Quantitative imaging of transcription in living Drosophila embryos reveals the impact of core promoter motifs on promoter state dynamics

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Genes are expressed in stochastic transcriptional bursts linked to alternating active and inactive promoter states. A major challenge in transcription is understanding how promoter composition dictates bursting, particularly in multicellular organisms. We investigate two key Drosophila developmental promoter motifs, the TATA box (TATA) and the Initiator (INR). Using live imaging in Drosophila embryos and new computational methods, we demonstrate that bursting occurs on multiple timescales ranging from seconds to minutes. TATA-containing promoters and INR-containing promoters exhibit distinct dynamics, with one or two separate rate-limiting steps respectively. A TATA box is associated with long active states, high rates of polymerase initiation, and short-lived, infrequent inactive states. In contrast, the INR motif leads to two inactive states, one of which relates to promoter-proximal polymerase pausing. Surprisingly, the model suggests pausing is not obligatory, but occurs stochastically for a subset of polymerases. Overall, our results provide a rationale for promoter switching during zygotic genome activation.
In all eukaryotes, transcription of active genes into RNA requires the controlled assembly of multiple protein complexes at promoters. This includes the sequential recruitment of general transcription factors to form the pre-initiation complex (PIC), followed by the recruitment of RNA Polymerase II (Pol II) at the transcriptional start site (TSS) of promoters. Among all these complexes, TFIIID stands out as the primary core promoter recognition factor that triggers PIC assembly. TFIIID includes TATA-binding protein (TBP), which binds upstream of the TSS, as well as 13 TBP-associated factors (TAFs), known to bind downstream promoter elements such as the Initiator element (INR) and the downstream promoter element (DPE) motifs.

After initiation, Pol II transcribes a short stretch of 30–80 nucleotides before pausing. This step is regulated by the TFIIID complex, as directly demonstrated in vitro and inferred from the over-representation of TFIIID-bound core promoter motifs (INR, DPE, pause button) in highly paused genes. Pausing durations are highly variable among genes. However, it is thought that exit from the paused state is a kinetic bottleneck in the transcription process. It is likely a consequence of Pol II pausing. We propose a renewed view of promoter motifs affecting this rate-limiting step is still unknown, particularly in the context of a developing embryo.

In parallel to these genomics approaches, single-cell imaging revealed that transcription is not a continuous process over time but occurs through stochastic fluctuations between periods of transcriptional activity and periods of inactivity, called bursting. Direct labeling of newly synthesized RNA with the MS2/MCP amplification ring has been the method of choice to observe these bursts in real time in living cells. A synthetic platform to image promoter dynamics in living embryos was developed to foster coordination in gene activation, plasticity, and development to avoid over-representation of TFIID-bound core promoter motifs (INR, DPE, pause button) in highly paused genes, which are highly variable among genes. However, relatively less attention has been given to the impact of core promoter motifs on transcriptional bursting.

To build a mechanistic understanding of bursting beyond a qualitative description of burst size and frequency, it is critical to consider the various timescales of the transcription process and to employ mathematical modeling explicitly describing various promoter states. Indeed, depending on the gene and the cellular context, transcriptional bursting has been shown to occur at multiple timescales, from seconds (e.g., polymerase clusters, Pol II firing), to minutes (e.g., TBP binding/unbinding, transcription factor binding) to hours (e.g., nucleosome remodeling, chromatin marks). These differents timescales can all occur at a single gene, and the term multiscale bursting was coined to describe such complex bursting kinetics. This multiscale bursting might explain why the simple two-state model (random telegraph), whereby promoters switch from an active to an inactive state, does not always suffice to reliably capture promoter dynamics.

While it is well understood that cis-regulatory sequences must impact transcriptional bursting, with enhancers primarily affecting burst frequencies, a detailed dissection of the impact of promoter motifs on promoter-state dynamics is lacking. In this study, we sought to examine the role of core promoter sequences on transcriptional bursting in *Drosophila* embryos. In particular, we focus on two core promoter motifs, the TATA box (TATA-WAWR) and the INR (TCAGTY in *Drosophila*) which represent pivotal core promoter contact points with the TFIIID complex, and are known to regulate both initiation and promoter pausing.

The *Drosophila* early embryo is an ideal system to decipher transcriptional bursting regulation since spatially distinct patterns of gene expression are deployed within a relatively short time frame in a multinucleated syncytium, rapidly dividing, that is highly amenable to quantitative imaging. At this early stage of development, the zygotic genome, initially transcriptionally silent, awakens progressively while cell cycle durations lengthen, a process known as the maternal-to-zygotic genome transition (MZT). This zygotic genome activation (ZGA) occurs gradually through two waves, a major wave during nuclear cycle 14 (nc14) and a minor wave occurring at earlier stages. Moreover, *Drosophila* developmental genes show a clear promoter code with well-defined promoter elements, with differential usage during ZGA.

We employed the MS2/MCP system to monitor nascent transcription. We implemented a machine-learning method that deconvolves single-nuclei mRNA production from live *Drosophila* embryos to detect all single polymerase initiation events. The waiting times between successive initiation events were further analyzed to infer the number of promoter states and the transition rates among these states. Our results show that TATA box-containing promoters are highly permissive to transcription through long ON durations and high Pol II firing rates. However, the presence of an INR in the core promoter necessitates the use of a three-state model, with a second inactive promoter state that is likely a consequence of Pol II pausing. We propose a renewed view of promoter pausing whereby only subsets of polymerases enter into a paused state.

### Results

**A synthetic platform to image promoter dynamics in living embryos.** To examine how the core promoter sequence influences gene expression variation, we developed a synthetic platform whereby several core promoters were isolated, cloned into a minigene, and inserted at the same site in the *Drosophila* genome (Fig. 1a). This approach allows for a direct comparison of core promoter activity since differences caused by variations in cis-regulatory context, genomic positioning, and mRNA sequence (particularly the 3'UTR) are eliminated. Core promoters were inserted immediately downstream of the *snail* (*sna*) distal minimal enhancer (*snaE*). The core promoters were selected based on the presence of known core promoter motifs, such as a TATA box in *sna* or an INR in *kruppel* (kr) and *Insulin-like peptide 4* (*Ilp4*). We also selected the *brinker* (*brk*) core promoter, which is devoid of any known canonical promoter motifs. None of these four promoters have a DPE, however, *Ilp4* possesses a bi-partite bridge element. These developmental genes are endogenously expressed during nc14, and precise TSS positions were established using embryonic CAGE (Cap Analysis of Gene Expression) datasets. For all transgenes, we used 100 bp of a promoter sequence, as previous work established that minimal promoter sequences are sufficient to establish pausing in vivo.

To track transcription, 24 MS2 stem loops were placed in the 5' UTR downstream of the promoter, followed by the sequence of the *Drosophila* yellow gene, a gene fragment used as a reporter because of its large size and lack of endogenous expression in the early embryo. MS2 stem loops in transcribed mRNA were bound by a maternally supplied MCP-GFP fusion protein, which allowed the detection of transcriptional foci as bright GFP spots. We imaged living *Drosophila* embryos at a high temporal resolution of one frame per 3.86 s and focused on the first 30 min of nc14. To ensure that all quantified nuclei experience similar peak levels of transcriptional activators, we restricted our analysis to a defined domain within the mesoderm (25 μm on either side of the presumptive ventral furrow; Fig. 1b). Signal intensity at the transcription site was retrieved for each nucleus after 3D detection and tracked throughout nc14 (with mitosis between nc13 and nc14 considered as time zero; see “Methods”
section) and then associated with their nearest nucleus. This generated individual 4D nuclear trajectories (Supplementary Fig. 2; “Methods”).

