A Genetic Assay for Transcription Errors Reveals Multilayer Control of RNA Polymerase II Fidelity

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Abstract

We developed a highly sensitive assay to detect transcription errors in vivo. The assay is based on suppression of a missense mutation in the active site tyrosine in the Cre recombinase. Because Cre acts as tetramer, background from translation errors are negligible. Functional Cre resulting from rare transcription errors that restore the tyrosine codon can be detected by Cre-dependent rearrangement of reporter genes. Hence, transient transcription errors are captured as stable genetic changes. We used this Cre-based reporter to screen for mutations of Saccharomyces cerevisiae RPB1 (RPO21) that increase the level of misincorporation during transcription. The mutations are in three domains of Rpb1, the trigger loop, the bridge helix, and in sites involved in binding to TFIIS. Biochemical characterization demonstrates that these variants have elevated misincorporation, and/or ability to extend mispaired bases, or defects in TFIIS mediated editing.

Introduction

Accurate transcription is an essential step in accessing the genetic information stored in genes. In eukaryotes, transcription of DNA into mRNA is carried out by RNA polymerase II (Pol II), an enzyme comprised of 12 subunits. While there has been extensive research on core Pol II, including biochemical and detailed structural information [1–7], less is known about how the accuracy of transcription is controlled. The net misincorporation rate is estimated to be about one per 10^5 bases [8,9]. This reflects initial post-incorporation stage [12,25,26]. Indeed, it has been demonstrated that in vitro TFIIIS preferentially promotes cleavage of the mismatched 3′ end of nascent RNA [27].

The interpretation of phenotypic changes associated with alterations in RNA polymerase is complicated by a mix of direct...
Author Summary

Mistakes made during the synthesis of messenger RNAs have been difficult to detect, both because mRNAs can be short lived, and because the translation of mRNAs into proteins has a much higher error rate that masks transcription errors. We present here a highly sensitive genetic screen that detects transcription errors and use it to identify mutations that increase the error rate of RNA polymerase II. The screen incorporates a new principle that allows transient transcription errors to cause permanent genetic changes. The screen is based on suppression of a missense mutation (cre-Y324C) in the active site of the Cre recombinase. Infrequent and transient transcription errors that restore the original codon for Y324 cause the Cre-dependent activation of a reporter gene. Background from translation errors is negligible because Cre acts as a tetramer in which all four subunits require the active site tyrosine. Transcription errors as low as ~10^-6 can be detected. We identify rpb1 mutations that define four classes, those that have increased (1) misincorporation, (2) extension of a misincorporated base, (3) both misincorporation and extension, and (4) those that block the activity of the transcription proofreading factor, TFIIS.

Results

Transcription fidelity assay based on Cre recombinase

Assays of transcription fidelity based on nonsense or missense suppression are problematic in part due to the high background caused by translational errors [10-12,30]. In the approach reported here, we rely on a requirement for an active tetramer to reduce the contribution of translation errors on the background. In addition, previous approaches have required a continual production of transcription errors to produce the monitored phenotype. In our approach, transient transcription errors can result in a stable genetic change. The system involves three parts, 1) a missense mutation in the active site tyrosine of Cre recombinase, 2) a Cre-dependent recombination reporter substrate with a very low Cre-independent rate of recombination and 3) promoters that give low level expression of the mutant cre allele. These three aspects of the system are discussed separately below.

Cre mutations

Cre functions as a homotetramer, recognizing two 34 bp DNA sequences termed loxP and promoting recombination of their flanking DNAs [31,32]. Depending on the orientation of the two loxP sites, the flanking DNA is either inverted (intrachromosomal antiparallel), excised (intrachromosomal direct), or exchanged (interchromosomal) [33]. The active site tyrosine (Y324) of Cre recombinase is essential for its activity [34]. All four subunits of the tetramer become covalently attached to strands of the DNA via those tyrosine 324 side chains during the recombination reaction. Chimeric enzymes assembled from mutant and WT subunits are catalytically inactive [35-39]. Translation errors that produce a WT monomer will not result in active Cre, whereas a transcription error that restores the WT codon can be translated into multiple WT subunits and assembled into an active Cre tetramer. The experiments presented here focus on the Y324C allele. This TAT G TGT mutation requires Pol II to misincorporate an adenine opposite a cytosine in the template strand in order to suppress the cre-Y324C defect (Figure 1A). This G A transcription error is among the most common mistakes made by E. coli RNAP in vitro and in vivo [40] and by S. cerevisiae [19].

Cre reporters

We created a reporter for Cre activity based on HIS3 into which we placed an artificial intron [41] carrying a loxP site, HIS3-AI2lox (Figure 1B, construct (i)). A Cre activatable derivative, his3-AI2loxMX, was made by insertion of a kanMX cassette (Figure 1B, construct (ii)) [42,43]. However, the Cre-independent His^6 background from this reporter was too high (~2×10^-6 His^+ among total cells). We created a HIS3 based Cre reporter system with very low Cre-independent background by placing the downstream loxP site, splice acceptor and C-terminal portion of a his3-AI2loxMX reporter in an inverted orientation at a position 3 kb distal of HIS3 beyond the DED1 gene (Figure 1B, construct (iv)). The N-terminal portion of his3-Allux is at the normal chromosome XV position for HIS3 and is marked by hygMX, while the C-terminal portion of his3-Allux was marked with a zeoMX cassette. Cells with this inverted lox reporter, designated his3-AI2loxMX, are His^+, hygromycin resistant and zeomycin resistant. If Cre is active, it can cause inversion of the interval between the two loxP sites to generate the functional HIS3-AI2lox1 gene (Figure 1B, construct (v)). These cells are His^6-, hygromycin resistant and zeomycin resistant. The frequency of Cre-independent His^6 cells among total cells with this system is <10^-6 (Figure 2C).

