External conditions inversely change the RNA polymerase II elongation rate and density in yeast

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Abstract

Elongation speed is a key parameter in RNA polymerase II (RNA pol II) activity. It affects the transcription rate, while it is conditioned by the physicochemical environment it works in at the same time. For instance, it is well-known that temperature affects the biochemical reactions rates. Therefore in free-living organisms that are able to grow at various environmental temperatures, such as the yeast Saccharomyces cerevisiae, evolution should have not only shaped the structural and functional properties of this key enzyme, but should have also provided mechanisms and pathways to adapt its activity to the optimal performance required. We studied the changes in RNA pol II elongation speed caused by alternations in growth temperature in yeast to find that they strictly follow the Arrhenius equation, and that they also provoke an almost inverse proportional change in RNA pol II density within the optimal growth temperature range (26–37 °C). Moreover, we discovered that yeast cells control the transcription initiation rate by changing the total amount of available RNA pol II.

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1. Introduction

Living cells have evolved within the constraints imposed by physical and chemical laws. This rule acts on every chemical reaction within a cell and has limited the concentrations, structures, affinities and specific locations of all enzymes and metabolites within ranges that are compatible with life and evolution. For instance, it has been recently demonstrated that metabolic enzyme thermostability is rate-limiting at superoptimal temperatures in Escherichia coli [1].

Chemical reactions are governed by the thermodynamics and kinetics laws that depend on concentrations of reactants and products, and also on environmental factors, such as pH or temperature. In the specific case of nucleic acid synthesis, reactants (nucleotide triphosphates, NTPs/dNTPs) are usually well above K\(_\text{m}\) [2], and the main product of the reaction is a long polymer to which the catalytic enzyme (polymerase) is continuously bound during its synthesis (processive synthesis). The use of highly processive enzymes is necessary because a high synthesis rate must avoid the delay that a distributive mechanism for monomer incorporation would cause [3].

Syntesis of mRNAs is usually the most important step in gene regulation in eukaryotes, and RNA polymerase II (RNA pol II) is the actual enzyme in this process. RNA pol II molecules are first recruited to gene promoters to then transcribe along the coding region of the gene (see [4] for a review). RNA pol II is a complex of 12 subunits that is helped by many other ancillary proteins during initiation and elongation [4]. During elongation, RNA pol II undergoes different modifications which mark their particular stage throughout the process [4]. All eukaryotic RNA pol II have a long C-terminal tail (CTD) in their largest subunit (Rpb1) with repeated heptads (from 26 in yeast to 52 in humans); e.g., Tyr-Ser-Pro-Thr-Ser-Pro-Ser. The CTD is a target for many reversible modifications, including phosphorylation, prolyl-isomerization and glycosylation, at various positions [4]. Unphosphorylated molecules form the PIC (pre-initiation complex) at gene promoters. Then Ser5 phosphorylation in the CTD is required to start early elongation. Ser2 phosphorylation starts later and progressively replaces Ser5 during elongation, thus rendering it a marker for elongating RNA pol II (see [4,3] for a review). Moreover, it should also be considered that differences in transcription depend not only on RNA pol II molecules, but also on the ancillary factors accompanying them along the gene.

Genes differ not only in the particular amino acid sequence they encode, but also in other sequence-dependent features. Genes have different lengths, G + C content, dinucleotide distribution, exon/intron
organization in general, and a chromatin structure in particular. All these features, which are intrinsic to the transcriptional apparatus or to the template that RNA pol reads, can influence elongation [3]. As explained before however, RNA pol II elongation can also be conditioned by extrinsic factors, such as temperature, NTP concentration, and pH.

The speed at which processive RNA pol II elongates has been found to be 1–2 kb/min on average in several eukaryotes [5–7]. However, it has been discussed that it varies within a broader range (1–6 kb/min, see [8]) and has been recently shown to be variable among and along human genes [9], thus raising the important question of the role that the elongation rate plays in regulating the TR. The elongation rate of a prokaryotic RNA pol has been kinetically modeled on the basis of the single one-nucleotide successive steps as a thermal ratchet molecular motor that depends on NTP concentrations and DNA sequences [10]. In prokaryotes, in vitro experiments have demonstrated that RNA pol is dependent on temperature [11].