Core promoters differentially affect transcriptional synchrony. We hypothesized that differences in core promoter motifs would result in variability in gene expression independently of specific enhancer contacts. To test this, we characterized single-nucleus transcriptional activity (Fig. 1c–e), as well as the initial timing of activation. As previously reported with fixed sample approaches21,58, we observed a spectrum of synchrony profiles (Fig. 1f), defined as the temporal coordination of activation among a spatially defined domain. The TATA-driven sna and INR-driven kr minimal promoters showed a rapid activation with a t50 (the time needed for 50% of nuclei within the region of interest to show GFP puncta) of 11 min and 14 min, respectively while the INR-driven llp4 promoter was delayed with t50 of 21 min. The brk core promoter did not surpass 24% of nuclei activated throughout nc14.

In addition to differences in synchrony, the overall mRNA production also varied between transgenes (Fig. 1g). Within our four developmental promoters, promoters showing more rapid reactivation in nc14 also had higher TS intensity in active nuclei,
Fig. 1 A synthetic transgenic platform to image promoter dynamics. a Schematic view of transgenes used to study transcriptional dynamics of sna, kr, lip4, and brk core promoters. A minimal sna enhancer was placed upstream of the core promoter followed by 24xMS2 repeats and a yellow reporter gene. Core promoter motifs are indicated in the inset. b Schematic of Drosophila embryo showing spatial restriction of analysis to presumptive mesoderm (purple). c Maximum intensity projection of representative 15 μm Z-stack of sna < snaPr < 24xMS2-y (snaE < sna) nc14 embryo showing MS2/MCP-GFP-bound transcriptional foci (GFP) and nuclei (His-RFP). Scale bar is 5 μm. d Sample single-nuclei trace showing GFP fluorescence during nc14. The surface of the green region indicates trace integral amplitude. e False-colored frames from live imaging of indicated promoters showing relative instantaneous fluorescence intensity in early and late nc14. Inactive nuclei are gray and highly active nuclei are in yellow. f Cumulative activation curves of all nuclei during the first 30 min of nc14 for sna (green), kr (purple), lip4 (red), and brk (blue). Time zero is from anaphase during nc13-nc14 mitosis. g Average instantaneous fluorescence of transcriptional foci of active nuclei during the first 30 min of nc14 for sna (green), kr (purple), lip4 (red), and brk (blue). Time zero is from anaphase during nc13-nc14 mitosis. h Distribution of individual trace integral amplitudes from first 30 min of nc14 for sna (green), kr (purple), lip4 (red), and brk (blue). The intensity amplitude at a given time may result from the overlap of several bursts and is a convolution of promoter active/inactive times, polymerase initiation frequency, and the duration of a single polymerase signal. The integral amplitude estimates the transcriptional activity and the total number of transcripts at a steady state; it is proportional to the probability of active state (pON) initiation rate (kINI) and to the duration of the signal. Solid lines represent median and dashed lines first and third quartiles, using a one-tailed Kruskal–Wallis test for significance with multiple comparison adjustments. Statistics: snaE < sna, 216 nuclei, 3 movies; snaE < kr, 243 nuclei, 4 movies; snaE < lip4, 114 nuclei, 2 movies; snaE < brk, 45 nuclei, 2 movies. See Supplementary Movies 1 and 5.

indicating increased instantaneous mRNA production. To examine the total mRNA output, we looked to the integral amplitude, or the area of the curve defined by the average MCP-GFP signal plotted over time (Fig. 1d). The integral amplitude showed that promoters mediating faster activation and higher instantaneous mRNA production had a higher total mRNA output (Fig. 1h).

Thus, we established an experimental setup and quantification pipeline that allows disentangling the contribution of minimal promoter sequences on transcriptional activation of individual, naturally synchronized nuclei within a well-defined, homogeneous spatial domain in live embryos. This initial quantitative comparison of four natural developmental promoters suggests that those with a canonical promoter motif (either TATA or INR) tend to produce higher levels of expression.

A machine-learning method to infer promoter-state transition rates. With current labeling and imaging technology, individual transcriptional initiation events cannot be easily and directly visualized, tracked, and quantified in the early Drosophila embryo. Live imaging of transcription typically shows spots comprised of several newly synthesized mRNAs resulting from the action of multiple polymerases. In order to calibrate fluorescent signals from live imaging, we used single-molecule hybridization experiments. Using the fluorescence of a single mRNA molecule, we estimated the average number of mRNA molecules present at the TS within the nucleus at a steady state (Supplementary Fig. 3). This calibration step allowed us to express single-nuclei TS intensities as an absolute number of transcribing polymerases (Fig. 2a).

We then used a recently described novel machine-learning method to infer the transcriptional bursting mechanism. This method involves three major steps: detection of successive initiation events for each nucleus, multiexponential parametric regression of the distribution of waiting times between successive events, and identification of Markovian promoter-state transition models.

In order to detect initiation events, we considered that each trace results from the convolution between (i) the signal from a transcriptionally active promoter (Fig. 2b) and (ii) the signal produced by a single polymerase (Fig. 2c). The deconvolution procedure uses a genetic algorithm to determine optimal Pol II positioning within the gene body (Fig. 2c) and thus the time between successive Pol II initiation events (Δt) (Fig. 2d). In this first step, some specific parameters are fixed, such as the temporal resolution (3.86 s), the speed of Pol II elongation (45 bp s⁻¹ in Drosophila embryos), the size of the transcript (5.6 kb), and the retention time of the mRNA at the transcription site (assumed to be small relative to the time required to produce a transcript).

Having obtained temporal maps of individual Pol II transcription events, the method then statistically analyzes the distribution of waiting times between two successive polymerases (Δt). A multiexponential regression fitting was applied to the distribution of Δt, indicating the number of rate-limiting transitions required to best fit the data ("Methods"). We used confidence intervals and a Kolmogorov–Smirnov test to rigorously determine the smallest number of rate-limiting steps fitting our experimental data (Supplementary Table 1). Following the principle of parsimony and to avoid overfitting, models with a larger number of steps and more parameters were not retained even if they also fit well. Importantly, this step uncovered the number of characteristic timescales of transcriptional fluctuations that are tantamount to the number of promoter states.

Finally, a transcriptional model of the promoter was established, with multiple states and timescales inferred from the parameters of the multiexponential Δt distribution (Supplementary Table 2). This step permitted the estimation of the kinetic parameters such as the kON (rate of switching from a transcriptionally nonpermissive state to a permissive one), kOFF (switching rate from a transcriptionally permissive to a nonpermissive state), and kON (the rate of Pol II initiation events once the promoter is in a transcriptionally permissive state) for the simplest two-state model. However, kinetic estimates of more complex models (three or more promoter states) can also be derived from the Δt distribution. The accuracy and robustness of the deconvolution method were tested by applying it to artificial data with known positions of transcription initiation events and known kinetic parameters (Fig. 2e–h). These artificial data were generated using the Gillespie algorithm and proved the robustness of the approach (see "Methods").

Importantly, this approach provides a time map of transcription events in a cell population in a model-independent manner. The number of promoter states is evaluated during the multiexponential fit procedure, but with no a priori constraints on the result. This contrasts with current methods which directly fit a particular transcription model to the data, such as methods based on the autocorrelation functions, maximal likelihood estimates, or Bayesian inference.

