpha his3Δ1 leu2Δ0 MET15 ura3Δ0 lys2Δ0 rpa14Δ::KAN-MX | This study |
| rpa14Δ in Figure 6 with the plasmids indicated in the legend | yCN225-1a | MATalpha his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 rpa14Δ::KAN-MX | This study |
| rpa14Δ RPA135-F301S in figure 5C | yTD38-3d | MATa his3Δ1 leu2Δ0 ura3Δ0 lys2Δ0 rpa14Δ::NATMX RPA135-F301S::URA3 | This study |
| rpa14Δ rpa49Δ RPA135-F301S in figure 5C | yTD39-8a | MATa his3Δ1 leu2Δ0 ura3Δ0 lys2Δ0 rpa14Δ::NATMX rpa49Δ::HPHMX RPA135-F301S::URA3 | This study |
| rpa49Δ in Figure 6 with the plasmids indicated in the legend | yTD53-1a | MATalpha his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0 KANMX6-pGAL::RPA12 | This study |
| rpa34Δ in Figure 6 with the plasmids indicated in the legend | yTD53-1a | MATalpha his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0 KANMX6-pGAL::RPA12 | This study |
| RPA12 alleles in figure 6 with the plasmids indicated in the legend | OGT30-1c | MATa his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 KANMX-pGAL::RPA12 | This study |
| rpa49Δ in Figure 6 with the plasmids indicated in the legend | OGT30-3c | MATalpha his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 rpa49Δ::HPHMX KANMX-pGAL::RPA12 | This study |
| OGT30-1a | MATa his3Δ1 leu2Δ0 lys2Δ0 rpa49Δ::HPH RPA135-F301S-TAP-HIS3 KANMX-pGAL::RPA12 | This study |
| OGT30-3a | MATa his3Δ1 leu2Δ0 lys2Δ0 rpa49Δ::HPH RPA135-F301S-TAP-HIS3 KANMX-pGAL::RPA12 | This study |
| yTD40-1a | MATa his3Δ1 leu2Δ0 lys2Δ0 rpa49Δ::HPH RPA135-F301S-TAP-HIS3 KANMX-pGAL::RPA12 | This study |
| yTD41-1a | MATa his3Δ1 leu2Δ0 lys2Δ0 rpa49Δ::HPH RPA135-F301S-TAP-HIS3 KANMX-pGAL::RPA12 | This study |
| RPA135-F301S in figure 6B | yTD42-1a | MATa his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 RPA135-F301S::URA3 KANMX-pGAL::RPA12 | This study |
| rpa49Δ RPA135-F301S in figure 6B | yTD43-1a | MATalpha his3Δ1 leu2Δ0 lysΔ0 ura3Δ0 rpa49Δ::HPH MX RPA135-F301S::URA3 KANMX-pGAL::RPA12 | This study |
| WT in figure 7 | yJPF159-1a | MATa his3Δ1 leu2Δ0 lys2- ura3Δ0 RPA135-TEV-ProA::KANMX6 | This study |
| rpa12 in figure 7 | yJPF165-1a | MATa his3Δ1 leu2Δ0 lys2- ura3Δ0 RPA135-TEV-ProA::KANMX6 HIS3MX::GAL::HA-RPA12 | This study |
### Supplementary table 3: Plasmids used in this study