Low level expression

In the experiments described here it was anticipated that the mutant Cre-Y324C protein should act as a dominant inhibitor of wild type Cre protein. That is, assembly of mutant subunits into the tetramer will block its function. Therefore, we expected that it is important to have very low levels of expression in order to favor assembly of tetramers from translation of one mRNA. This condition promotes the detection of transcription errors that restore the Tyr codon in the cre-Y324C transcript. We used two approaches to have low levels of transcription (Figure 1A). First we placed the cre gene under the control of the GAL1 promoter but grew the cells in the presence of glucose. Under these glucose repressed conditions, expression was expected to be rare. In the second approach, we placed the cre-Y324C allele under the control of the HO promoter so that it would be expressed only in that half of the cell population that had already divided once before [44,45].

Proof of principle

Our initial experiments used the PGAL1-cre-Y324C expression system as the transcription error substrate (inserted into the BUD5 gene) and the his3-AI2loxINV Cre activity reporter. For cells with a WT Pol II (GRY3739) the frequency of His^6 cells among the total population is similar in the absence of Cre or with the PGAL1-cre-Y324C construct (Figure 2A&C). In this assay, patches of cells grown on rich media (YPD) were replica plated onto synthetic complete medium lacking histidine. The starting strains cannot
grow, but rare His\(^+\) cells within the patch can grow into small colonies or papillae. The $P_{GAL1}$-cre-Y324C strain lacking TFIIS ($dst1A$ GRY3742) has an elevated level of His\(^+\) cells (Figure 2A&C) that is not observed when the cre gene is absent. We interpret the elevated level of papillation in the TFIIS defective strain as a reflection of transient Cre activity caused by uncorrected G $\rightarrow$ A errors during transcription in the mutant cre-Y324C gene that restore the UAU tyrosine codon in the mRNA (Figure 1A).
Figure 2. Suppression of cre-Y324C is elevated in strains with Pol II fidelity defects. (A) PGAL1-cre-Y324C strains grown in glucose on plates lacking histidine show increased numbers of cells capable of growing microcolonies in strains with known transcription fidelity defects. Two representative patches are shown for each variant. (B) PGAL1-cre-Y324C used to identify new transcription fidelity mutants. (C) Mean frequency of His⁺ cells relative to total viable cells in PGAL1-creY324C or PGAL1-creY324C, his3-A12loxINV strains with Pol II mutants. Cultures were grown overnight in 1 ml of YPD at 30°C and plated on SC-His and 10-fold serial dilutions onto YEPEDA plates. Plates were incubated at 30°C for 3 days and individual colonies were scored. Error bars show standard error.

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demonstrate that the His⁺ cells were not a result of an elevated reversion rate for the cre-Y324C mutation in TFIIS defective cells, 11 independent His⁺ cells from the dst1 strain were crossed to a cell carrying an ade6-A12loxMX reporter and shown to be still defective for Cre recombination in vivo. In addition, the cre gene sequenced from 12 more His⁺ colonies still carried the cre-Y324C mutation. To determine whether the increase in His⁺ cells might be the result of elevated Cre-independent recombination between the loci sites in TFIIS defective strains, we compared the frequency of His⁺ cells in the absence of the cre gene and observed that WT and dst1 strains gave similar levels. These results confirm that TFIIS has a role in vivo in editing RNA errors and promoting transcription fidelity.

RPB9 has been identified as a nonessential subunit of Pol II that promotes accurate start site selection, efficient elongation, and contributes to transcription fidelity [11,23,46-50] and regulates mismatch extension [24]. Consistent with a role for the Rpb9 subunit in promoting transcription fidelity in vivo, we find that a null allele of RPB9, rpb9-A0 (G3574), shows an elevated His⁺ papillation rate in the his3-A12loxINV assay (Figure 2A&C).

In order to test a collection of rpb1 mutations, we created a strain (G3585) with the PGAL1-cre-Y324C substrate, the his3-A12loxINV reporter and the rpb1-natMX null allele complemented by RBPI on a URA3 selectable plasmid (pJDI220). We made rpb1 variants on a LEU2 based plasmid (pJS757) and substituted the mutant alleles for the RBPI wild type allele by transforming the URA3 plasmids and selecting for loss of the URA3 plasmid with 5-FOA which selects against Ura⁺ cells [51]. Cells with the rpb1-E1230K (rpb2I-24) mutation (pJS932), which blocks the ability of TFIIS to bind to Pol II [52], show an elevated number of His⁺ papillae (Figure 2A&C). This is consistent with the results presented above for the TFIIS defective strain (dst1A0). We previously described rpb1-E1103G (pJS781) as causing increased misincorporation during transcription [19]. That allele was also among those identified as having elevated misincorporation rates in vitro [3]. When put into the his3-A12loxINV reporter strain with cre-Y324C under the control of the GAL1 promoter, it showed a modest increase in His⁺ papillation (Figure 2A&C).