The sna promoter as a model to investigate the impact of core promoter on transcriptional dynamics. To investigate the quantitative dynamics of the sna promoter, we applied this method to hundreds of nuclei pooled from snaE < sna embryos. A heatmap of positions of single initiation events for each nucleus

ARTICLE NATURE COMMUNICATIONS | https://doi.org/10.1038/s41467-021-24461-6

NATURE COMMUNICATIONS | (2021) 12:4504 | https://doi.org/10.1038/s41467-021-24461-6 | www.nature.com/naturecommunications
revealed the sna promoter drove frequent transcriptional initiation events. Silent periods with no initiating polymerase were very rare (Fig. 2i–j, white bars), consistent with the results obtained from fixed embryos.\(^{21,66}\). We found that a biexponential (sum of two-exponential functions) fitting correctly fit the survival function of polymerase waiting times (Fig. 2k). The biexponential fitting is only compatible with a two-state model (Fig. 2k). Thus, the transcriptional activity of the sna promoter can be described by the random telegraph model, with a simple random switch between an inactive OFF state and a permissive ON state, from which transcription initiates at a given rate (\(k_{\text{INI}}\); Fig. 2l). The probability to occupy the permissive ON state was high in the case of snaE < sna.\(^{21}\) The \(k_{\text{INI}}\) for sna was estimated at one polymerase per 9 s (Fig. 2l), consistent with the inferred initiation rate at its endogenous locus\(^{66}\) and in the range of recent
estimates of initiation frequencies for other developmental genes in Drosophila and in mammalian cells. Collectively this shows that we can locate individual initiation events, determine the number of promoter states, select a promoter model by the principle of parsimony, and then estimate average promoter switching rates in vivo.

The TATA box regulates the ON and OFF duration but not the number of states. To determine how the TATA box influences transcriptional kinetics in vivo, we developed a series of snai core promoter mutants (Fig. 3a) where the TATA box was replaced with either the TATA-like sequence of kr (snaiTATAlight) that is bound by TBP (Supplementary Fig. 1) or a non-TATA sequence where the first four bases are mutated (snaiTATAmut). Surprisingly, the strong TATA mutation did not completely abolish the activity of the snai promoter, although this mutant promoter was devoid of any known core promoter motifs. In comparison to the snai promoter, the synchrony of the snai TATA mutants was reduced in a graded manner relative to the fidelity of the TATA box (Fig. 3b, Supplementary Movies 1–3). The snai promoter reached synchrony at ~11 min into nc14, while the snaiTATAlight promoter reached synchrony at 24 min and the snaiTATAmut promoter never reached synchrony. This implies that the TATA box influences synchrony of activation, most likely by promoting and stabilizing TBP binding (Supplementary Fig. 1).

Similar to the activation profiles, instantaneous TS intensities were directly correlated with TATA box sequence fidelity (Fig. 3c). Interestingly, while both the snai promoter and snaiTATAlight promoters appeared to reach a steady state, the snaiTATAmut was unable to do so. This may have been the result of MCP-GFP signal being too low to consistently surpass the detection threshold, as single-molecule FISH experiments indicated most nuclei in the ventral region were at least weakly active in nc14 snaiTATAmut embryos (Supplementary Fig. 4a–c). The total mRNA production of each promoter measured by the integral amplitude was similarly affected (Fig. 3d), with the snai promoter driving the highest total mRNA expression while the TATA box mutants expressed reduced amounts of mRNA correlated to the fidelity of the TATA box.

We next asked whether the snaiTATAlight and snaiTATAmut transgenes still followed a two-state model. For each nucleus of each of these three transgenic genotypes, we located single Pol II initiation events during the first 30 min of nc14 (Fig. 4a–f and Supplementary Fig. 4d–j). By examining the survival curve of Pol II waiting times, the dynamics of the snai TATA mutants appeared to be accurately recapitulated by the two-state random telegraph model, similarly to the unmutated snai promoter (Supplementary Fig. 4g–i, Supplementary Table 1). We then analyzed which kinetic parameters changed in the mutant promoters. Extracting kinetic parameters for the snaiTATAlight transgene revealed a relatively mild effect on initiation rates (1 Pol II per 13 s for snaiTATAlight instead of 1 Pol II per 9 s) (Fig. 4g–i and Supplementary Table 2). In contrast, we observed a strong increase in the TOFF (Fig. 4g–k) for the snaiTATAlight promoter relative to the snai promoter (Fig. 4g–i, k and Supplementary Fig. 5b). The strong TATA mutation led to a similar trend, although due to the weak activity of this promoter, the number of analyzed nuclei for this mutant is lower than for the other promoters. The kinetic parameters of TATA mutant promoters led to an overall reduction in burst size, defined as the number of transcripts produced during an active period (Fig. 4l). The reduction was primarily due to a decrease in the duration of the ON periods, consistent with a destabilized TBP/TATA box interaction or reduced TBP recruitment to the promoter.

![Fig. 3 Decoding the role of the TATA box on promoter dynamics.](image)

**Fig. 3 Decoding the role of the TATA box on promoter dynamics.** a TATA box mutations of the snai promoter. The snaiTATAlight mutation corresponds to the sequence of the non-canonical TATA box from the kr promoter. b Cumulative active nuclei percentage within the mesodermal domain for snai (dark green), snaiTATAlight (green), and snaiTATAmut (light green). c Average instantaneous fluorescence of transcriptional foci of active nuclei during the first 30 min of nc14 for snai (dark green), snaiTATAlight (green), and snaiTATAmut (light green). Time zero is from anaphase during nc13-nc14 mitosis. False-colored panels on the right are colored according to instantaneous fluorescence intensity, with inactive nuclei shown in gray and highly active nuclei in yellow. d Distribution of individual trace integral amplitudes from first 30 min of nc14. The solid line represents median and dashed lines the first and third quartiles, using a one-tailed Kruskal–Wallis test for significance with multiple comparison adjustments. Statistics: snai < snai, 216 nuclei, 3 movies; snai < snaiTATAlight, 353 nuclei, 6 movies; snai < snaiTATAmut, 21 nuclei, 3 movies. See Supplementary Movies 1-3.

The impact of TATA on the duration of ON/OFF promoter states observed in Drosophila embryos is also in agreement with results from live imaging of HIV-1 transcription in human cells, and with genomic studies in cultured mammalian cells, where TATA-driven promoters are associated with large burst sizes. They differ from those obtained with mutations of the native
The INR motif induces a third rate-limiting promoter state. At the biochemical level, transcriptional activation is a multistep process requiring the orchestration of many factors, and yet TATA-driven promoters can be modeled with a simple two-state model corresponding to one kinetic rate-limiting step in this complex process. We hypothesized that additional rate-limiting steps were likely to exist and could be identified using our imaging-based methodology. As the TATA box did not affect the number of promoter states for the *sna* promoter, we examined the effect of another core promoter motif, the INR. In *Drosophila* cells and embryos, the INR is associated with stably paused genes, genome wide. Moreover, analysis in cell culture indicated that genes with increased pausing stability tend to harbor an INR motif in the core promoter, and in particular a preferential G at the +2 position. We, therefore, reasoned that manipulating the INR motif in embryos might affect pausing and therefore may induce changes in rate-limiting promoter states.