The nascent transcription rate (nTR) depends on the density of RNA pol II molecules and their speed. Density directly depends on the rates of initiation events and abortive elongation that cause RNA pol II to drop off [4]. When the remaining factors are fixed, density and the elongation rate are mutually inverse parameters. Thus, if an external factor that affects the elongation rate is changed, it will be observed as a reciprocal inverse change in RNA pol density. Current techniques to determine the nTR, such as chromatin immunoprecipitation of RNA pol II molecules or run-on analysis, are based on the determination of RNA pol II densities [12]. When doing comparative analyses between samples, presumably elongation rates do not vary [13–15]. In some cases they have been assumed to change with temperature [16]. However, this assumption has not been experimentally tested and it is not known if cells react in any way to a change in the RNA pol II elongation rate when it happens.

In this study, we experimentally show that in vivo RNA pol II in yeast elongates at different speeds depending on external factors, such as temperature or NTP concentration. The change in RNA pol II density is mainly reciprocal to that in speed, and the TR does not alter. Nonetheless, we also observed minor changes in the TR within the optimal yeast growth temperature range (26–37 °C), which are brought about by an alteration in the initiation rate, mainly in response to changes in the total cell amount of RNA pol II.

2. Materials & methods

2.1. Yeast strains, media and growth conditions

For most experiments, we used the Saccharomyces cerevisiae THJ2400 strain (MATα, pGAL1–YLR454W::URA3, ural3A, genetic background S288C), which was grown in liquid YPD medium (1% yeast extract, 2% peptone, 2% glucose), YPGal (1% yeast extract, 2% peptone, 2% galactose) or synthetic complete media lacking uracil and pH.

For cell growth rate determination, cells were grown in 250 mL flasks in 100 mL of liquid YPD medium at the corresponding temperatures from O.D.₆₀₀ = 0.1 to the mid-late log phase. The growth rate was calculated as the inverse of the slope of the line of best fit obtained when representing the log₂ value of the measured O.D. against time.

The growth rate of each relevant segment of the reference (ORF-Free Chr 5) was extrapolated to determine the RNA pol II elongation rate when it happens.

2.2. RNA pol II elongation rate assays

The procedure to determine the RNA pol II elongation rate was adapted from Mason and Struhl [17]. Cells were grown in the specified galactose media until O.D.₆₀₀ = 0.5 at the corresponding temperature. Upon glucose addition, 50 mL of culture aliquots were harvested at the required time points. After collection, cells were treated with 1% formaldehyde for 15 min at room temperature. Then cells were resuspended in lysis buffer (50 mM HEPES-KOH pH 7.5, 140 mM NaCl, 1 mM EDTA, 0.1% sodium deoxycolate, 1 mM PMFS, 0.15% benzamidine, 1 tablet each of 50 mL of Protease inhibitor cocktail tablets, Roche) and were broken with glass beads by Fastprep (MP Biomedicals). The lysate was sonicated in a Bioruptor device to yield DNA fragments with an average size of 350 bp. Immunoprecipitation was performed by incubating chromatin with an antibody (8WG16, Covance) against the C-terminal domain of the RPB1 subunit of the RNA pol II in magnetic beads (Dynabeads Pan Mouse IgG, Invitrogen) for 90 min with rotation. The immunoprecipitated material was eluted from the beads by incubating twice at 65 °C for 10 min in elution buffer (50 mM Tris–HCl pH 8, 10 mM EDTA, 1% SDS). To revert the crosslinking, samples were incubated in elution buffer for 15 h at 65 °C by agitation and for 90 min at 37 °C with 0.05% Proteinase K (Roche). DNA was obtained by phenol–chloroform extraction. A quantitative analysis DNA was done by qPCR using the oligonucleotides listed in Table 1. Background noise and immunoprecipitation homogeneity were tracked by a reference DNA fragment contained in the non-coding region of Chromosome V (ORF-Free Chr 5). Relative RNA pol II occupancy was determined by dividing the immunoprecipitation efficiency of each relevant segment of the gene by the immunoprecipitation efficiency of the reference (ORF-Free Chr 5). The relative occupancy values are expressed as the ratio of the RNA pol II occupation upon glucose treatment divided by occupation in galactose. Having obtained the kinetics of the RNA pol II dissociation on each gene fragment, we calculated the linear tendencies and extrapolated the line to the X-axis to obtain the time (min) at which the last wave of RNA pol II abandoned that specific segment. The RNA pol speed was calculated as the position (kb) divided by the obtained time (min) value (see Supplementary Fig. S1).