Genetic screen for new rpb1 mutants

To use the his3-A12loxINV reporter as an effective screen for new rpb1 mutations that reduce fidelity, we addressed the problem of how to get widespread but low level expression of the cre-Y324C gene. The ideal condition would be the one that produced one transcript per cell. The GAL1 promoter in the presence of glucose is repressed [53], but low levels of transcription in a population probably reflects some cells with transcripts and most cells with none. The promoter for the HO gene is under complex regulation so that it is only expressed late in the G1 phase of the cell cycle, only in cells that have the α or α mating phenotypes (not a/α “diploid” cells), and only in cells that have divided at least once before, so called mother cells [44,45]. We placed the cre-Y324C gene under the control of the HO promoter at its normal position on chromosome IV. Only half of the cells (mothers) in the population will be expressing PHO-cre-Y324C, and half of those will be expressing it for the first time. We reasoned that this would create a condition where rare transcripts that have an error that restores the tyrosine codon would produce active Cre recombinase that would not be in competition with the inactive monomers for the assembly of active Cre tetramer.

To make it easy to introduce rpb1 variants into this system, we made a yeast strain (G35258) with the P_HO-cre-Y324C substrate, the his3-A12loxINV reporter, and the rpb1-natMX deletion complemented by RBPI on a URA3 2-micron based plasmid (pJS725). As above, this makes it possible to substitute in rpb1 variants on a LEU2 based vector by plasmid shuffling, selecting against the URA3 RBPI with 5-FOA [51]. Figure 2B illustrates the clear distinction between RBPI and the transcription fidelity mutant rpb1-E1103G in the number of His⁺ papillae with the his3-A12loxINV reporter and the P_HO-cre-Y324C substrate. The increase is dependent on the cre-Y324C gene. Similarly, rpb1-E1230K, which blocks TFIIS function, gives a clearly elevated signal in this assay.

To identify additional rpb1 alleles that cause reduced transcription fidelity in vivo, we screened a mutagenized pool of a LEU2 CEN based plasmid carrying RBPI (pJS757). The pool was transformed into G35258 selecting Leu⁺ colonies picked and arrayed as patches. The patches were replica plated to FOA plates to select against the URA3 RBPI plasmid. From the FOA plates the patches were replica plated to SC-His media to detect patches with elevated levels of Cre-mediated His⁺ papillae. Mutant candidates were struck for single colonies, and retested as patches. The LEU2 based plasmid from those that repeated the elevated level of His⁺ papillae was recovered into E. coli and retransformed into G35258 to confirm that the mutation causing the elevated Cre activity was on the plasmid. Those that passed that test were sequenced to identify the rpb1 mutation responsible for the transcription infidelity phenotype. In all, over 12,000 transformants were tested from which eight rpb1 variants were identified. These included re-isolating rpb1-E1103G, plus seven new alleles: rpb1-G8235, -A1076T, -A1076V, -M1079I(T1548I), -K1132E, -T11411(G888D,I1237T), and -S1229F. The phenotype of the rpb1-T11411(G888D,I1237T) variant was shown to be a consequence of the T11411 substitution by testing variants with each of the separate mutations. Patches demonstrating the phenotypes of these variants are shown in Figure 2B. In addition, we show results for an allele rpb1-T1113P, isolated from a screen for alleles that elevate transcription slipage [15].

Quantification of the increase in transcription errors was accomplished by measuring the frequency of His⁺ cells as determined by growth on medium lacking histidine (SC-His) normalized to growth on rich media. The mean frequency from 8 or more cultures was determined for each strain and shows that the mutations identified by our screen increase the frequency of cells with the His⁺ phenotype 7–50 fold compared to the wild type strain (Figure 2C).

In vitro measurement of transcription fidelity

In the in vivo screen, increase of the frequency of His⁺ papillae by the mutations in RPB1 can be indirect. For instance, it could be caused by the increase of cre expression because of an increase in initiation by the mutant Pol II. It is noteworthy that the His⁺
frequency is much higher in the $P_{HOG1}$-cre system than in the $P_{GAL1}$-cre system for the $rpb1$-E1103G allele. In contrast the His+ frequencies for the $rpb1$-E1230K allele are similar in the two systems. Whether this reflects a differential level of sensitivity for these promoters to these defects has not been determined.

Furthermore, a higher net frequency of error-containing RNA may be caused not only by the higher frequency of misincorporation (as in the $rpb1$-E1103G mutant), but also by higher efficiency of mismatch extension and/or on lower efficiency of RNA editing (like in a $DST1$-deficient strain or in the $rpb1$-E1230K mutant, defective in TFIIS binding) [52]. Below we describe the results of three in vitro tests designed to address the biochemical basis for the low-fidelity phenotype of the alleles identified in the genetic screen: 1) the ability of Pol II to select cognate NTP, 2) the capacity to extend a mismatch, and 3) the ability to remove misincorporated NMP by TFIIS-mediated editing.