| Name                     | Description                                                                 | Source                                      |
|--------------------------|-----------------------------------------------------------------------------|---------------------------------------------|
| pUC19-HPH                | Plasmid bearing HPH-MX4                                                     | (Berger et al., 2007)                       |
| pFA6-kanMX6              | Plasmid bearing KAN-MX6                                                     | (Longtine et al., 1998)                     |
| pFA6a-KanMX6-GAL::3HA    | Plasmid used for GAL promoter insertion                                      | (Longtine et al., 1998)                     |
| pFA6a-HA-KURA3           | Plasmid used for epitope switching                                          | (Sung et al., 2008)                        |
| pNOY80                   | Plasmid CEN6.ARS4, URA3, RPA135                                             | (Yano et al., 1992)                        |
| pVV190                   | Plasmid pFL44-A190 (2μ URA3 RPA190)                                         | (Van Mullem et al., 2002)                   |
| pRS316                   | CEN6 A12, URAS, URA3                                                       | (Sikorski & Hieter, 1989)                   |
| pCR4-HIS3                | HIS3                                                                       | (Sikorski & Hieter, 1989)                   |
| pGL190_3                 | (2μ URA3 RPA190-E1274K) selected from randomly mutagenized pVV190         | This study                                  |
| pGL190_11                | (2μ URA3 RPA190-C1493R) selected from randomly mutagenized pVV190         | This study                                  |
| pGL190_23                | (2μ URA3 RPA190-LJ262P) selected from randomly mutagenized pVV190         | This study                                  |
| pGL135_6prim             | (CEN4 URA3 RPA135-R379G) selected from randomly mutagenized pNOY80       | This study                                  |
| pGL135_54                | (CEN4 URA3 RPA135-Y252H) selected from randomly mutagenized pNOY80       | This study                                  |
| pGL135_33                | (CEN4 URA3 RPA135-F301S) selected from randomly mutagenized pNOY80       | This study                                  |
| pTD1_3b_135TAP           | Plasmid pNOY80 bearing RPA135-TAP-HIS3 obtained by homologous recombination using a PCR-amplified fragment generated with oligos 835 and 836 and genomic DNA of strain 20223825 as template. | This study                                  |
| pTD2_6c_135TAP           | Plasmid pNOY80 bearing RPA135-F301S-TAP-HIS3 obtained by homologous recombination using a PCR-amplified fragment generated with oligos 835 and 836 and genomic DNA of strain 20223825 as template. | This study                                  |
| pTD5                     | Plasmid pTD1_3b_135TAP deleted of URA3 using NsiI and SdaI digestion and self-ligation. | This study                                  |
| pTD6                     | Plasmid pTD2_6c_135TAP deleted of URA3 using NsiI and SdaI digestion and self-ligation. | This study                                  |
| Ycp50-26                 | URA3 ARS/CEN RPA49                                                         | P. Thuriaux, unpublished (Sikorski & Hieter, 1989) |
| pRS316-A12               | pRS316 vector ligated with a PCR-generated fragment using oligos 1554 and 1555 and yeast genomic DNA as template cut BamHI-XbaI, and cloned at same site. | This study                                  |
| pRS316-A12-AvrII         | PCR mediated mutagenesis to introduce AvrII site in pRS316-A12 using oligos 1714 and 1715. | This study                                  |
| pRS316-A12-S6L           | S6L Allele of RPA12 isolated as a suppressor of the rpa49A growth defect, selected from a PCR-mediated random mutagenesis of pRS316-A12-AvrII. | This study                                  |
| pRS316-A12-S6L-KAN       | Plasmid obtained using pRS316-A12-S6L modified by homologous recombination using a PCR-amplified fragment generated with oligos 1682 and 1559 and pFA6-kanMX6 as template. | This study                                  |
| pRS316-A12-T49A          | T49A Allele of RPA12 isolated as a suppressor of the rpa49A growth defect, selected from a PCR mediated random mutagenesis of pRS316-A12-AvrII. | This study                                  |
| pRS316-A12-DCter         | Plasmid obtained using pRS316-A12 modified by homologous recombination using a PCR-amplified fragment generated with oligos 1371 and 1559 and pFA6-kanMX6 as template. | This study                                  |
| pTD9                     | Plasmid obtained using pRS316-A12-T49A modified by homologous recombination using a PCR-amplified fragment generated with oligos 1371 and 1559 and pFA6-kanMX6 as template. | This study                                  |
| pTD10                    | Plasmid obtained using pRS316-A12-S6L modified by homologous recombination using a PCR-amplified fragment generated with oligos 1371 and 1559 and pFA6-kanMX6 as template. | This study                                  |
| Plasmid          | Description                                                                 | Source                  |
|------------------|-----------------------------------------------------------------------------|-------------------------|
| pCJPF4           | pFL36cII with LEU2 marker, CEN4, containing RPA49 coding region              | This study              |
| pCJPF4-GAL49-1   | Plasmid obtained using pCJPF4 modified by homologous recombination using a PCR-amplified fragment generated with oligos 624 and 625 and pFA6a-KanMX6-GAL::3HA as template. | This study              |
| pMAX1            | Plasmid including Pol I promoter used for in vitro assays                   | (Pils et al. 2016)      |
| pUC 19 tail g- TER elongated | Plasmid containing tailed template DNA containing 18 S rDNA for non-specific in vitro transcription | (Merkl et al. 2014)    |
| Sequence                                                                 | N°  |
|-------------------------------------------------------------------------|-----|
| gctgctttatagaaccaatctgacgcacaaacccagagagcaaccaacaagtcgtatatacgatgttaatcagtagtttaaatgcgagctcgtttaaac   | 208 |
| ceactactcaggtgcgttttaacacccatgttcactattgcagtcgttatcaacctttttgcacttttcttagtagagaattcgagctcgtttaaac   | 624 |
| cactttcaatttcgattccggacacactttttcgagacgatcctttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttt
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Figure 1
Figure 2.
Figure 3.
Figure 4.
Figure 5.
Figure 6.
|    | Tailed Template | Promoter-Dependent |
|----|-----------------|--------------------|
| ΔA12 | 4 nM 12 nM | 4 nM 12 nM |
| WT  | 4 nM | 4 nM |

Figure 7.
Figure 8

Free monomeric

ITC

Pol I \textit{rpa49\Delta}

Pol I \textit{rpa49\Delta} + suppressors
Suppl. Figure 1: Genetic interaction mapping (GIM) to identify suppressor mutations in SGR1 and SGR2. (A) Schematic representation of GIM interaction assay. Enrichment ratio between control and SGR are used as read-out to map genetic interactions. (B) Relative enrichment of each barcode (black cross) along chromosome (in red) are used to map genetic linkage. Green curve represents mean in a sliding windows of 20 barcodes. Local maximum in such curve was used to identify locus of interest: RPA49 as positive control (upper panel, chr 14); RPA135 in SGR1 (middle panel, chr 16); RPA190 in SGR2 (lower panel, chr 15).