2.3. In vitro transcription

For the average intensity normalization in GRO experiments, we performed an in vitro transcription of the PCR fragments amplified

<table>
<thead>
<tr>
<th>Primer sequences for the quantitative PCR analysis.</th>
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<tr>
<td>GAL1-YLR454W</td>
</tr>
<tr>
<td>+3/+46</td>
</tr>
<tr>
<td>GY0-A: 5′-GATGTTTCCGATTAAGTCTCTGACAA-3′</td>
</tr>
<tr>
<td>GY0-B: 5′-GCCGTTAAAGCTACAGTTCGACAA-3′</td>
</tr>
<tr>
<td>GY0-C: 5′-GCTCCATAAGAAAGTCACTGCAAA-3′</td>
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<tr>
<td>GY0-D: 5′-CAGGGCTTTTGGGTTTTTCA-3′</td>
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<tr>
<td>GY1: 5′-CCACCATCAAGCAACAGAC-3′</td>
</tr>
<tr>
<td>GY2: 5′-GCCAAAATAAGACGTGATTCGTTTGA-3′</td>
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<td>GY4: 5′-GTGGGACATTTAAACGGCG-3′</td>
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<tr>
<td>ORF-Free Chr 5</td>
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<tr>
<td>Positions 9754/9837 (intergenic) of chromosome V</td>
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<td>CHBS-A: 5′-GTCCCTTIAAACAGGCTACAGTGAT-3′</td>
</tr>
<tr>
<td>CHBS-B: 5′-GTGGGACATTTAAACGGCG-3′</td>
</tr>
<tr>
<td><strong>Table 1</strong></td>
</tr>
<tr>
<td><strong>GAL1-YLR454W</strong></td>
</tr>
</tbody>
</table>
from the thr gene of Bacillus subtilis using T7 RNA polymerase (Ambion) and [α-33P] UTP (Perkin Elmer).

2.4. Genomic run-on (GRO) assays

The GRO protocol was adapted from [13]. In all cases, cells were grown to O.D.600 ~ −0.5. Radioactively-labeled RNA was hybridized to macroarrays containing probes for the 769 ORFs of the S. cerevisiae genome (see Suppl. Table S1) and the 32 internal controls of the thr B. subtilis gene (ATCC clone # 87484) at random positions, produced in the DNA chips service S.C.S.I.E. (University of Valencia). The same amount of in vitro transcribed B. subtilis clones was added to each GRO sample. Macroarrays were exposed to an Imaging plate (BAS-MP, Fujifilm) and were scanned in a Fuji FLA3000 Phosphorimager. The signal intensity of the spots was quantified with the Array Vision™ image analysis software. sARM values (background-corrected Artifact Removed Median density) were used for the analysis and were normalized by the average intensity value of the internal controls to avoid technical variation. The RNA pol II total density value, under all the tested conditions, was calculated as the median of the intensities of all the genes analyzed.

2.5. Western blot

Cells were grown in synthetic medium to O.D.600 ~0.5 at their respective temperatures in biological triplicates and were counted in a Coulter-counter device. To prepare protein extracts, cultures were harvested by centrifugation and resuspended in lysis buffer (50 mM Tris–Base, 150 mM NaCl, 5 mM MgCl2, 1% NP40, pH = 7.5) containing phosphatase inhibitors (1 mM Na-orthovanadate, 60 mM β-glycerol phosphate, 50 mM NaF) and Complete Mini Protease Inhibitor Cocktail Tablets (Roche). Cells were broken with glass beads by FastPrep. Protein concentration was determined by the Bradford Assay (Bio-Rad). Then 1 μg of total protein was redissolved on a 7% SDS-PAGE gel (7% acrylamide, 0.1% Bis-Acrylamide, 0.2% SDS, 0.75% M Tris–Base) and was transferred onto a nitrocellulose membrane (GE Healthcare) in transfer buffer (25 mM Tris–Base, 192 mM Glycine, 0.1% SDS, 20% Methanol). The total RNA polymerase II protein expression was detected using the 8WG16 antibody, while phosphorylated RNA polymerase II was detected using the anti-RNA polymerase II CTD repeat YSPTSPS (phospho S2) antibody [HSJ] (Abcam 24758). This antibody, however, has been described to recognize both Ser2 and Ser5 phosphorylations [18]. Therefore, we assume that it quantifies all forms of elongating RNA pol II. The Pkg1 protein expression was detected as an endogenous control using the anti-Pkg1 antibody (Invitrogen). The anti-mouse HRP-conjugated secondary antibody was purchased from Promega. Immunoblots were developed using the ECL Prime kit (GE Healthcare). Band intensities were analyzed with the Image Quant TL program (GE Healthcare).