### Competition fidelity assay

To determine whether any of the new alleles of $rpb1$ have direct impact on cognate NTP selection, we tested in vitro the level of misincorporation errors by the Pol II variants. Misincorporation rates and transcription fidelity may depend on a sequence context [4]. Therefore, we employed a DNA corresponding to the region surrounding codon 324 (TGT) in the cre-Y324C gene as the template for the assembly of the promoter- and factor-independent Pol II elongation complexes [54] used in these fidelity assays. The elongation complex (U10) was stalled before incorporating AMP and the frequency of GMP to AMP transition error was directly measured using a “competition” assay for transcription fidelity [20], which provides fidelity values nearly identical to those obtained by conventional fidelity assay for the wild type and mutant variants of Pol II [20,55]. A similar assay has been described for T7 DNA polymerase [56]. The competition assay employs differential mobility in denaturing polyacrylamide gels of short RNA species of the same length, but different composition to distinguish cognate from misincorporation events. To determine the frequency of misincorporation, U10 (schematically depicted in Figure 3A) was incubated with a mix of GTP in a low (20 nM) concentration and different higher (0.5, 0.75 and 1.5 mM) concentrations of ATP. In these conditions, the reaction products of both cognate (GMP) and non-cognate (AMP) incorporation were separated (Figure 3B, lanes 3–13) and quantified (Figure 3C). To determine the misincorporation frequency of AMP in place of GMP in the presence of equal concentrations of cognate and non-cognate substrates, the ratio of non-cognate to cognate products was normalized to (divided by) the ratio of non-cognate (ATP) to cognate (GTP) substrate concentrations. The resulting misincorporation frequency was similar for different concentrations of ATP tested suggesting that it faithfully reflects fidelity of substrate selection in this position, independent of the substrate concentration. This assay is particularly useful way to qualitatively compare the relative misincorporation of various alleles. It is clear that Pol II carrying the $rpb1$-E1103G mutation misincorporates more frequently than wild type Pol II (Figure 3B, lane 8), consistent with its previously reported defect in NTP selection [19]. The $rpb1$-A1076T, $A1076V$, and $T1113P$ substitutions also noticeably increase the frequency of misincorporation (lanes 5, 6, and 9). The $RPB1$-A1076 residue is located in the trigger loop, a mobile
element of the catalytic subunit implicated in transcription fidelity maintenance by genetic and biochemical analyses [3,19]. In contrast, mutations located in the TFIIS-binding domain (\textit{rpb1-K1132E}, \textit{rpb1-T1141I} and \textit{rpb1-S1229F}) and one mutation in the trigger loop (\textit{rpb1-M1079I}) do not appear to change GMP to AMP transition frequency \textit{in vitro}.

Mispaired base extension assay

Next, we tested whether the newly identified \textit{rpb1} alleles alter the ability of Pol II to extend the mismatch by incubating the complex with 1 mM non-complementary ATP and 0.5 mM next cognate UTP to allow misincorporation and mismatch extension. Under these conditions, wild type Pol II does not easily extend the A11 mismatch and pauses at the U12 and A13 positions (Figure 4A, lane 3). Apparently, the mismatch continues to present an obstacle to efficient transcript elongation, consistent with the recent report by Knippa and Peterson [24]. Inefficient mismatch extension, similar to wild type Pol II was observed in the Pol II variants carrying mutations in the TFIIS-binding domain (\textit{rpb1-K1132E}, \textit{rpb1-T1141I} and \textit{rpb1-S1229F}) and one mutation in the trigger loop (\textit{rpb1-M1079I}) do not appear to change GMP to AMP transition frequency \textit{in vitro}.

\textit{rpb1-M1079I} mutant, which does not display increased frequency of misincorporation, provides an explanation for the identification of this allele as error-prone in the \textit{in vivo} screen. Evidently, relatively fast mismatch extension interferes with the post-incorporation error removal by

\textbf{Figure 4. Mismatch extension by Pol II variants carrying mutations in Rpb1.} (A) Products of simultaneous misincorporation and mismatch extension. The U10 complex was incubated with a mixture of 1 mM ATP and 0.5 mM UTP, or 1 mM ATP, 0.5 mM UTP and 0.5 mM GTP where indicated. The original U10 RNA, A*11 mismatch and mismatch extension products are indicated at left. The arrows at right mark the mismatch (A*11) and mismatch extension products. (B) the fraction of U14 mismatch extension product from the sum of A11, U12, A13 and U14 products was quantified; results of three independent experiments were averaged. The error bars show standard deviation. (C) Products of consecutive misincorporation and mismatch extension. U10 complexes were incubated in 1 mM ATP for 10 minutes to obtain complexes containing an RNA with a 3’ mismatch (A*11). A sample of A*11 complex for each Pol II variant was taken, and 0.5 mM UTP was added to initiate extension of the mismatch. The extension products were analyzed at the indicated times after the addition of UTP. RNA A*11 and extension products are indicated at left. (D) Mismatch extension was quantified as in (B) and plotted versus time. The error bars show standard deviation of three independent experiments. doi:10.1371/journal.pgen.1004532.g004
TFIIS-mediated cleavage or the proteolytic degradation of irreversibly arrested Pol II, thus increasing the fraction of the full-length transcripts containing the error. The in vivo and in vitro properties of Pol II carrying the rpb1-M1079I substitution provide direct experimental evidence that recognition of incorporated mismatches by core Pol II significantly contributes to fidelity maintenance.