To examine the role of the INR core promoter motif, we created a series of transgenes in which we manipulated the INR without changing the enhancer or the downstream gene sequence (Fig. 5a). The *sna* core promoter does not have an INR, while the core promoter of *kr* has a natural INR sequence with a G at +2 and...
is stably paused\textsuperscript{16,18}. Moreover, the kr promoter possesses a non-canonical TATA box (TATAlight, Fig. 1a). We exchanged the TSS region of sna with the INR of the kr promoter (sna+INR). We also created a transgene (Fig. 5a) whereby the INR of kr was replaced with the TSS region of sna (kr-INR1). Interestingly, neither loss of the INR in the kr-INR1 transgene nor gain in the sna+INR transgene strongly affected synchrony (Fig. 5b and Supplementary Movies 1, 4, 5). The addition of an INR to sna did not dramatically affect mRNA output (Fig. 5d). Both the sna+INR and kr-INR1 transgenes had higher instantaneous activity than the cognate wild-type promoters (Fig. 5c). Moreover, a second mutation of the INR in kr (kr-INR2) had similar synchrony and mRNA production (average instantaneous intensity) as in the wild-type kr promoter (Supplementary Fig. 6a–c). We then applied the deconvolution procedure to each genotype (Fig. 6a and Supplementary Figs. 6d–k and 7a, d–k). Surprisingly, our analysis of polymerase initiation events indicated that, for sna+INR nuclei, a two-state model was not sufficient to fit the data (Fig. 6a, b, Supplementary Fig. 7i, and Supplementary Table 1). Instead, the sna+INR survival function was well fitted by adding an extra exponential term. Thus, a three-state model appropriately recapitated sna+INR promoter dynamics (Fig. 6a, b and Supplementary Fig. 7i–i’). Similarly, the kr promoter also required a three-exponential fitting (Fig. 6b and Supplementary Fig. 6h–i’). In contrast, removing the INR from this promoter (kr-INR1 promoter) led to a two-exponential fitting for the survival function (Fig. 6b and Supplementary Fig. 6i–i’). Similar results were obtained with a second INR mutant (kr-INR2) (Supplementary Fig. 6j–j’). Furthermore, the natural INR-driven Ilp4 promoter was also associated with a three-exponential fitting, and mutation of its INR sequence (Ilp4-INR) resulted in a reversion to a two-exponential fit (Fig. 6b and Supplementary Fig. 8d, e, h, i’). These results, therefore, suggest that the INR motif is associated with a third rate-limiting promoter state.

**Pol II pausing is associated with a third promoter state.** Given the correlation between the presence of an INR motif and the number of characteristic timescales and thus the number of promoter states (three in the presence of an INR), we reasoned that one of these states could be polymerase pausing. To test this hypothesis, we impaired the establishment of pausing by reducing the expression of the largest subunit of the pause-inducing negative elongation factor complex (NELF), NELF-A. Previous work demonstrated that depletion of NELF by RNAi globally reduced promoter-proximal pausing in *Drosophila* cells\textsuperscript{19}, and that loss of the human C-terminal NELF-A "tentacle" blocked stabilization of pausing\textsuperscript{69}. We combined
the inducible RNAi/GAL4 system with our MS2/MCP-GFP reporter to examine the transcriptional dynamics of the three-state kr promoter. We performed live imaging of the kr-MS2 transgene and compared a control white RNAi knockdown with embryos harboring a maternal depletion of Nelf-A (evaluated to >85% knockdown by RT-qPCR, Supplementary Fig. 9a, Supplementary Movies 7 and 8). Remarkably, reducing Nelf-A transcript levels led to a change in kr promoter dynamics, with a reversion to a two-promoter-state dynamic and much less frequent long waiting times (Fig. 6b and Supplementary Fig. 9b, arrowheads, Supplementary Table 1). These data, combined with the effect of INR mutations in cis, led us to propose that one of the two inactive promoter states observed with the three-state promoters could be associated with promoter-proximal polymerase pausing.
To confirm that the INR motif was truly altering Pol II pausing, we performed Pol II ChIP-qPCR on our promoter transgenes from staged embryos. We evaluated the strength of pausing by computing the pausing index. The pausing index (PI) was obtained by quantifying the Pol II binding at the promoter relative to the gene body17 (Fig. 6d). As shown previously, in our transgenic embryonic extracts encompassing various stages from nc13 to late nc14, the wild-type sna promoter is paused18,21,30, but the addition of an INR motif significantly increased its PI (Fig. 6e), consistent with INR-mediated lengthening of the pause duration observed in cell culture15. To examine the inverse scenario, we also quantified the PI of the kr and kr-INR1 transgenic promoters. Similar to what is observed in the endogenous promoter18, the kr transgene was highly paused (Fig. 6e). Mutation of the INR in the kr-INR1 transgene reduced pausing (Fig. 6e). Likewise, the Ilp4 promoter from our transgene showed a high level of pausing, which decreased upon mutation of its INR sequence (Fig. 6d). Both the sna+INR and kr promoters also show increased NELF-E enrichment at these promoters by ChIP-qPCR on embryonic extracts (Supplementary Fig. 9g). Taken together, characterization of the INR core promoter motif suggests the presence of a strong INR motif results in stabilization of pausing in vivo.

**Pol II does not undergo systematic pausing.** Next, we envisaged how pausing could be modeled by three distinct promoter states. The current view of pausing is that all polymerases systematically enter into pausing, a step followed by either a productive elongation or a termination17,71–73. We therefore initially tested if our data could be explained by a model consisting of three promoter states (inactive OFF, permissive ON, and paused) where all productive polymerases undergo pausing prior to transitioning to the permissive ON state (Fig. 6c). This model was clearly incompatible with our data, as the fitting exceeded the bounds of the 95% confidence interval (Supplementary Figs. 6d and 7i†). Recent work on the prototypic model of a highly paused promoter, the HIV-1 promoter, also found that modeling transcriptional dynamics with obligatory pausing was not in agreement with live-imaging data in HeLa cells, and instead proposed an alternative model of pausing where only a fraction of polymerases are subject to pausing in a stochastic manner59. We asked if this alternative non-obligatory pausing model could be applicable to our findings (Fig. 6c). According to the goodness of fit, nonsystematic pausing was compatible (Supplementary Figs. 6h† and 7i†). Our machine-learning method enabled the estimation of the kinetic transition rates between these states, summarized in Supplementary Table 2 and Supplementary Fig. 5a–c.

We conclude the presence of an INR motif translates into a longer paused state that creates an additional rate-limiting step during transcription in vivo. In our kinetic model, the inactive states could in principle equally correspond to pausing. These two-promoter states are discernible by their distinct timescales, on the order of seconds and minutes, respectively (Supplementary Table 2). Because the NELF knockdown dramatically decreases the frequency of long waiting times (Supplementary Fig. 9e–f), we favor the longer-lived inactive state, which lasts on the order of minutes, as the potential pausing state. The second short-lived inactive state, lasting on the order of seconds and present for both TATA- and INR-containing promoters, would then correspond to a nonpermissive promoter state, possibly TFIID-unbound as its frequency increases in TATA mutants (Fig. 4k and Supplementary Table 2). As the sna and kr endogenous promoters drive a high level of gene expression (Supplementary Fig. 1), it is reasonable to hypothesize that such a nonpermissive state is transient and infrequent (Supplementary Table 2).

**INR control of promoter switching kinetics.** Like the natural sna promoter, the sna+INR promoter showed a high probability of occupying the transcriptionally permissive ON state, with a long lifetime of 170 s. This ON duration was slightly reduced compared to the sna transgene (Fig. 6f, i), possibly implying competition between the natural canonical TATA box and the added INR in the sna+INR combination (see below). The major difference between these two promoters was the existence of two distinguishable inactive states linked to the presence of the INR sequence. One of these was short-lived (17 s) and one long-lived (302 s) (Fig. 6f, i, Supplementary Fig. 5b, c, and Supplementary Table 2). The stable long-lived paused state was not observed with the sna promoter but did occur in the presence of the INR-containing kr promoter.