3. Results

3.1. The RNA pol II elongation rate varies with the growth temperature

Enzymatic reactions increase velocity with temperature according to the Arrhenius dependence of the catalytic constant [19]. For RNA synthesis, this dependence affects the individual catalytic steps of nucleotide incorporation, but it should not necessarily be converted into a quantitative identical effect on the elongation rate. To test if this was actually the case for yeast RNA pol II, we used the assay described by Mason and Struhl [17]. In this assay, the last elongation wave of RNA pol II after the shut-off of the GALI promoter was followed by qPCR over time using different primer sets along the 8 kb-long YLR454W gene to determine by extrapolation the time that RNA pol molecules require to elongate from the promoter to the primer pair (see Fig. S1). Although, in principle, only one primer pair was necessary to measure the elongation rate, we used four pairs placed at 2, 4, 6 and 8 kb from the TSS. This allowed us to obtain a more robust result by averaging the RNA pol speed calculated from several independent reference points.

For the experiment that determines how temperature influences the elongation rate, we utilized two different growth conditions: rich and synthetic media. The results for three independent experiments were repetitive and similar in both growth media. As expected, Fig. 1A depicts that the RNA pol II speed increased linearly with growth temperature from 0.67 to 1.31 kb/min between 23 °C and 37 °C. This increase (about 0.5 kb/min every 10 °C) perfectly fitted the Arrhenius equation (Fig. 1B), which suggests that it is mostly the result of the effect of temperature on overcoming a fixed activation energy step.

3.2. RNA pol II density inversely correlates with RNA pol II speed

A change in the elongation rate should be concomitantly seen as a change in RNA pol II density (i.e., not altering the TR) if the rate of initiation and the drop-off index are constant. Therefore, we decided to check whether these premises are fulfilled exactly in yeast. A plot of the time zero ChIP values from the YLR454W experiment vs. temperature shows that RNA pol II density decreased with temperature (Fig. 2A). Since the particular fusion gene used can have specific effects on transcription initiation (i.e., a temperature effect on the GALI promoter [20]), we decided to use the genome-wide techniques to avoid any bias in the density calculations.

To do this, we performed a genomic run-on experiment [13] to measure the total elongating RNA pol II density in the ORF-containing genes. This approach offers the advantage of avoiding having to count RNA pol
and RNA pol III contribution, which account for ~70% of yeast cell transcription [21]. As run-on counts all the elongating RNA pol molecule in the regions covered by probes, and since our macroarrays contain whole-length ORF sequences for almost 800 genes of the S. cerevisiae genome, the sum of all their signals is a good estimator of the total RNA pol II average density in canonical genes.

Fig. 2B and Table 2 illustrate how the median run-on signal drastically dropped from 23 °C to 37 °C over the set of canonical RNA pol II genes. This result was then confirmed by evaluating the amount of elongating RNA pol II by Western blot. The quantification of phosphorylated polymerase using the H5 monoclonal antibody confirmed a severe decrease of elongating RNA pol II from 23 °C to 37 °C (Fig. 2C & D, Table 2). It is interesting to note that the drop in density did not exactly compensate the increase in speed with temperature, which caused a bell-shaped TR curve (Fig. 3). As the growth rate can impose different constraints to yeast physiology, including variations in the TR, we checked the exact growth rates for our particular S. cerevisiae strain at several common laboratory temperatures ranging from 18 °C to 37 °C. Fig. S2 shows that the growth rate did not significantly differ within the 26–37 °C range, whereas it was 70% slower at 23 °C than it was at 30 °C. These results are similar to those obtained by other authors for the BY4741 yeast strain [22]. In that case however, the growth rate at 37 °C was slightly better than it was at 30 °C. The behavior of the TR at temperatures under 26 °C can be influenced by major changes in growth rates, which precludes a quantitative comparison. Finally, although the observed growth rates were similar between 30 °C and 37 °C, the TR slightly lowered (by around 25%) at 37 °C (Fig. 3).