TFIIS-dependent editing

The effect of the newly identified mutations on interaction of Pol II with TFIIS has been tested by treating the A*11 elongation complex carrying the 3’-end mismatch with TFIIS, and observing the extent of 3’-RNA cleavage (Figure 5). The 3’-mismatched AMP and the complementary penultimate UMP are removed after one minute incubation with TFIIS in the major fraction of the wild type complexes, resulting in appearance of a 9-nt RNA (C9). Note that the C9 product was barely detectable in the initial A*11 elongation complexes (lane 3 in Fig. 5 A and B). The shorter RNA cleavage products (A8, A7, and G6) are also detected, especially after incubation with higher concentration (600 nM) TFIIS (Figure 5B, lane 4). The appearance of the 3’ cleavage products as short as 6 nt is somewhat unexpected, considering that the RNA-DNA hybrid in Pol II elongation complex is 8-bp long [57]. Nevertheless, we use appearance of the short cleavage products, along with the major C9 product, to judge the efficiency of TFIIS-induced RNA cleavage.

As expected, substitutions in the TFIIS binding site (rpb1-S1229F and rpb1-E1230K) dramatically decreased Pol II susceptibility to TFIIS-induced mismatch removal. The C9 cleavage product appears only when the elongation complexes are treated with the higher concentration (600 nM) TFIIS, and no shorter products are detected (compare lane 4 with lanes 13 and 14 in Figure 5A&B). A less pronounced, but still substantial decrease in the sensitivity to TFIIS is observed for rpb1-G823S, K1132E, and rpb1-T1141I mutants (lanes 5, 11, and 12). It is likely that K1132 and T1141 substitutions directly reduce binding of TFIIS. The G823S substitution in the bridge helix may alter TFIIS binding, but alternatively could be involved in TFIIS-induced transcript cleavage, and/or affect Pol II backtracking. The effect of T1141I substitution was weaker than the effects of the other three substitutions in the TFIIS binding site (note the appearance of the short cleavage products in Figure 5B, lane 11). It is interesting that Rpb1-G823S, which shows a decreased sensitivity to TFIIS similar to Rpb1-K1132E substitution, also slightly promotes misincorporation and mismatch extension in vitro (Figures 3 and 4). All other mutants tested for susceptibility to TFIIS were similar to wild type Pol II (Note the A8, A7 and G6 products in Figure 5B, lanes 4 and 6–10).

Discussion

We present here a solution to problematic issues about the measurement of transcription fidelity in vivo. Previous results suggested that G to A transitions represent the major class of errors generated by yeast Pol II [19] and by the E. coli RNA polymerase in vitro and in living cells [58]. Using that information, we developed a second generation assay specific for the detection of G to A transcription errors. The assay is based on a mutation of the active site TAT tyrosine codon of the Cre recombinase to TGT. Rare Pol II errors that restore UAU to that position allow the production of active Cre protein whose activity is detected by inversion of a 4.8 kb interval to restore a selectable phenotype (Hisr). The requirement that each of the four subunits of the Cre tetramer is active eliminates the background from translation errors. The assay converts a transient Cre activity into a stable genetic change allowing the detection of Cre activity at frequencies less than 10−5. This sets a lower bound on net transcription error rates, but it is likely that it underestimates the real level of uncorrected G to A substitutions during transcription because the efficiency of assembling an active Cre tetramer and the efficiency of the recognition and recombination of the lox sites in the reporter gene are unknown.

Identification of Rpb1 regions involved in control of fidelity in vivo

The random mutagenesis of the entire RPB1 (RP021) gene yielded three groups of amino acid residues that are highly clustered in the X-ray structure of Pol II [18]. The rpb1-A1076T/ V, rpb1-M1079I, and rpb1-E1103G mutations alter the trigger loop, a domain of Pol II that closes over the active site and has been demonstrated to influence fidelity in vitro [3,19]. The rpb1-T1113P mutation targets a residue in the immediate vicinity of E1103 (Figure 6A). The rpb1-G823S allele (Figure 6B, right
Figure 6. Rpb1 domains involved in control of transcription fidelity. (A) Clusters of residues implicated in transcription fidelity control by genetic screening and biochemical analyses. RNA, DNA, substrate NTP, and the closed trigger loop (TL) are from pdb 2E2H; this structure is aligned with the X-ray structure of TFIIS taken from the backtracked Pol II/TFIIS complex (pdb 3PO3, shown in light red). (B) Location of Rpb1-A1076, M1079, E1103 and T1113 residues between the open (yellow; pdb 3PO3) and the closed (cyan; pdb 2E2H) states of the TL. On the right: Rpb1-G823 (yellow) is located next to the site of the bridge helix bending. The long-distance movement of L1081 residue of the TL associated with the bridge helix bending and the loop opening is shown. The upper right corner shows a view along the long axis of the bridge helix. (C) The cartoon illustrates...
the major conformational changes near the active site associated with the TL closure and opening and identifies the four hinges in the trigger loop and bridge helix. At the bottom: an alignment of the TL and the bridge helix amino acid sequences from the yeast Pol II and T. thermophiles RNAP with all hinges and L1081 residue highlighted. The TL and bridge helix crosstalks are indicated by double-headed arrows.