In the case of kr and Ilp4 developmental promoters, the paused state lasted ~151 and 105 s, respectively, but was reached relatively rarely (Fig. 6g, j, Supplementary Fig. 5c, and Supplementary Table 2). Pausing was not observed upon two types of mutations of the kr INR motif (Fig. 6f, Supplementary Fig. 6i–j†, m, and Supplementary Table 2). Thus our data favor a model where the paused state lasts several minutes (in the case of kr, sna+INR, and Ilp4), but occurs infrequently. Interestingly, the sna promoter is also moderately paused in our ChIP assay and at its endogenous locus74, but this promoter is not regulated by a three promoter-state dynamic. We hypothesize that pausing is highly unstable for this promoter at this stage and does not constitute a rate-limiting step that we can capture with our live-imaging assay (see “Discussion”).

In addition to two separable inactive states, the kr transgene exhibited a highly probable permissive state (0.83) with a duration of 94 s (Fig. 6g, Supplementary Fig. 5a, and Supplementary Table 2). When the INR of kr was mutated (kr-INR1), there was a significant increase in ON duration (Fig. 6f) and Supplementary Table 2). A similar effect was obtained with an alternative mutation of the INR (kr-INR2) (Supplementary Figs. 5a and 6m and Supplementary Table 2).

Interestingly, in all the transgenes, the differences in the number of states and the active/inactive durations were not accompanied by a change in Pol II firing rates. The sna, sna+INR, kr, and kr-INR1 mutant all exhibited an estimated kINI of 1 Pol II every 8 s (Fig. 6f–g, i–j and Supplementary Table 2). These estimates agree with recently documented polymerase initiation rates of Drosophila developmental promoters30,32,75. Our results suggest the rate of initiation is not associated with the presence or absence of an INR motif. Instead, the INR motif leads to a regulation of transcriptional bursting via two inactive promoter states, one of which is associated with stabilized pausing.

Taken collectively, our results demonstrate that transcription dynamics are differentially regulated by the INR and the TATA box. Developmental promoters containing a strong INR motif can travel through multiple inactive states due to an extra regulatory step at early elongation via Pol II pausing. Our live-imaging data favor a model whereby pausing occurs on the order of minutes but is not an obligatory state reached by all engaged polymerases. Instead, we propose that when pausing constitutes a rate-limiting step, it occurs for only a subset of polymerases with a low frequency.

**Promoter sequences and timing of initiation.** A second property that we found to be common to all promoters examined, was the regime of waiting times between mitosis and first transcriptional activation. The analysis of the lag time between mitosis and onset of activation is “non-stationary” and was previously modeled using a different modeling paradigm, based on mixed gamma distributions of the random time to transcription activation after mitosis in nc1457. This analysis estimates two parameters, the number of promoter states (a) and transition times between them (b). We applied this
modeling framework to our data by only focusing on the distribution of waiting times between mitosis and first activation (illustrated in Supplementary Fig. 10a). Interestingly, a multieponential distribution becomes a mixed gamma distribution when the time parameters of the exponentials are even. Conversely, the equality of these time parameters justifies the use of a mixed gamma regression. Using the more general multieponential regression, we proved that regardless of the promoter sequence, the transition times between states during this lag regime are homogeneously distributed (Supplementary Fig. 10b, c). This is in contrast with the transition times between states in the stationary bursting regime, which are heterogeneous. This suggests that the states and transitions involved in the two regimes (nonstationary and stationary) are distinct and have separate regulations. This further supports a previous study that demonstrated the delay in postmitotic transcription activation is dependent on the enhancer sequence.

Interplay between TATA and INR. Recent single-cell RNA-seq quantification suggests that promoters exhibiting both TATA and INR motifs produce higher burst sizes than when only one motif is present. However, systematic interrogation of human promoters with a synthetic biology approach shows that TATA and INR additively but not synergistically increase gene expression. A synergetic and/or additive effect does not seem to be evident in Drosophila, as TATA-containing promoters have relatively infrequent INR motifs or exhibit an INR sequence devoid of a G at +2, shown to be associated with long pause durations.

To quantify TATA box and INR interplay on rate-limiting promoter switching states in vivo, we compared the sna + INR and snaTATAlight transgenes to that of a combined mutant snaTATAlight+INR promoter (Supplementary Fig. 7a). Remarkably, after applying the deconvolution procedure to snaTATAlight+INR nuclei (Supplementary Fig. 7g), the distribution of waiting times between initiation events required a fit with two exponentials similarly to snaTATAlight (Supplementary Fig. 7k–k’). We note however that the statistical tests allocating the number of promoter states for this particular genotype were not robust (Supplementary Table 1). In terms of promoter kinetics, the snaTATAlight+INR behaved similarly to the snaTATAlight promoter (Supplementary Fig. 5a, b). Interestingly, the snaTATAlight+INR has a p(OFF) nearly twice that of the snaTATAlight promoter (Supplementary Table 2), indicating that the INR may act to maintain a nonpermissive promoter state even in the absence of polymerase pausing.

To gain more insight into TATA/INR interplay, we also generated a kr mutant promoter that does not contain a noncanonical TATA (kr-TATA; Supplementary Fig. 6a). Data from this promoter required a three-state model (Supplementary Fig. 6k–k’), similar to the kr promoter. In the kr-TATA promoter, the duration and probability of the paused state increased while the ON state appeared shorter and less probable (Supplementary Fig. 5a, c and Supplementary Table 2). This indicates that the loss of TBP binding may increase pausing induced by the INR motif, potentially providing a mechanistic rationale for the relative scarcity of dual TATA/canonical INR promoters in the Drosophila genome. Taken together, these results indicate that TATA and INR are not simply synergistic or additive but display more complex interactions.

Discussion

The spatiotemporal organization of gene expression is critical to the development of a functional living organism. While we have accumulated knowledge on how enhancers precisely regulate gene expression, we know relatively little about the impact of core promoters on transcriptional bursting in a developing embryo. Here, we investigated how minimal 100 bp sequences of developmental promoters control transcriptional states and their switching kinetics. We found that a classical two-state model does not suffice to capture promoter dynamics of stably paused promoters containing a strong INR motif but does fit with TATA-containing promoters.

The development of a novel numerical deconvolution method revealed startling insights into the variation of transcriptional initiation between minimal promoter sequences in vivo. Our experimental setup allowed us to unmask gene expression variations, which are only dictated by variations in core promoter sequences. Indeed, cell cycle duration and synchrony, the concentration of input transcription factors, and chromatin states were intentionally kept constant. Our assay revealed that a similar transcriptional activity profile could be obtained from two very distinct promoter sequences via distinct modes of transcriptional initiation. While sna promoter dynamics could be explained by a simple two-state model, the kr promoter required a three-state model, with two distinct inactive promoter states, a short-lived and a longer-lived one. These distinct transcriptional regimes are likely due to differences in enhancer–promoter specificity, as evidenced by the synchronous and high transcriptional activation reached with these two promoters. However, knowing the wide-spread preferential promoter code for specific enhancers, it will be interesting to use our pipeline to examine which rate-limiting step of promoter dynamics is tuned by enhancer–promoter choice.

How could such a small number of states be compatible with the numerous promoter-binding events occurring during transcription initiation? Biochemical studies reveal structural states, while imaging-based approaches unmask key rate-limiting kinetic states. Whole-genome methylation footprinting disentangled five transcription initiation states in Drosophila cultured cells. Remarkably, the authors demonstrated that TATA-containing promoters were frequently found in PIC-bound configuration (PIC alone or PIC + Pol II). We, therefore, propose that the permissive state in Drosophila embryos corresponds to a state where promoter DNA is bound by the PIC. The inactive state, which we found to be very transient, could then possibly represent a TBP-unbound configuration consistent with TBP dynamics observed in human cells.

By using the sna promoter as a model, we found that TATA box directly impacts promoter occupancy of the active state and allows large transcriptional bursts by promoting long ON durations. How could the presence of a TATA box permit these long ON durations? Slow TBP protein turnover at human TATA box-containing genes may foster stable long ON durations. However, this hypothesis needs to be confirmed by quantification of TBP kinetics during Drosophila ZGA.