To further verify the inverse relationship of RNA pol II speed and density, we decided to investigate another cause of RNA pol II speed reduction: the effect of the [NTP] depletion caused by the 6-azauracil (6AU) I and RNA pol III contribution, which account for ~70% of yeast cell transcription [21]. As run-on counts all the elongating RNA pol molecule in the regions covered by probes, and since our macroarrays contain whole-length ORF sequences for almost 800 genes of the S. cerevisiae genome, the sum of all their signals is a good estimator of the total RNA pol II average density in canonical genes.

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### Table 2

<table>
<thead>
<tr>
<th>Temperature</th>
<th>SerP</th>
<th>SD</th>
<th>Run-on</th>
<th>SD</th>
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<td>0.12</td>
<td>1.42</td>
<td>0.03</td>
<td>1.44</td>
</tr>
<tr>
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<td>0.00</td>
<td>1.00</td>
<td>0.00</td>
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</tr>
<tr>
<td>34 °C</td>
<td>0.98</td>
<td>0.16</td>
<td>0.82</td>
<td>0.19</td>
<td>0.90</td>
</tr>
<tr>
<td>37 °C</td>
<td>0.60</td>
<td>0.10</td>
<td>0.49</td>
<td>0.06</td>
<td>0.55</td>
</tr>
</tbody>
</table>

Fig. 3. The RNA pol II transcription rate according to growth temperature. The elongation rate (Fig. 1A) and density (Table 2 average) data were multiplied to obtain a relative TR at three different growth temperatures. SD bars are shown.
In our case, we noted that the results were similar to those previously published [17]. As seen in Fig. 4A, we found that 50 μg/mL of 6AU led to a reduction in RNA pol II speed in a wt strain after a 30-minute treatment, and this reduction was estimated to be about 60%. Next we determined by run-on the RNA pol II density in the cells in which a decrease in the NTP pool caused the elongation rate to lower (Fig. 4B). The experiment was done in an \textit{imd2} mutant strain to avoid the partial recovery of NTP pools, which occurs in a wild-type strain due to the induction of the \textit{IMD2} gene during nucleotide depletion [23,24]. We used the same 6AU concentration as in the experiment of Fig. 4A. This concentration allows \textit{imd2} strain growth, be it at a lower rate than in its absence, and it provokes constant [NTP] depletion. Fig. 4B depicts how density increased by 40% after 30 min in 6AU as compared to time 0 in an \textit{imd2} mutant. Taken together, the fact that the addition of 6AU lowers the RNA pol II elongation rate (Fig. 4A) and increases Pol II density (Fig. 4B) supports the idea that the RNA pol II elongation rate is reflected by its density. This means that, given the possible variations in RNA pol II speed, the TR cannot always be compared between two samples if it has been estimated directly from RNA pol density values.

Another factor that might influence RNA pol II density is the drop-off, which might vary with temperature and cause differences in the density observed. We checked this hypothesis by looking at the distribution of the RNA pol II molecules along the \textit{YLR454W} gene for each growth temperature. Figs. 5 and S3 reveal that the distribution along

**Fig. 4.** The RNA pol II elongation rate and density in the presence of 6AU in wt and \textit{imd2} cells. A) The RNA pol II elongation rate was measured using the \textit{GAL-YLR454W} construction [17] in a wt strain before (−6AU, left panel) and after (+6AU, right panel) the addition of the nucleotide-depleting drug 6AU (50 μg/mL). Values are normalized to the time zero value for each probe. Decay curves were obtained from the cells growing in SC–URA at 30 °C after changing from galactose to glucose medium. Elongation rates (1 kb/min and 0.4 kb/min, respectively) were calculated from the decay curves as distance/time, as described in Fig. S1, and were averaged for all the probes to obtain a final speed value for each growth condition. B) Histogram showing the total RNA pol II density calculated by run-on (GRO assay) in an \textit{imd2 Δ} strain growing at 30 °C in the absence of (left), or 30 min after the addition of, 50 μg/mL 6AU (right). The SD bars from three replicates are shown.

**Fig. 5.** Analysis of the RNA pol II distribution on the \textit{YLR454W} gene according to growth temperature. The same experimental data as in Fig. 1A were used to obtain the RNA pol II relative distribution along the \textit{YLR454W} gene (probes 2, 4, 6 and 8 kb; see Fig. S1) by taking the values of 30 °C as 1. Fig. S3 is a representation of these data without the normalization by the 30 °C values.
the coding region, from 2 to 8 kb probes, decreased from 5′ to 3′ for all three temperatures, which was expected given previously published results [17]. This decrease, however, did not reveal a significantly different distribution for 23 °C or 37 °C if compared to 30 °C.