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Panel) alters the bridge helix, a domain that interacts with the active site and with the trigger loop. Substitutions in the bridge helix of E. coli RNA polymerase have been demonstrated to reduce the fidelity of transcription in vitro [59]. Notably, the changes identified in this work directly target the flexible hinge regions of the trigger loop and the bridge helix that were identified by molecular dynamics simulations based on X-ray crystallography and confirmed by mutational analysis of yeast Pol II [20]. The hinges, H1 and H2 in the trigger loop and H3 and H4 in the bridge helix, according to the nomenclature from [20] (Figure 6C) undergo conformational changes associated with NTP binding, sequestration, catalysis and translocation. The trigger loop and bridge helix residues forming the hinges have been implicated in transcription control [19,55,59,60].

Another cluster of mutations alters the TFIIS binding site (Figure 4A, residues shown in red) [61]. The rpb1-S1229F mutation likely reflects a defect in the interaction of Rpb1 and TFIIS, similar to the well characterized rpb1-E1230K allele [52]. The rpb1-K1132E and rpb1-T1141I substitutions also alter positions close to where TFIIS binds [61]. These mutations may impair TFIIS-dependent error editing by decreasing TFIIS binding, similar to mutations of E1230 and S1229. Alternatively, they might affect the backtracking capabilities of Pol II, a mechanism related to Pol II translocation and likely affected by the G823S substitution. The precise characterization of TFIIS cleavage mechanisms affected by mutations identified here is beyond the scope of this work. Most importantly, our results represent a direct demonstration of proofreading function of TFIIS in living cells. Although there is firm evidence that TFIIS plays a major role in correction of transcription errors in vitro, several attempts to demonstrate the similar activity in vivo were inconclusive [10,11]. Our results are consistent with those by Koyama and co-workers showing a major contribution of TFIIS to Pol II fidelity in yeast [12,21].

**Pol II fidelity is under multilayer control in vivo**

Our in vivo approach combined with the biochemical in vitro validation identified at least three intrinsic and one factor-dependent mechanism for faithful transcription in living cells. The TFIIS-dependent mechanism has been discussed above. The first intrinsic mechanism includes regulation of the trigger loop movement. Interaction of Rpb1 E1103 residue with the H2 hinge (residues 1095–1099) of the trigger loop has been previously proposed to delay the trigger loop closure thus slowing down transcription elongation and supporting fidelity maintenance [3,19,23,62]. The observation that rpb1-T1113P substitution renders transcription error-prone indicates that the T1113 residue plays the same or similar role as E1103. Notably, the recent molecular dynamic simulations of the Pol II trigger loop opening and closure revealed the potential interaction of T1113 with the trigger loop predicting that substitutions of T1113 residue should increase transcription elongation rate [63]. Our work provides direct proof that the mechanism dependent on E1103 and T1113 interaction with the trigger loop acts in vivo.

The second potential mechanism includes H1 hinge of the trigger loop (Rpb1-A1076/M1079) and H4 hinge of the bridge helix (Rpb1-G823) that may crosstalk through the adjacent L1081 (wedging) residue of the trigger loop [64] (Figure 6B&C). Because conformational changes of the bridge helix and the trigger loop are implicated in translocation [64–66] our identification of error-prone mutants in the mobile hinges of these two structural elements suggests a possible connection of the Pol II translocation cycle and cognate NTP selection. The mechanism of this link remains to be established.

The third intrinsic mechanism revealed by our work does not immediately follow from the extensive in vitro studies of transcription fidelity. It involves inhibition of the mismatch extension with the next cognate NMP. This mechanism, apparently mediated by the trigger loop and bridge helix, promotes Pol II pausing or arrest after misincorporation. Our finding that several mutations selected in the Cre-based genetic screen promote mismatch extension in vitro strongly argues that slow extension of a mismatch in the nascent RNA plays a major role in faithful transcription in living cells by allowing correction to occur. The rpb1-M1079I allele identified here appears of special interest, because it does not affect NTP selection by Pol II, but clearly promotes mismatch extension. Because the net frequency of in vivo transcription error occurrence correlates with the propensity of E. coli RNA polymerase to backtrack within a given sequence context [58], we are currently investigating the effect of rpb1-M1079I substitution on the mismatch-induced transcription arrest.

The slow mismatch extension may enable correction of the error by TFIIS-mediated transcript cleavage [27]. The arrest may also provide time for elimination of the flawed transcript by ubiquitin-mediated proteolytic degradation of Pol II [67]. One can imagine rpb1 alleles that impair accuracy in nucleotide selection, but, due to a reduced transcription elongation rate [3,20,60] provide increased opportunity to detect and remove misincorporated RNPs. Thus, slow elongation that results in poor mismatch extension could reduce production of the full-length error-containing Cre mRNA, counter-acting the defect in substrate selection. Accordingly, faster elongation might further decrease overall fidelity of the mutants with impaired NTP selectivity, such as rpb1-A1076T, A1076V, E1103G and T1113P [19].