In its endogenous context, the snail promoter is among the first genes to be transcribed during ZGA, in a particularly constrained environment with extremely short cell cycles (<15 min). Remarkably, the majority of genes expressed during this critical period are highly expressed, short, and intron-less with a canonical TATA box and are generally nonpaused. Subsets of these, including snail, are considered “dual promoters”, as they gradually acquire pausing as development proceeds. Thus, it is possible that the kinetic bottlenecks regulating transcription of developmental promoters evolve as developmental timing proceeds, with paused polymerase gradually emerging as a rate-limiting step. In our study, the promoter model sna shows a moderate level of pausing (as measured by ChIP) but is regulated by a simple two-state model in nc14. This could indicate that pausing occurs on a timescale indistinguishable from the OFF state and is thus embedded within it. Strikingly, early transcription in zebrafish embryos occurs in similarly constrained rapid cell cycles where a subset of zygotically activated genes also
display a T/A-rich WW-box motif. Thus, regulation of transcription via a unique rate-limiting step (OFF to ON transition) that is TBP-dependent might be conserved between fly and vertebrate embryos.

In this work, we provided quantitative evidence that INR-containing promoters are associated with a three-state model (ON, paused, OFF), contrasting with TATA promoters.

Interestingly, genomics studies revealed that transcription initiation code evolves during ZGA in flies and vertebrates. Given the results of this study and those obtained in Drosophila cultured cells, it is tempting to link the gradual emergence and stabilization of pausing during ZGA with the presence of an INR, in particular an INR with a G at +2 position. In light of our results, we propose that the switch in promoter usage from TATA-driven to INR-driven during ZGA may lead to a change in transcriptional dynamics from two to three states, to include an elongation-mediated checkpoint. Acquisition of an extra rate-limiting step might help control cell-to-cell expression variability as well as fine-tune gene expression levels.

Taken collectively, this study establishes our promoter imaging assay and novel mathematical deconvolution and modeling methodology as a valuable tool to probe gene expression and novel mathematical deconvolution and modeling methodology as a valuable tool to probe gene expression dynamics during development. Quantitative analysis of promoter dynamics at high temporal resolution opens the door to a deeper insight into the molecular mechanisms underlying transcriptional regulation in vivo. Future studies involving direct manipulation of pause initiation or duration in living embryos using approaches such as optogenetics will help to establish a broader understanding of the nature of promoter states and the role of Pol II pausing in vivo.

Methods

Drosophila stocks and genetics. All crosses were maintained at 25 °C. Transgenic lines were maintained as homozygous stocks. For live imaging, homozygous males carrying the transgene of interest were crossed with homozygous females bearing the transgene of interest were crossed with homozygous females carrying the MCP-GFP-His2A-RFP construct. For smiFISH, homozygous flies were crossed to yw in order to facilitate single-molecule detection. nosGAL4-VP16 was recombined with MCP-GFP-His2A-RFP57. UAS-white RNAi (UAS5573) and UAS-Nef-A RNAi (UAS2987) were obtained from the Bloomington Stock Center (University of Indiana, Bloomington).

Cloning and transgenesis. The smo distal enhancer-24xMS2-y minigene has been previously described.12,13. Promoters (Supplementary Fig. 1 and Supplementary Table 3) were amplified from genomic DNA using Q5 polymerase (New England Biolabs) and inserted between the enhancer and 24xMS2 sequences using restriction enzyme-mediated ligation. Mutations were performed with the QuikChange II Site-Directed Mutagenesis kit (Agilent Technologies) or synthesized (Twist Bioscience) and inserted using restriction-mediated cloning. All constructs were sequenced to ensure appropriate insertion. Transgenic flies were generated using PhiC31-mediated recombination (Best Gene, Inc.), and all constructs were inserted into the same genetic background and genomic position (BL 9750). Stocks are available upon request.

Live imaging. Embryos were permitted to lay for 2 h prior to collection for live imaging. Embryos were hand dechorionated and mounted on a hydrophilic membrane prior to immersion in oil to prevent desiccation, followed by the addition of a coverslip. Live imaging was performed with an LSM 880 with Airyscan module (Zeiss). Z-stacks comprised of 30 planes with a spacing of 0.5 µm were acquired at a time resolution of 3.86 s per stack in fast Airyscan mode with laser power measured using a ThorLabs PM100 optical power meter (ThorLabs Inc.), and maintained across embryos at 3.8 µW for constitutive MCP-GFP-His2A-RFP expression, and 5.0 µW for RNAi analyses. All movies were performed with the following settings: GFP excitation by a 488-nm laser and RFP excitation by a 561-nm were captured on a GaAsP-PMT array with an Airyscan detector using a ×40 Plan Apo oil lens (NA = 1.3) and a 3.0 zoom on the ventral region of the embryo centered on the presumptive ventral midline. Resolution was 512 × 512 pixels with bidirectional scanning. Airyscan processing was performed using 3D Zen Black v3.2 (Zeiss).

Live-imaging analysis. Visualization and analysis of the time series were performed using a custom-made software developed in Python that permits visualization of each analysis step and manual correction if necessary (Supplementary Fig. 2). Activation time traces were collected starting from the Airyscan-processed Z-series described above. Green (MS2) and red (His2A) channels were clipped to consider only after the start up to the progression of anaphase across the region of interest. Nuclei were maximum intensity projected and pre-smoothed with a Gaussian filter and then thresholded with an Otsu threshold value. The resulting connected components of the binary images were then labeled and touching nuclei segmented with a watershed algorithm. The software enabled manual correction of the segmentation. Nuclei were finally tracked across the time frames using a minimum distance criterion plus a user-defined distance threshold. Nuclei appearing for only a few frames or those touching the border were removed from the analysis to exclude sources of errors. The transcription sites were averaged based on the proximity of sister chromatids and vertebrates (Supplementary Fig. 2). Activation time traces were collected starting from the Airyscan-processed Z-series described above. Green (MS2) and red (His2A) channels were clipped to consider only after the start up to the progression of anaphase across the region of interest. Nuclei were maximum intensity projected and pre-smoothed with a Gaussian filter and then thresholded with an Otsu threshold value. The resulting connected components of the binary images were then labeled and touching nuclei segmented with a watershed algorithm. The software enabled manual correction of the segmentation. Nuclei were finally tracked across the time frames using a minimum distance criterion plus a user-defined distance threshold. Nuclei appearing for only a few frames or those touching the border were removed from the analysis to exclude sources of errors. The transcription sites were averaged based on the proximity of sister chromatids and vertebrates (Supplementary Fig. 2). Activation time traces were collected starting from the Airyscan-processed Z-series described above. Green (MS2) and red (His2A) channels were clipped to consider only after the start up to the progression of anaphase across the region of interest. Nuclei were maximum intensity projected and pre-smoothed with a Gaussian filter and then thresholded with an Otsu threshold value. The resulting connected components of the binary images were then labeled and touching nuclei segmented with a watershed algorithm. The software enabled manual correction of the segmentation. Nuclei were finally tracked across the time frames using a minimum distance criterion plus a user-defined distance threshold. Nuclei appearing for only a few frames or those touching the border were removed from the analysis to exclude sources of errors. The transcription sites were averaged based on the proximity of sister chromatids and vertebrates (Supplementary Fig. 2).
Mathematical modeling of burst parameters and multiexponential regression fitting

Burst deconvolution and Pol II positioning. The Pol II positions were found by combining a genetic algorithm with a local optimization procedure.