As mentioned before, density directly depends on the rates of initiation events and abortive elongation that cause RNA pol II to drop-off [4]. Since we have demonstrated that drop-off is not variable in our particular case, the 25% drop in the TR observed at higher temperatures (Fig. 3) suggests that the cause is a decrease in transcription initiation events. This led us to investigate if this reduction is caused by a drop in the total amount of RNA pol II or only in the proportion of that involved in elongation.

3.3. RNA pol II amount varies with growth temperature

To further investigate the cause of the decreased RNA pol II density at high temperatures, we measured by Western blot the RNA pol II amount per cell in the cells grown at different temperatures using an antibody against total RNA pol II. Fig. 2C and D shows that the amount of total RNA pol II/cell lowered from 30 °C to 37 °C, similarly to elongating RNA pol II density. It is interesting to note that this reduction was specific for RNA pol II because total protein/cell did not reduce (Fig. 2D). Conversely at 23 °C, the amount of RNA pol II was lower than at 30 °C. As we have previously shown, both run-on and H5 antibody data, which quantifies only actively elongating polymerase, indicate that the amount of elongating RNA pol II continuously diminished with an increase in temperature (Fig. 2B–D). Therefore, yeast cells seem to adjust the total RNA pol II amounts per cell to accomplish an appropriate initiation rate on gene promoters within the range of the similar growth rates tested in these experiments (30–37 °C). At lower temperatures and growth rates (23 °C, Fig. 52), the level of RNA pol II/cell did not increase further. In this case, it seems that the proportion between elongating and total RNA pol II molecules increases.

4. Discussion

The TR is a fundamental parameter of every living cell because the synthesis of all its biomolecules relies directly or indirectly on the expression of its genetic information. The rate at which genes are transcribed is the first, and probably one of the most important, regulated steps along the flux of genetic information. RNA polymerases are, therefore, highly regulated machines in both prokaryotes and eukaryotes.

In recent years, several different techniques, mostly genome-wide, have been developed to measure the TR in both yeast and higher eukaryotes (revised in [12]). Many calculate RNA pol II density by analyzing either the DNA sequence where it is placed onto (chromatin immunoprecipitation, ChIP, i.e., [25,26]) or the nascent RNA sequence being either the DNA sequence where it is placed onto (chromatin immunoprecipitation, ChIP, i.e., [25,26]) or the nascent RNA sequence being transcribed (genomic run-on, GRO; i.e., [13,15,27]). In this kind of studies, the TR is decreased with an increase in temperature (Fig. 2B–D). Therefore, yeast cells seem to adjust the total RNA pol II amounts per cell to accomplish an appropriate initiation rate on gene promoters within the range of the similar growth rates tested in these experiments (30–37 °C). At lower temperatures and growth rates (23 °C, Fig. 52), the level of RNA pol II/cell did not increase further. In this case, it seems that the proportion between elongating and total RNA pol II molecules increases.

In this study, we used two external factors that should presumptively affect the elongation rate and demonstrate that RNA pol II is affected by them at exactly the magnitude predicted by the Arrhenius equation (temperature) or within expectations (NTP depletion). Published RNA pol elongation models [10,38] have predicted its kinetic behavior based on a general ratchet model for processive enzymes. Each individual step depends on thermal fluctuations and [NTP]. In fact in single molecule studies, RNA pol from E. coli has been reported to increase the elongation rate in vitro with a temperature fitting the Arrhenius equation, including pauses, between 21 °C and 37 °C [11,29]. From those results, it can be anticipated that the RNA pol II elongation rate will increase with temperature. However, this had not been demonstrated in eukaryotic cells in vivo to date, that is, not until this study.

A yeast cell growing under optimal conditions (i.e., 30 °C in YPD) probably performs transcription at its best. If temperature changes, but the growth rate does not alter (i.e., at 37 °C), at first sight, one might expect the TR to remain constant. This can be achieved by keeping the initiation rate constant. In this way, the average number of elongating RNA pol IIs will decrease in exactly the same proportion in which their elongation rate increases because they spend less time in the elongation phase.