In conclusion, we developed a reliable experimental approach to monitor transcription fidelity in vivo. Using this tool we will characterize the role of other core Pol II subunits, as well as known transcription elongation factors, such as TFIIF, Sp4/5 and Sp6 in transcription fidelity maintenance. This methodology will allow for isolation of mutants affecting transcription fidelity and thus will promote identification of new genes and mechanisms related to the accuracy of transcription. These experiments also highlight the complications associated with assigning a specific phenotype solely to the fidelity of transcription. The rpb1-E1103G mutation increases the transcription error rate. When combined with a defect in TFIIS (ds1DA), which corrects transcription misincorporation errors, the double mutant is dead. It is tempting to conclude that the death is the direct result of an error catastrophe of too many transcription errors resulting in too many incorrect RNAs and defective proteins encoded by them. However, TFIIS also has a role in transcription initiation at some promoters and the lethality of the double mutant could reflect some more complicated interplay of the properties of the defective RNA polymerase and the changes in the transcriptome caused by the TFIIS defect. Similarly, it was possible that an
increase in the frequency of His\(^+\) cells in our genetic assay reflected changes in the expression level of the cre reporter genes caused by a change in the transcriptome unrelated to reduced fidelity. A recent survey of changes in the yeast transcriptome caused by RNA polymerase variants further emphasizes this biological feature [28]. They found changes in transcription start sites, altered relative transcript levels, and changes in splicing efficiency. Thus, it was important to include a biochemical characterization of the mutant polymerases to see if there is a fidelity defect. We characterized misincorporation, mispaired base extension, and mispaired base removal and found defects in one or more of these processes in each of the mutants. These mutants provide an opportunity to elucidate the cellular consequences of error prone transcription. The work described here is a step to understanding the biological role of faithful information transfer from DNA to RNA.

| Plasmids: | | 
| --- | --- | 
| pJX220 | RPB1 | URA3 2-micron circle plasmid |
| pJS670 | RPB1 | LEU2 CEN plasmid |
| pJS725 | RPB1 | URA3 2-micron circle plasmid |
| pJS757 | RPB1 | LEU2 CEN plasmid |
| pJS871 | rpb1-E1103G | LEU2 CEN plasmid |
| pJS932 | rpb1-E1230K (pap21-24) | LEU2 CEN plasmid |
| pJS1006 | rpb1-A1076T | LEU2 CEN plasmid |
| pJS1007 | rpb1-G823S | LEU2 CEN plasmid |
| pJS1008 | rpb1-S1229F | LEU2 CEN plasmid |
| pJS1009 | rpb1-A1076V | LEU2 CEN plasmid |
| pJS1010 | rpb1-M1079I, T1548I | LEU2 CEN plasmid |
| pJS1025 | rpb1-K1132E | LEU2 CEN plasmid |
| pJS1126 | rpb1-T1141I | LEU2 CEN plasmid |
| pJS1212 | rpb1-T1113P | LEU2 CEN plasmid |

| Yeast Strains: | | 
| --- | --- | 
| GRY2852 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), met15-\(\Delta\), ura3-\(\Delta\), bud5::PGAL1-cre-WT, rpb1::natMX, [pJDI220 (RPB1 URA3)]. |
| GRY2855 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), met15-\(\Delta\), ura3-\(\Delta\), bud5::PGAL1-cre-Y324C, rpb1::natMX, [pJDI220 (RPB1, URA3)]. |
| GRY3175 | MAT\(^a\), his3, leu2-\(\Delta\), ura3, his3, lys2-\(\Delta\), trp1-361, pep4::HIS3, prb1-\(\Delta\)1.6R |
| | rpb1::kanMX, RPB3::6XHis, [pJS725 (RPB1, URA3)]. |
| GRY3258 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), ura3-\(\Delta\), rpb1::natMX, P\(_{PGAL1}\)::cre-Y324C, [pJDI220 (RPB1, URA3)]. |
| GRY3284 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), lys2-\(\Delta\), ura3-\(\Delta\), rpb1::natMX, P\(_{PGAL1}\)::cre-Y324C, [pJS781 (rpb1-E1103G, LEU2)]. |
| GRY3739 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), met15-\(\Delta\), ura3-\(\Delta\), bud5::PGAL1-cre-Y324C, rpb1::natMX, [pJS670 (LEU2, RPB1)]. |
| GRY3740 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), met15-\(\Delta\), ura3-\(\Delta\), bud5::PGAL1-cre-Y324C, rpb1::natMX, [pJS670 (LEU2, RPB1)]. |
| GRY3742 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), met15-\(\Delta\), ura3-\(\Delta\), trp1, bud5::PGAL1-cre-Y324C, rpb9::kanMX |
| GRY3743 | MAT\(^a\), his3-AI2floxINV (hygMX, zeoMX), leu2-\(\Delta\), met15-\(\Delta\), ura3-\(\Delta\), trp1, bud5::PGAL1-cre-Y324C, rpb9::kanMX |

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Methods

Plasmids

See Table 1. pJS757 is a LEU2-based CEN vector carrying the RPB1 ORF, as well as 594 bases upstream and 501 bases downstream of the ORF [16]. pJS725 and pJDI220 are UR3A-based 2-micron circle plasmids with the same RPB1 insertion as pJS757. RPB1 was randomly mutagenized by passing a pool of plasmid pJS757 twice through the mutator (mutS mutD mutT) E. coli strain XL-1 Red (Stratagene).