Before the initiation of the analysis algorithm, several key parameters were established. The Pol II elongation speed was fixed at 45 bp s⁻¹ (see ref. 60). The reporter construct transcript was divided into three sections consisting of the pre-MS2 fragment (41 bp), 24xMS2 loops (1292 bp), and post-MS2 fragment containing the yellow reporter (4526 bp). The retention time was assumed to be small in relation to the time needed to produce a transcript, and so was fixed at 0 s. The temporal resolution of each movie was 3.86 s per frame. This frame rate is sufficient to detect processes that occur on the order of seconds.

The possible polymerase positions were discretized using a step of 30 bp (or equivalently 2/3 s). This step was chosen, as it is smaller than the minimum polymerase spacing and large enough to have a reasonable computation time. For a movie of 35-min length, this choice corresponds to a maximum number of 3150 positions. The algorithm was implemented in Matlab R2020a using Global Optimization and Parallel Computing Toolboxes for optimizing Pol II positions in parallel for all nuclei in a collection of movies. The resulting positions were stored for analysis in the further steps of our computational pipeline. At this step, the density of Pol II initiation events can be visualized by binning time and checking the occurrence of Pol II activation in each bin. This was rendered as a heatmap in which rows represent a single-nucleus time series and the number of activation events per 30-s bins (or equivalently 1350 bp) is indicated by the color (Fig. 2e).

The deconvolution step is common to all of the MS2 data analysis pipelines. A detailed description of the algorithm can be found in ref. 59.

Multiexponential regression fitting of the survival function and model reverse engineering using the survival function. Data from several movies corresponding to the same genotype was first pooled together. The entry and exit of each track corresponding to a unique nucleus were defined using a threshold representing 1/5 of the maximum intensity for the specific track, in order to restrict the analysis to the stable part of the signal (Fig. 2a, gray box). Waiting times were extracted as differences between successive Pol II positions from all the resulting traces and the corresponding data was used to estimate the nonparametric cumulative distribution function by the Meyer–Kaplan method. This also permits the calculation of a 95% confidence interval for the experimental survival function that is further used to judge the quality of a parametric multiexponential regression fitting.

Then, a multiexponential regression fitting produced a set of 2N−1 distribution parameters, where N is the number of exponentials in the regression procedure (3 for N = 2 and 5 for N = 3). The regression procedure was initiated with multiple initial guesses and followed by local gradient optimization of the following objective function:

\[ O = \alpha \sum_{n=1}^{N} \left( S(t_n) - S(t_n)^{2} \right)^2 + \frac{1}{\alpha} \sum_{n=1}^{N} \left( \log(S(t_n)) - \log(S(t_n)^{2}) \right)^2, \]

(1)

where \( S(t_n) \) and \( S(t_n)^{2} \) are the theoretical (multiexponential) and empirical (estimated by the Meyer–Kaplan method) survival functions, respectively, and \( \alpha \) is a parameter satisfying \( 0 \leq \alpha \leq 1 \) and representing the weight of linear scale differences in the objective function. We chose an intermediate value \( \alpha = 0.6 \) for all our parameter estimates (these estimates are nevertheless robust with respect to \( \alpha \)).

The optimization resulted in a best-fit solution with additional suboptimal solutions (local optima with objective function value larger than the best fit). A multiexponential regression is considered acceptable if the predicted survival function is within the confidence bounds of the experimental survival function. This provides a method to select the number \( N \) of exponentials in the regression: we progressively increase \( N \) starting with \( N = 2 \) until an acceptable regression is reached.

The 2N−1 distribution parameters can be computed from the 2N−1 kinetic parameters of a N-state transcriptional bursting model. Conversely, a symbolic solution for the inverse problem was obtained, allowing computation of the kinetic parameters from the distribution parameters and reverse engineering of the transcriptional bursting model. In particular, it is possible to know exactly when the inverse problem is well-posed, i.e., there is a unique solution in terms of kinetic parameters for any given distribution parameters in a domain.

The transcriptional bursting models used in this paper are as follows (see Tantale et al.59 for a detailed description):

For \( N = 2 \), there were three distribution parameters and three kinetic parameters.

The distribution parameters are \( A_1, \lambda_1, \lambda_2 \), defining the survival function

\[ S(t) = A_1 e^{\lambda_1 t} + (1 - A_1) e^{\lambda_2 t}. \]

(2)

The solution of the inverse problem for the ON−OFF telegraph model (Fig. 2k and Fig. 6b) is

\[ k_1 = -S_1, \]

(3)

\[ k_1^* = S_1 - S_2, \]

(4)

\[ k_2 = \frac{S_2 S_1 - S_2^2}{S_1 (S_1 - S_2)}, \]

(5)

\[ S_1 = A_1 + A_2, \]

(6)

\[ S_2 = A_2 + A_2^2, \]

(7)

\[ S_3 = A_1 + A_2^2, \]

(8)

\[ A_1 = 1 - A_2. \]

(9)

where \( k_1, k_1^*, k_2^* \) are the initiation rate, the OFF to ON and ON to OFF transition rates, respectively. Thus, the duration of the OFF and ON states can be calculated as:

\[ T(\text{OFF}) = \frac{1}{k_1^*}, \]

(10)

\[ T(\text{ON}) = \frac{1}{k_1}. \]

(11)

For this model, the probability to be in the state ON and OFF is:

\[ P(\text{ON}) = k_1^*/(k_1^* + k_2^*), \]

(12)

\[ P(\text{OFF}) = 1 - P(\text{ON}). \]

(13)

For \( N = 3 \), there were five distribution parameters and five kinetic parameters.

The distribution parameters are \( A_1, A_1, A_2, A_2, A_2 \), defining the survival function

\[ S(t) = A_1 e^{\lambda_1 t} + A_2 e^{\lambda_2 t} + (1 - A_1 - A_2) e^{\lambda_3 t}. \]

(14)

The inverse problem has a unique solution for the three-state model (non-obligatory pause) with two OFF states (OFF and PAUSE) and one ON state (Fig. 6c).

\[ k_1 = -S_1, k_2^* = \frac{1}{2} \left[ -L_1 + S_1 - \sqrt{(S_1 L_2 - S_2)^2 - 4 L_2 S_1} \right]/S_1, \]

(15)

\[ k_3^* = \frac{1}{2} \left[ -L_2 + S_2 - \sqrt{(S_1 L_2 - S_2)^2 - 4 L_2 S_1} \right]/S_1, \]

(16)

\[ k_1 = \frac{1}{2} \left[ -L_1 + S_1 + \sqrt{(S_1 L_2 - S_2)^2 - 4 L_2 S_1} \right]/S_1, \]

(17)

\[ k_3 = \frac{1}{2} \left[ -S_2 - L_2 + S_2 + S_1 L_2 - L_3 + \frac{S_2}{S_1} \right]/S_1, \]

(18)

where

\[ S_1 = A_1 + A_2 + A_2, \]

(19)

\[ S_2 = A_2 + A_2 + A_2, \]

(20)

\[ S_3 = A_1 + A_2 + A_2, \]

(21)

\[ A_1 = 1 - A_1 - A_2, \]

(22)

\[ L_1 = \lambda_1 + \lambda_3, \]

(23)

\[ L_1 = \lambda_1^2 + \lambda_1 + \lambda_2, \]

(24)

\[ L_1 = \lambda_1^2 + \lambda_2 + \lambda_3, \]

(25)
Thus, duration of the ON, OFF, and PAUSE states can be calculated as:

\[ T(\text{OFF}) = \frac{1}{k_i^+} \]  

(26)

\[ T(\text{PAUSE}) = \frac{1}{k_i^+ + k_i^-} \]  

(27)

\[ T(\text{ON}) = \frac{1}{k_i^- + k_i^-} \]  

(28)

For this model, the steady-state probability to be in a given promoter state is

\[ P_{\text{OFF}} = \frac{k_i^- k_i^+}{k_i^- k_i^+ + k_i^- k_i^+ + k_i^- k_i^+} \]  

(29)

\[ P_{\text{PAUSE}} = \frac{k_i^+ k_i^-}{k_i^- k_i^+ + k_i^- k_i^- + k_i^+ k_i^-} \]  

(30)

\[ P_{\text{ON}} = \frac{k_i^- k_i^-}{k_i^- k_i^+ + k_i^- k_i^- + k_i^+ k_i^-} \]  

(31)

The alternative three-state model with systematic pause (Supplementary Fig. 5) satisfies the following relation among distribution parameters:  

\[ A_1 \lambda_1 + A_2 \lambda_2 + (1 - A_1 - A_2) \lambda_3 = 0. \]  

(32)

This means that only four and not five distribution parameters are free, which further constrains the three-exponential fittings. In order to infer this model, a constrained fitting was performed but the bad quality of fitting recommended rejection of the model (Supplementary Fig. 5).