To check this hypothesis, we used a genomic run-on [13] to measure the total RNA pol II density in ORF-containing genes. We discovered that reduced RNA pol II density compensated the increased elongation rate. However the drop in density was slightly higher (25%) than required at 37 °C, which means that the initiation rate lowered (Fig. 3). Since the elongation rate is a feature of every single RNA pol. Moreover, it has been shown to be variable along and among genes in human cells [9]. However, it can be averaged for the whole length of the “gene highway” that it travels along with single molecule techniques (see refs. [6,7]) or, more commonly, for the whole set of RNA pol molecules that transcribe many copies of the same gene in a cell population over a given time lapse [32–34].

In many instances, RNA pol II density has been interpreted as a measure of the nTR by assuming an average constant elongation rate. In order to be rigorous, this assumption needs the experimental verification of absence of variation in the elongation rate or in the drop-off rate between the compared samples. In fact, the changes in RNA pol II speed have been calculated in vivo or in vitro in very few cases [17,35]. Most of the studies conducted in yeast have been done at 28–30 °C. In a previous work [16], we used a correction of the elongation rate in the TR based on the Arrhenius equation during a heat-shock response at 37 °C; however, it was not experimentally verified. Here we confirm that our prediction was correct, and we show that the current protocols used to evaluate the TR are all based on the determination of RNA pol densities [12] and require an independent estimation of the existence, or not, of the external factors that induce changes in the elongation rate. In E. coli, it has been shown that RNA pol elongates at a maximum speed of 80 to 90 nucleotides (nt) per s in mRNA operons [36] and at a speed of 45 nt/s for mRNA operons at 37 °C [37]. The speed of RNA pol II in eukaryotes is believed to be slower, 25 nt/s on average, but varies considerably (13–100 nt/s), which could be either a real biological property or caused by the experimental method to determine it [5]. In yeast growing at 28–30 °C, measuring the last wave of RNA pol II after blocking the transcription [17] determined an average of 18–25 nt/s (1–1.5 kb/min), which is similar to that established by single molecule live tracking techniques [6,7].
expected [see the Discussion in [30,39,40]]. This conclusion is based on the comparison of genomic run-on data with single-gene data for elongation rate and drop-off. Accordingly, we cannot exclude that specific subsets of genes control their TR response to temperature at post-initiation steps. In this regard, we have previously shown that some yeast regutors, like those encompassing ribosome-related and mitochondrial-related genes, can be regulated at the level of RNA pol II activity during elongation [41,42].

What is the mechanism that a eukaryotic cell uses to globally control the initiation rate? The objective to increase/decrease the number of initiating RNA pol II molecules can be accomplished by changing the total amount of RNA pol II in the cell or by acting in any (or several) successive steps from the formation of closed PIC to the productive elongation by phosphorylated RNA pol II [3,4]. In E. coli cells, the RNA polymerase concentration increases with a higher growth rate [31] and the percentage of transcription-engaged RNA pol molecules also rises [37]. Therefore, this prokaryote controls the TR, as far as its growth requirements are concerned, by increasing both the total RNA pol available and the fraction of it devoted to transcribe, be it with differences between the rRNA and mRNA operons [31]. In this case, the increase in the growth rate is studied at a constant temperature, but the elongation rate in in vitro studies increases with temperature by the same relative amounts as the general growth rates [11]. Here we demonstrate in yeast that the increase in the growth rate from 23 °C to 30 °C involved an increment in the total amount (Figs. 2C & D), but not in the in vivo elongating RNA pol II (Fig. 2B–D & Table 2). Furthermore, from 30 °C to 37 °C, we observed that the total RNA pol II amount diminished per cell (Fig. 2D monodisks) and, probably as a result of this, the phosphorylated RNA pol II lowered (Fig. 2D squares). This reduction in elongating pol II may correspond to the principle of reducing the number of costly machineries, which become faster due to the additional energy supply at higher temperatures, as previously seen with ribosomes [43,44].

5. Conclusions

We have found an in vivo dependence of the RNA pol II elongation rate and density on transcribed genes with the growth temperature in yeast cells. The variation of elongation rate and density with temperature is reciprocal, as expected [30]. Accordingly, RNA pol II density cannot be directly used to compare TRs without verifying constant elongation rates. Our results also imply that global variations in the TR seem to be mainly due to the control at the transcription initiation level. Within the usual growth temperature range in S. cerevisiae, initiation of transcription is, in turn, controlled by adjusting the total amount of RNA pol II molecules present in the cell.

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