Yeast strains

The yeast strains are related to the BY4741 and BY4742 [68] See Table 1.

Cre expression

The cre gene used in these experiments has a seven amino acid nuclear localization site from the large SV40 T-antigen added at the N-terminus [69]. The P_{GAL1}-cre gene was made by fusing the NLS-cre ORF to the P_{GAL1} promoter inserted into the BUD5 gene at an autochthonous EcoRI site. The P_{HIS3}-cre gene was made by overlap PCR and integrated at the normal position for HO. The insertions were made by selecting for replacement of URA3, previously inserted at those locations.

Reporters for Cre activity

Direct repeat of loxP: The his3-A12floXkanMX reporter (Figure 1B, construct (ii)) has an artificial intron inserted into HIS3 at an MspI site in codon 70. The 1710 base insertion includes two loxP sites in direct orientation flanking the kanMX cassette. kanMX is in the same orientation as HIS3 so that the terminator blocks transcription through the rest of HIS3 resulting in cells that are His-.

The Cre recombinase can remove the 1502 bases including kanMX plus one loxP resulting in a spliceable intron (HIS3-A12lox Figure 1B, (ii)) and cells that are His+. The ade6-A12floXkanMX reporter has the same artificial intron inserted after the tenth codon of ADE6 causing cells to be ade6-.

The Cre recombinase can remove kanMX and make a functional ADE6-A12lox gene. Inverted repeat of loxP: The his3-A12floXN reporter, and Ashley Denney for the rpb1-T1113P allele.

Specifically, cells from 3–4 ml of stationary (2–3 days) culture were pelleted, washed once with cold water and resuspended in lysis buffer (150 mM Tris- acetate, pH 7.9, 50 mM potassium acetate, 5 mM MgCl2, 10 μM ZnCl2, 2 mM 2-mercaptoethanol, 0.5 mM EDTA and protease inhibitors). The cells were disrupted using a Precellys homogenizer according to manufacturer’s instructions, the lysate was removed from the glass beads, and KCl was added to 1M final concentration. The debris was precipitated for 15 min at 14,000 rpm in Eppendorf table-top microcentrifuge at 4°C. The supernatant was added to 50 μl of Ni-NTA agarose pre-washed with transcription buffer (TB) containing 20 mM Tris-HCl, pH 7.9, 5 mM MgCl2, 10 μM ZnCl2, 2 mM 2-mercaptoethanol and 1000 mM KCl (TB1000). Pol II was immobilized on the beads for 30–50 min at 4°C, and the beads were extensively washed with TB1000 and TB40 (TB with 40 mM KCl). The elongation complex was assembled and purified with the immobilized Pol II exactly as described [54] using the following oligonucleotides: Cre RNA C9: 5’ GUC AUG AAC 3’ (phosphorylated with T4 polynucleotide kinase in the presence of γ-P32-labeled ATP); NDS-Cre-Cys ([Non Transcribed DNA Strand]) 5’GGCTGGACATGTAATTGTGCATGAC GGTATCCTGAA CCGTGATAGTGAAGA 3’; and TDS-Cre-Cys ([Transcribed DNA Strand] which is an exact complement of NDS (Cre-Cys). All NTPs used for walking and misincorporation assays were additionally purified [19]. The elongation complex containing 10-nm RNA (U10) was obtained by 5 min incubation of the assembled and purified complex with 5 μM UTP and washed 4 times with 1 ml of TB40. For the TH1S sensitivity assay, mismatched A11 elongation complex was obtained by 5 min incubation of U10 with 1 mM ATP purified by 4 washes with TB40, eluted from the beads with 100 mM imidazole in the presence of 0.2 mg/ml acetylated BSA, diluted 10-fold with TB40 to decrease imidazole concentration and concentrated using Microcon AmiconUltra (Millipore) concentrator to 30 μl. Reactions were stopped by addition of equal volume of gel loading buffer containing 10 M urea and 50 mM EDTA. The reaction products were resolved in 19:1) and processed at 50 W constant power. The shorter 20 cm was within 3 cm of the bottom of the gel. For other experiments the 20×20 cm gels were used and electrophoresis has been performed in 1× TBE at constant power of 65 W until the bromphenol blue tracing dye was within 3 cm of the bottom of the gel. For other experiments the 20×20 cm gels were used and electrophoresis was performed at 50 W constant power.

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Author Contributions

Conceived and designed the experiments: JNS JDI MLK MK. Performed the experiments: JNS JDI BKS DRG IH MLK. Analyzed the data: JNS JDI MLK MK IH DRG. Contributed reagents/materials/analysis tools: JNS JDI MLK MK. Wrote the paper: JNS JDI MLK MK.

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