Testing the method with artificial data. The entire computational pipeline was tested using artificial data (see also Tantale et al.\textsuperscript{39}). Artificial traces were generated by simulating the model using the Gillespie algorithm with parameter sets similar to those identified from data. The simulations generated artificial polymerase positions, from which a first version of the signal was computed by convolution. The results are provided in Fig. 2e–h.

A modified Kolmogorov–Smirnov test for the parametric distribution. A one-sided one-sample Kolmogorov–Smirnov (KS) test was used to check if the waiting times follow the parametric fitted distribution. The output of this test is a P value that is large if the waiting times follow the fitted distribution, and small if not. The KS test is based on differences between estimated and empirical probabilities, being thus sensible to errors in the larger probabilities of shorter waiting times, and less sensitive to rare, very long waiting times. Our fitting procedure combines linear and logarithmic scales to find a balance between short and long time scales (see Tantale et al.\textsuperscript{39}). Although relatively small (in terms of the objective function), the resulting errors on short times are large enough to be considered significant by the KS test for all models. In order to be able to distinguish between models, we have limited the analysis to times larger than 10–20 s.

Error intervals. Distribution parameters result from multixponential regression fitting using gradient methods with multiple initial data. These optimization methods provide the best fit (global optimum) but also suboptimal parameter values. Using an overfit ratio (a number larger than one, in our case 2) to restrict the number of suboptimal solutions, we define boundaries of the error interval as the minimum and maximum parameter value compatible with an objective function less than the best-fit times the overflow.

Mathematical modeling of postmitotic gaps. The distribution of the postmitotic gap was estimated using the equation below. The fitting suggests that the timescale parameters of the postmitotic gap are even, i.e., \( \lambda_i = \lambda_{i'} \). In this limit, the five parameters, three-exponential distribution defined by the equation below, becomes the simpler, three parameters mixed gamma distribution described as

\[ S(t) = p_1 e^{-t} + p_2 \left( 1 - \gamma \left( \frac{a}{b} \right) \right) \left( 1 - p_3 + p_3 \left( 1 - \gamma \left( \frac{a}{b} \right) \right) \right) \]  

(33)

where

\[ \gamma(x,y) = \int_0^x s^{x-1} e^{-s} ds \]  

(34)

is the lower incomplete gamma function, and \( p_1, p_2, p_3, a, b \) are the probabilities of one step, two steps, and the mean step duration, respectively.

Chromatin immunoprecipitation. Homozygous embryos were collected and fixed in 1.4% formaldehyde for 20 min prior to dry storage at \(-80^\circ C\). Embryos were dissociated on ice in RIPA buffer (150 mM NaCl, 1% IGEPAL CA-630, 0.5% sodium deoxycholate, 0.1% SDS, 50 mM Tris-HCl pH 7.8) supplemented with protease and phosphatase inhibitors (Roche). Chromatin shearing was performed in a pre-chilled Bioruptor Pico (Diagenode) for five cycles of 30 s/30 s. Samples were divided into equal volumes and incubated overnight at 4°C with either rabbit anti-Rbp3 (10 μg) or rabbit anti-Nel-F (10 μg) (gifts from J. Zeitlinger\textsuperscript{28}) or normal serum. A second incubation overnight at 4°C with Protein G-magnetic Sepharose beads (GE Healthcare) was used to pull down protein:DNA complexes prior to washing in low salt (120 mM NaCl, 0.1% SDS, 0.5% Triton X-100, 20 mM Tris-HCl pH 8.0) and high salt (500 mM NaCl, 0.1% SDS, 0.8% Triton X-100, 20 mM Tris-HCl pH 8.0) buffers, elution, incubation with protease K and RNase A, and DNA retrieval using the QIAquick PCR cleanup kit (Qiagen). qPCR analysis was performed using Light Cycler 480 SYBR Green I Master Mix (Roche) using primers listed in Supplementary Table 5. Analysis was performed using Microsoft Excel and Prism (Graphpad v8.0.1) with a Mann–Whitney test to determine significance.

Quantitative RT-PCR. To test RNAi-mediated knockdown of Nelf-A expression, 0–2 h embryos were homogenized in Triozl (Invitrogen) and RNA was extracted per the manufacturer’s directions. Reverse transcription was performed using the Superscript IV system (Invitrogen) with oligo(dT)\textsuperscript{20} priming prior to qPCR analysis. nos-GAL4-VP16; UASwhite RNAi embryos were used as the control and all measurements were performed in biological and technical triplicate. qPCR analysis was performed using Light Cycler 480 SYBR Green I Master Mix (Roche) using primers listed in Supplementary Table 5. Analysis was performed using Microsoft Excel and Prism (Graphpad 8.0.1) with a one-tailed Student’s t test to determine significance.

Image analysis software. Live imaging analysis software is available at http://www.igmml.cnrs.fr/segment-track/ along with a video tutorial. Code for the mathematical analysis of burst parameters and multixponential regression fitting is available\textsuperscript{39}.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. The data that support this study is available from the corresponding author upon reasonable request. Source data are provided with this paper.

Received: 4 August 2020; Accepted: 31 March 2021; Published online: 23 July 2021

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Acknowledgements
We thank Julia Zeitlinger, Jean-Christophe Andrau, Jeremy Dufourt, and all members of the Lagha lab for their critical reading of the manuscript and constructive discussions. We are grateful to F. Juge for the insightful discussion and H. Faure-Gautron for technical assistance. We acknowledge the MRI imaging facility, a member of the national infrastructure France-BioImaging supported by the French National Research Agency (ANR-10-INBS-04, «Investments for the future»). This work was supported by the ERC SyncDev starting grant to M.L and CNRS; by CNRS grant PEPS MIGHTY to O.R. and E. B., and by ANRS grants ECTZ62561 to E.B. O.R. acknowledges support from the French National Research Agency (ANR-17-CE40-0036, project SYMBIONT). Machine-learning calculations were performed on the HPC facility MESO@LR run by the University of Montpellier.

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Conception and design: M.L.; acquisition of the data: M.D., V.P., and C.F.; analysis: M.L., M.D., V.P., and O.R.; software: A.T. and O.R.; modeling: O.R.; interpretation of the data: M.L., M.D., V.P., and O.R.; writing: V.P. and M.L. with input from O.R. and E.B.; visualization: M.D., V.P., and O.R.; supervision: M.L.; fund acquisition: M.L. O.R. and E.B. conceived and developed the machine-learning deconvolution procedure and the facultative model of paused polymerase. All authors read the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-021-24461-6.
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Peer review information Nature Communications thanks David Suter and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Peer reviewer reports are available.

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